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# Nonmydriatic Ultra-Wide-Field Scanning Laser Ophthalmoscopy (Optomap) versus Two-Field Fundus Photography in Diabetic Retinopathy

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## Abstract

The purpose of this study was to investigate the diagnostic properties of a 2-laser wavelength nonmydriatic 200° ultra-wide-field scanning laser ophthalmoscope (SLO) versus mydriatic 2-field 45° color fundus photography (EURODIAB standard) for assessing diabetic retinopathy (DR). A total of 143 consecutive eyes of patients with different levels of DR were graded regarding DR level and macular edema based on 2-field color photographs or 1 Optomap Panoramic 200 SLO image. All SLO images were nonmydriatic and all photographs mydriatic. Grading was performed masked to patient and clinical data. Based on photography, 20 eyes had no DR, 44 had mild, 18 moderate and 42 severe nonproliferative DR, and 19 eyes had proliferative DR. Overall correlation for grading DR level compared to Optomap SLO was moderate with kappa 0.54 ( $p < 0.001$ ), fair-to-moderate in macular edema grading with kappa 0.39 ( $p < 0.001$ ), and substantial for grading clinically significant macular edema (kappa 0.77). The wide-field SLO offers a wider field of view and can potentially better differentiate lesions by applying the 2 laser wavelengths. However, these advantages over 2-field fundus photography need to be confirmed in further studies.

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## Introduction

Diabetic retinopathy (DR) and particularly diabetic macular edema (DME), microvascular complications of diabetes mellitus, are leading causes of visual impairment in developed countries. According to the World Health Organization (WHO) almost 2% of the world population is blind due to DR, and 10% are visually disabled [1]. Taking the rise from 170 million people affected by diabetes mellitus in 2000 to an estimated 366 million in 2030 into account, the absolute number of patients suffering from vision deterioration due to DR complications will increase significantly [2]. After 20 years all people with insulin-dependent diabetes mellitus and 60% with insulin-independent diabetes mellitus have DR [3].

DME is the most common cause for vision decrease and the presence of DME increases the risk for this by 30–50% [4, 5]. With the introduction of VEGF-targeted intravitreal therapeutic approaches, an even earlier detection of the DR grade including the presence of DME is needed.

Over the last decades many efforts to develop appropriate screening tools and protocols have been made. The rationale for the development of nonmydriatic screening

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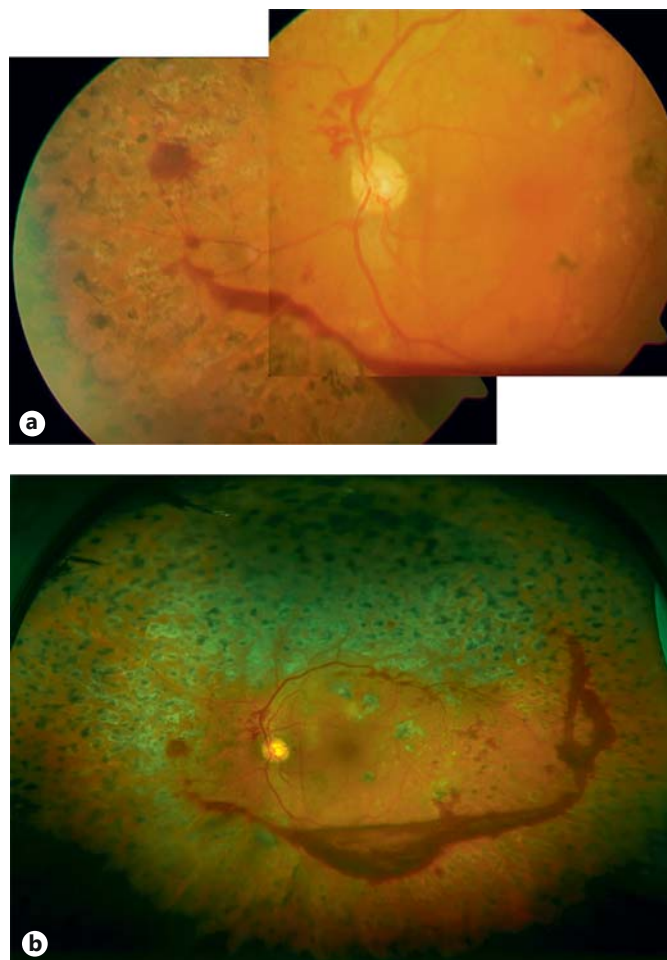
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techniques based on fundus photography and telemedicine programs included patient's comfort and needs (particularly in many screening programs in developed countries), as well as user-friendly application and handling in rural areas (where there is less availability of an ophthalmological expert) and the need for teleophthalmological networks.

The recognized 'gold standard' in obtaining retinal photographs is the Early Treatment Diabetic Retinopathy Study (ETDRS) standard 7-field 30° color fundus photography [6]. However, although ETDRS photography and grading provide a reliable and proven method of documentation and assessment of DR-related fundus alterations, it is a time-consuming examination requiring skilled photographers and pharmacological pupil dilation. The complexity of this single examination might be one of the reasons why only an estimated 60% of the diabetic population in the USA receives recommended annual eye screenings [7].

In order to make this procedure more patient and user friendly several alterations in obtaining fundus photographs, including nonmydriatic cameras, digital video imaging and fewer-field photography, have been evaluated [8–10]. Among these, particularly 2-field 45° retinal photographs (1 macular-centered field and 1 disc/nasal field), are a well-established and validated screening standard as used and proposed in the EURODIAB IDDM complications study [11].

Ultra-wide-field scanning laser ophthalmoscopy (SLO) is a novel nonmydriatic fundus imaging device (Optomap Panoramic 200; Optos plc., Dunfermline, Scotland, UK) which allows imaging not only of the posterior pole of the retina in undilated pupils, but even extending over the equator. It covers 180–200° with no need for pupil dilation, which has theoretical advantages over standard fundus photography. Moreover, it is well known that the SLO technique is less susceptible to media opacities (especially cataracts) [12] and to a decrease in pupil diameter. On the other hand, SLO optical resolution is limited and has numerically less pixels than fundus photography – but it yields a higher contrast [13]. We have previously shown that the Optomap SLO performance in assessing DR can compete with clinical examination [13]. However, assessment by photographic reading center grading is known to be superior to clinical examination [14, 15]. In this patient series, we therefore sought to assess the value of wide-field SLO in DR screening application versus photography. This study compares the diagnostic properties of nonmydriatic 200° ultra-wide-field SLO (Optomap) versus a photographic DR screening standard of two 45° mydriatic fundus photographs.



**Fig. 1.** Example of 2-field photography (a) and corresponding wide-field SLO imaging (b) of a patient. Despite panretinal laser coagulation a preretinal hemorrhage can be seen at the lower temporal vessel arcade with significant neovascularization. On wide-field SLO imaging some lashes (lower part of the images) and parts from imaging optics (left and right periphery) are present.

## Methods

### Patients

Consecutive patients were recruited from the outpatient clinic of the Department of Ophthalmology, Ludwig-Maximilians-University, Munich. Patients were included if they had diabetes (based on WHO criteria) for at least 3 years. Eyes were excluded if there were eye diseases involving the posterior pole other than DR (such as age-related macular degeneration), but not for media opacities. Graders did not participate in clinical examination of the patients and had no access to clinical data.

### Optomap Imaging

After informed consent, Optomap imaging was performed without pupil dilation before, and independently of, the clinical examination. The study conformed to the principles expressed in

**Table 1.** Agreement in DR grading based on EURODIAB photography versus Optomap fundus photographs

Optomap grading	Photography (EURODIAB) classification of DR					Total
	no DR	mild NPDR	moderate NPDR	severe NPDR	PDR	
No DR	17	5	0	0	1 <sup>1</sup>	23
Mild NPDR	3	26	2	0	1 <sup>1</sup>	32
Moderate NPDR	0	8	11	9	0	28
Severe NPDR	0	5	4	27	6	42
PDR	0	0	1	6	11	18
Total	20	44	18	42	19	143

EURODIAB photography-based grading of DR vs. assessment from Optomap images. The numbers give the number of eyes in each category. Overall correlation for DR level grading compared to Optomap was moderate with kappa of 0.54 ( $p < 0.001$ ).

<sup>1</sup> Both cases were verified to have no active PDR in regrading and in clinical examination: 1 patient had a fibrosed neovascularization of the disc – while the fibrosis and panretinal laser photocoagulation was not imaged in 2-field photography. Figure 2 shows the corresponding images. The other patient had no DR but due to low imaging quality in photography a disc neovascularization was suspected.

the Declaration of Helsinki and approval was obtained from the Institutional Review Board. Optomap imaging consisted of taking several images, where the best image per eye was saved on the server for grading. The instrument takes 1 image in approximately 0.25 s, thus avoiding motion artifacts. Total scanning time approximates 3–5 min including patient positioning and was performed by one of the authors (K.L.). Basic operation of the Optomap Panoramic 200 (an SLO) involves scanning with 2 laser wavelengths: 1 green (532 nm) and 1 red (633 nm) laser. The 2 images are then either viewed separately or superimposed by the software to yield semirealistic color imaging (see fig. 1). The instrument requires a small optical path of only 2 mm and by a special mirror design is able to obtain wide-field images of approximately 180–200° without pupil dilatation. The optical resolution with the instrument used in this study was  $3,900 \times 3,072$  pixels for that angle, resulting in approximately 15–21 pixels per degree. Due to the SLO principle, images of high contrast and sharpness were obtained [12].

#### Two-Field 45° Fundus Photography

Prior to retinal photography, the patient's pupils were dilated using tropicamide 1% and additional epinephrine 10% if required – repeated if pupils did not reach at least 6 mm in diameter. Color retinal photographs, with a suitable high-quality retinal digital camera (Zeiss FF450; Carl Zeiss Meditec AG, Jena, Germany) were taken by a certified ophthalmic photographer. A 5.0-megapixel charge-coupled device sensor (Sony 3CCD; Sony, Tokyo, Japan) was utilized in this study. Focusing and alignment of the image were performed using the ocular tube of the camera in addition to a previewing camera. The two 45° fields, according to EURO-

DIAB [11], were obtained as follows: (1) macular field: positioned such that the exact center of the optic disc lay at the nasal end of the horizontal meridian of the field of view and (2) disc/nasal field: such that the optic disc was positioned 1 disc diameter in from the temporal edge of the field, on the horizontal meridian.

#### Grading of Images

All retina images were loaded from the server to a viewing station (equipped with a conventional cathode ray 17-inch noncalibrated color monitor) via network and assessed with the Optomap viewing software (Optomap U-revu, version 1.0). This software allows basic image manipulations such as changing contrast and brightness and zooming. It also offers viewing both in the composite color image and the single-color wavelengths. The images obtained by the different wavelengths were utilized to better identify and differentiate lesions (especially red-free image [16]). The grader (L.C.) had not participated in the examination of the patients and was masked to all additional information such as visual acuity, duration of diabetes or clinical symptoms. The grader, however, could decide not to grade due to insufficient image quality, which was defined as not covering at least the central 60° and both the macula and the optic disc in adequate quality. Nongradable images were reassessed by a second grader (A.S.N.) to reach consensus on gradability. The level of DR and macular edema were assessed using the International Clinical Diabetic Retinopathy (ICDR) Severity Scale [17]. Independently of this assessment the presence of clinically significant macular edema (CSME) was graded according to ETDRS classification [18].

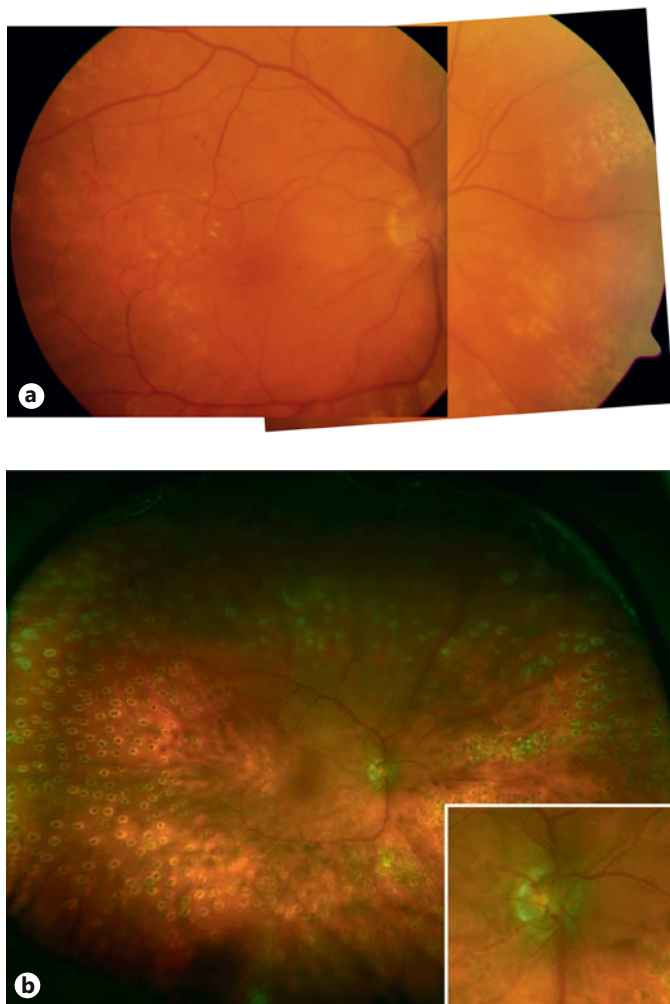
#### Statistics

All data were collected in an MS-Excel 2000 spreadsheet (Microsoft Corporation, Redmond, Wash., USA) and analyzed using SPSS 17.0 for Windows (SPSS Inc., Chicago, Ill., USA). In all tests  $p < 0.05$  was considered significant. Kappa statistics were calculated and assessed as proposed by Altman [19]: <0.20 poor, 0.21–0.40 fair, 0.41–0.60 moderate, 0.61–0.80 substantial and 0.81–1.00 almost perfect strength of agreement. Unweighted kappa was used, thus avoiding any bias by weighting.

#### Results

A total of 143 eyes, 74 of which were left eyes, of 74 patients were included in the study. Mean ( $\pm$  SD) patient age was  $60 \pm 12.1$  years (range 24–75 years). Diabetes duration ranged from 1 to 31 years (mean  $13.4 \pm 10.1$  years). Of all patients, 53% were using insulin while 47% were on oral medication. Mean HbA<sub>1c</sub> was  $7.51 \pm 1.46\%$  (range 8–10%).

Grading of DR based on the international classification scale is presented in table 1. According to the 45° photos, 20 (13.9%) eyes had no DR, 44 (30.8%) had mild, 18 (12.6%) moderate and 42 (29.4%) severe nonproliferative DR (NPDR), and 19 (13.2%) eyes had proliferative DR (PDR). The strength of agreement between these 2 imaging methods was moderate with a kappa statistic of 0.54 ( $p < 0.001$ ). For the 2 cases not identified as PDR by



**Fig. 2.** Example of 2-field photography (a) and corresponding wide-field SLO imaging (b) of a patient with PDR and fibrotic neovascularization at the optic disc. The patient was falsely graded to be active PDR based on fundus photography (a), while SLO imaging clearly showed the fibrosis, especially when viewed at higher magnification as shown in the lower right inset (b).

Optomap imaging (table 1), these were confirmed clinically and on regrading were determined to be nonproliferative. Figure 2 illustrates 1 case.

Based on the international clinical classification EURODIAB photography yielded macular edema absent in 61 eyes (42.6%), mild in 62 (43.3%), moderate in 14 (9.7%) and severe in 6 eyes (4.2%; table 2). Concordance regarding macular edema grading was fair-to-moderate with a kappa value of 0.39 ( $p < 0.001$ ). Table 2 shows results and corresponding Optomap grading in detail. On photography, CSME was graded to be present in 38 eyes (26.6%), while 105 (73.4%) of the eyes did not have CSME. Table 3

**Table 2.** Concordance in grading severity of DME according to international classification based on EURODIAB photography versus Optomap

Optomap grading	Photography (EURODIAB) classification of DME				Total
	no ME	mild ME	moderate ME	severe ME	
No ME	54	21	2	0	77
Mild ME	5	21	4	0	30
Moderate ME	1	16	7	3	27
Severe ME	1	4	1	3	9
Total	61	62	14	6	143

The numbers give the number of eyes in each category. Concordance regarding DME grading was fair-to-moderate with a kappa of 0.39 ( $p < 0.001$ ). ME = Macular edema.

**Table 3.** Concordance in grading CSME based on EURODIAB photography versus Optomap

Optomap grading	Photography (EURODIAB) assessment of CSME		
	CSME present	no CSME	Total
CSME present	18	9	27
No CSME	20	96	116
Total	38	105	143

The numbers give the number of eyes graded in each category. Sensitivity: 48%; specificity: 91%.

shows corresponding Optomap results. Both imaging methods have substantial agreement in the detection and evaluation of CSME (kappa value 0.77).

## Discussion

The current study demonstrates that nonmydriatic ultra-wide-field SLO (Optomap) correlates moderately with mydriatic 2-field 45° photography. This adds to previous data, which have shown validity of assessments compared to clinical fundus examination and a lower rate of nongradable images for Optomap [13]. In an environment that demands cost reduction, practice efficiency and effective disease screening, nonmydriatic digital fundus screening offers many advantages. One important advantage inherent to nonmydriatic digital

techniques is patient comfort (no need for dilating pupils with inability to drive a car) and facilitation of remote diagnostic image generation and interpretation. These factors are likely to improve patient preference and compliance in screening programs [20]. Another benefit of the Optomap nonmydriatic camera is its use of the SLO technique, as the image quality in opaque media is significantly better compared to standard cameras and thicker cataracts are more often seen in low-income and rural areas [21]. Additionally, remote interpretation could permit screening of underserved and low-income rural and urban populations at risk for DR [15]. The Digital Diabetic Screening Group suggests that a nonmydriatic digital photography protocol is superior to mydriatic ophthalmoscopy and roughly comparable to 7-field ETDRS mydriatic photography.

Most fundus camera-based screening systems for DR take one or two 45° or 60° images. This has been validated against the best known standard for fundus photography in DR, the 7-field ETDRS photography, which covers 75–65° of the central retina by montaging seven 30° photographs. The ETDRS standard is known to have good validity – given the fact that photographic grading was validated against long-term outcomes [22] – but is difficult to perform and relatively unsuitable for routine. While two 60° fundus photographs (1 macula-centered and 1 optic disc-centered) cover 80% of this area and make it unlikely that areas of neovascularization will be missed [23], most screening programs today cover significantly less retinal area. These programs still offer good screening characteristics [24], although the relatively small covered area may theoretically reduce sensitivity. However, these standards appear to be sufficient for screening purposes as carried out elsewhere when the standard using two 45° photographs, as in our study (EURODIAB protocol), was validated successfully against 7-field ETDRS photography, the diagnostic gold standard [11]. Without any doubt, a larger field of view offers better detection characteristics; however, coverage of the nasal retina appears to be more important than the field of view in total. In terms of area coverage, the ultra-wide-field SLO investigated here (see fig. 1) clearly exceeds the ETDRS standard, which could offer additional advantages. For instance, in 1 of the cases falsely classified as PDR based on photography (table 1), Optomap imaging quality and field of view could exclude proliferations.

The SLO technique showed only a fair-to-moderate agreement with photography regarding macular edema grading (tables 2, 3). It should be noted though that non-stereoscopic fundus photography is equally limited re-

garding detection of thickening as none of these imaging modalities create 3-dimensional photographs. Therefore, although a discrepancy between Optomap and standard photography (EURODIAB) graded pictures exist (table 3), these results have to be considered with caution as standard photographs do not represent a common standard in diagnosing macula edema. The results are particularly surprising since a previous study comparing Optomap photographs to stereo biomicroscopy with three independent graders revealed a specificity of 100% and a sensitivity of 94% compared to only 48% specificity and 91% sensitivity in comparison to standard photography in this study [13] (table 3). Another study compared stereoscopic digital fundus photography against contact lens biomicroscopy for the detection of CSME with a specificity of around 50–90% and a sensitivity of 90–99%, a result very comparable to ours when comparing Optomap against standard photography [25]. Taken together, there is no doubt that photographic assessment is suitable for detecting sight-threatening diseases and the course of these diseases (as in PDR), regardless of whether it is based on standard photography or SLO technique. Photography, however, cannot reliably detect 3-dimensional changes (as in macular edema). Therefore, grading of macular thickening always requires stereoscopic assessment by biomicroscopy and/or a cross-sectional technique such as OCT. Considering these different results in terms of the ability of photographic assessment for determining CSME, the real value of solely grading fundus images with regard to CSME by Optomap cannot be fully determined.

Limitations of the Optomap system have been described earlier and include misdiagnosing and missing lesions, which were not relevant in this study [26]. The percentage of nongradable images, which is a relevant marker in screening applications, cannot be assessed in this report. Only patients receiving both imaging technologies were included and in those no eyes had to be excluded for nongradable images. While this is a limitation of our study, in a similar patient series evaluated previously for Optomap nongradable and ‘nonobtainable’ images, the numbers were approximately <10% of all patients [13].

In summary, the nonmydriatic ultra-wide-field SLO images are of sufficient quality to assess the DR level validly. Compared to a screening standard (EURODIAB) of two 45° photographs, inter-method agreement was only moderate. The additional field of view from Optomap may offer additional advantages and yield other incidental fundus abnormalities.

## Acknowledgments

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## Disclosure Statement

The authors have no commercial interest in any of the materials and methods used in this study.

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