BMJ Open Diabetes Research & Care

Corneal confocal microscopy compared with quantitative sensory testing and nerve conduction for diagnosing and stratifying the severity of diabetic peripheral neuropathy

Maryam Ferdousi, ¹ Alise Kalteniece, ¹ Shazli Azmi, ¹ Ioannis N Petropoulos, ² Anne Worthington, ¹ Luca D'Onofrio, ³ Shaishav Dhage, ¹ Georgios Ponirakis ⁶, ⁴ Uazman Alam, ¹ Andrew Marshall, ¹ Catharina G Faber, ⁵ Giuseppe Lauria, ⁶ Handrean Soran, ^{1,7} Rayaz A Malik ¹, ^{1,2}

To cite: Ferdousi M,
Kalteniece A, Azmi S, et al.
Corneal confocal microscopy
compared with quantitative
sensory testing and nerve
conduction for diagnosing
and stratifying the severity of
diabetic peripheral neuropathy.
BMJ Open Diab Res Care
2020;8:e001801. doi:10.1136/

Received 30 July 2020 Revised 28 October 2020 Accepted 21 November 2020



© Author(s) (or their employer(s)) 2020. Re-use permitted under CC BY-NC. No commercial re-use. See rights and permissions. Published by BMJ.

For numbered affiliations see end of article.

Correspondence to

Dr Rayaz A Malik; ram2045@qatar-med.cornell.

ABSTRACT

Introduction Diabetic neuropathy can be diagnosed and assessed using a number of techniques including corneal confocal microscopy (CCM).

Research design and methods We have undertaken quantitative sensory testing, nerve conduction studies and CCM in 143 patients with type 1 and type 2 diabetes without neuropathy (n=51), mild neuropathy (n=47) and moderate to severe neuropathy (n=45) and age-matched controls (n=30).

Results Vibration perception threshold (p<0.0001), warm perception threshold (WPT) (p<0.001), sural nerve conduction velocity (SNCV) (p<0.001), corneal nerve fiber density (CNFD) (p<0.0001), corneal nerve branch density (CNBD) (p<0.0001), corneal nerve fiber length (CNFL) (p=0.002), inferior whorl length (IWL) (p=0.0001) and average nerve fiber length (ANFL) (p=0.0001) showed a progressive abnormality with increasing severity of diabetic neuropathy. Receiver operating characteristic curve analysis for the diagnosis of diabetic neuropathy showed comparable performance in relation to the area under the curve (AUC) but differing sensitivities and specificities for vibration perception threshold (AUC 0.79, sensitivity 55%, specificity 90%), WPT (AUC 0.67, sensitivity 50%, specificity 76%), cold perception threshold (AUC 0.64, sensitivity 80%, specificity 47%), SNCV (AUC 0.70, sensitivity 76%, specificity 54%), CNFD (AUC 0.71, sensitivity 58%, specificity 83%), CNBD (AUC 0.70, sensitivity 69%, specificity 65%), CNFL (AUC 0.68, sensitivity 64%, specificity 67%), IWL (AUC 0.72, sensitivity 70%, specificity 65%) and ANFL (AUC 0.72, sensitivity 71%, specificity 66%).

Conclusion This study shows that CCM identifies early and progressive corneal nerve loss at the inferior whorl and central cornea and has comparable utility with quantitative sensory testing and nerve conduction in the diagnosis of diabetic neuropathy.

INTRODUCTION

Corneal confocal microscopy (CCM) has been used to identify nerve fiber degeneration in

Significance of this study

What is already known about this subject?

Corneal nerve loss may allow the diagnosis of diabetic neuropathy.

What are the new findings?

- There is progressive loss of corneal nerves at the central and inferior whorl region with increasing severity of diabetic neuropathy.
- Inferior whorl length was reduced while central corneal nerve fiber length was still preserved in patients without diabetic neuropathy, consistent with an early distal axonopathy.
- Corneal confocal microscopy has comparable utility with quantitative sensory testing and nerve conduction studies in the diagnosis of diabetic neuropathy.

How might these results change the focus of research or clinical practice?

 Corneal confocal microscopy may be used to diagnose and stratify the severity of diabetic neuropathy.

patients with diabetic neuropathy. 1-5 It is a rapid, non-invasive nerve imaging technique with high reproducibility and moderate to high sensitivity and specificity in the diagnosis of diabetic peripheral neuropathy (DPN). 36-10 To date, most studies have undertaken central corneal nerve assessment. However, recent animal and small human studies suggest that quantification of corneal nerve morphology at the inferior whorl may detect earlier nerve fiber damage. 11-14 Given that DPN is a lengthdependent distal axonopathy, nerves at the inferior whorl are expected to be affected before central corneal nerves. 15 Indeed, we have previously shown greater corneal nerve loss at the inferior whorl compared with the



BMJ Open Diab Res Care: first published as 10.1136/bmjdrc-2020-001801 on 21 December 2020. Downloaded from http://drc.bmj.com/ on January 1, 2021 by guest. Protected by copyright

central cornea, 13 14 16 even though central corneal nerve fiber length (CNFL) is considered to be optimal for diagnosing DPN. 17-19 While Pritchard et al 13 reported a comparable diagnostic utility for central and inferior whorl corneal nerve length, Petropoulos et al¹⁴ showed that inferior whorl length (IWL) increased the sensitivity in diagnosing DPN. More recently, Kalteniece et al^{16} combined CNFL and IWL and showed that it was more reliable than either alone, in identifying patients with diabetic neuropathy. Corneal nerve loss in larger images incorporating nerves from the center and inferior whorl has been demonstrated in patients with recently diagnosed type 2 diabetes.²⁰ Furthermore, in our recent longitudinal study, a reduction in IWL occurred more rapidly than CNFL in patients with diabetic neuropathy.²¹

We have compared the utility of quantifying corneal nerve loss at the inferior whorl and central cornea with quantitative sensory testing and nerve conduction in the diagnosis and assessment of the severity of DPN.

RESEARCH DESIGN AND METHODS Study participants

One hundred and forty-three patients with diabetes and 30 control participants underwent a comprehensive assessment of peripheral neuropathy and CCM. Each participant provided informed consent prior to participation in the study.

Patients were excluded if they had a history of connective tissue or infectious disease, malignancy, deficiency in B₁₀ or folate, chronic renal and liver failure, current or active diabetic foot ulceration, previous ocular trauma, systemic disease other than diabetes that could cause neuropathy or affect the cornea, corneal surgery, and a history of or current contact lens wear.

Clinical and peripheral neuropathy assessment

Each participant underwent assessment of body mass index (BMI), blood pressure, glycated hemoglobin (HbA1c) and lipid profile. The simplified Neuropathy Disability Score (NDS), which assesses vibration, pinprick, temperature perception, and presence or absence of ankle reflexes, was used to stratify patients into three groups: no (NDS=0-2), mild (NDS=3-5) and moderate to severe (NDS=6–10) neuropathy.²²

Vibration perception threshold (VPT) was assessed using a neurothesiometer (Scientific Laboratory Supplies, Wilford, Nottingham, UK), and cold perception threshold (CPT) and warm perception threshold (WPT) were tested on the dorsolateral aspect of the nondominant foot (S1) using a TSA-II NeuroSensory Analyzer (Medoc, Ramat Yishai, Israel). Electrodiagnostic studies were undertaken by a consultant neurophysiologist using a Dantec 'Keypoint' system (Dantec Dynamics, Bristol, UK) equipped with a thermistor (DISA temperature regulator, Denmark) to keep the limb temperature at 32°C-35°C and sural nerve conduction velocity (SNCV) was tested.

Ophthalmic assessment

Examinations of the anterior ocular segment using slit-lamp biomicroscopy and CCM examination using laser scanning CCM (Heidelberg Retina Tomograph III Rostock Cornea Module; Heidelberg Engineering, Heidelberg, Germany) were performed for both eyes according to our established protocol.²³

Six images (three per eye) from the central sub-basal nerve plexus and four images (two per eye) from the inferior whorl region were selected and manually quantified using CCMetrics (The University of Manchester, Manchester, UK). Images were selected by a single expert in a masked fashion taking into account the quality, depth and variability following our previously established protocol.³ ¹⁴ Corneal nerve fiber density (CNFD; total number of main nerves per square millimeter) (no./mm²), corneal nerve branch density (CNBD; total number of branches per square millimeter) (no./mm²), CNFL (total length of main nerves and nerve branches per square millimeter) (mm/mm²), IWL (total length of nerves per square millimeter) (mm/mm²) and average nerve fiber length (ANFL=CNFL+IWL/2) (mm/mm²) were quantified.

Statistical analysis

Analysis was carried out using SPSS V.22.0 for Windows. The Shapiro-Wilk test was employed to assess whether data were normally distributed. Fisher's exact test was used to test the association between two categorical variables. Analysis of variance and analysis of covariance with least significant difference correction were used for multiple comparisons between groups. All data were expressed as mean±SEM. P<0.05 was considered significant. Graphs were created using GraphPad Prism (V.7.0c for Windows; GraphPad Software, La Jolla, California, USA). Receiver operating characteristic (ROC) curves were used to define the optimum cut-off points with the highest sensitivity and specificity in the diagnosis of DPN, and Youden's index (J=sensitivity+specificity-1) was measured.

RESULTS

Clinical, demographic and laboratory findings

All participant groups were matched for gender (p=0.4) and ethnicity (p=0.3). Age (p=0.003), duration of diabetes (p=0.01), HbA1c (p<0.0001), BMI (p=0.04) and lowdensity lipoprotein cholesterol (p<0.0001) were significantly different between all participant groups. There was no difference in high-density lipoprotein cholesterol (p=0.4) or triglycerides (p=0.7) between groups (table 1).

Peripheral neuropathy assessment

VPT was significantly higher in patients with no $(12.2\pm1.24, p=0.02), \text{ mild } (15.05\pm1.2, p<0.0001) \text{ and}$ moderate to severe (25.56±1.35, p<0.0001) neuropathy compared with controls (7.1 ± 1.80) (table 2). CPT was significantly lower only in patients with mild (19.52±1.47,



Table 1 Demographic and clinical findings in controls and in patients with diabetes with no, mild and moderate to severe neuropathy

Parameter	Controls (n=30)	No neuropathy (n=51)	Mild neuropathy (n=47)	Moderate to severe neuropathy (n=45)	ANOVA p value
Age (years)	54.51±2.3	57.68±1.6	60.16±1.7*	64.1±1.48*†	0.003
Gender (female/male)	15/15	20/31	18/29	16/29	0.4
Ethnicity (European/non-European)	23/7	42/9	31/16	36/9	0.3
Duration of diabetes (years)	0	15±2	18±2	24±3	0.01
HbA1c (mmol/mol)	36.4±0.65	56.21±2.52*	55.99±1.88*	55.37±2.05*	<0.0001
BMI (kg/m²)	27.05±0.94	29.09±0.86	28.73±1.0	31.62±1.41*	0.04
Triglycerides (mmol/L)	1.63±0.21	1.41±0.1	1.53±0.14	1.45±0.17	0.7
LDL-C (mmol/L)	2.6±0.14	2.14±0.12*	1.9±0.15*	1.7±0.08*†	<0.0001
HDL-C (mmol/L)	1.67±0.1	1.58±0.07	1.47±0.08	1.49±0.1	0.4

All data are expressed as mean±SEM.

p=0.02) and moderate to severe (18.99 \pm 1.55, p=0.01) neuropathy compared with controls (25.38 \pm 2.06). WPT was significantly higher in patients with no (41.65 \pm 0.6, p=0.01), mild (43.47 \pm 0.6, p<0.0001) and moderate to severe (43.62 \pm 0.7, p<0.0001) neuropathy compared with controls (38.87 \pm 0.9). SNCV was significantly lower in patients with no (43.14 \pm 1.56, p=0.001), mild (43.5 \pm 1.69, p=0.002) and moderate to severe (40.19 \pm 1.79, p<0.0001) neuropathy compared with controls (52.09 \pm 2.1).

Corneal confocal microscopy

CNFD was significantly lower in patients with no $(26.61\pm1.05, p<0.0001)$, mild $(24.47\pm1.09, p<0.0001)$ and moderate to severe $(22.4\pm1.14, p<0.0001)$ neuropathy

compared with controls (33.71 \pm 1.3) (figure 1, table 2). CNBD was significantly lower in patients with no (64.07 \pm 4.39, p=0.01), mild (58.49 \pm 4.76, p=0.002) and moderate to severe (45.60 \pm 4.50, p<0.0001) neuropathy compared with controls (81.52 \pm 5.54). CNFL was significantly lower in patients with mild (20.84 \pm 1.004, p=0.01) and moderate to severe (19.27 \pm 1.04, p=0.001) neuropathy compared with controls (25.07 \pm 1.27). IWL was significantly lower in patients with no (24.9 \pm 1.26, p=0.001), mild (22.28 \pm 1.31, p<0.0001) and moderate to severe (19.03 \pm 1.36, p<0.0001) neuropathy compared with controls (31.69 \pm 1.66). ANFL was significantly lower in patients with no (24.1 \pm 0.98, p=0.009), mild (21.56 \pm 1.02,

Table 2 ANCOVA with LSD correction for CCM and other measures of peripheral neuropathy in controls and in patients with diabetes with increasing severity of neuropathy

Parameters	Control (n=30)	No neuropathy (n=51)	Mild neuropathy (n=47)	Moderate to severe neuropathy (n=45)	P value
VPT (V)	7.1±1.80	12.2±1.24*	15.05±1.2*	25.56±1.35*†‡	<0.0001
CPT (°C)	25.38±2.06	21.68±1.44	19.52±1.47*	18.99±1.55*	0.06
WPT (°C)	38.87±0.9	41.65±0.6*	43.47±0.6*†	43.62±0.7*†	<0.001
SNCV (m/s)	52.09±2.1	43.14±1.56*	43.5±1.69*	40.19±1.79*	<0.001
CNFD (no./mm ²)	33.71±1.3	26.61±1.05*	24.47±1.09*	22.4±1.14*†	<0.0001
CNBD (no./mm ²)	81.52±5.54	64.07±4.39*	58.49±4.76*	45.60±4.5*†	<0.0001
CNFL (mm/mm ²)	25.07±1.27	23.31±0.96	20.84±1.004*	19.27±1.04*†	0.002
IWL (mm/mm ²)	31.69±1.66	24.9±1.26*	22.28±1.31*	19.03±1.36*†	<0.0001
ANFL (mm/mm²)	28.38±1.3	24.1±0.98*	21.56±1.02*	19.15±1.06*†	<0.0001

Data presented as marginal mean±SEM.

^{*}Significant difference compared with controls.

[†]Significant difference compared with no neuropathy.

ANOVA, analysis of variance; BMI, body mass index; HbA1c, glycated hemoglobin; HDL-C, high-density lipoprotein cholesterol; LDL-C, low-density lipoprotein cholesterol.

^{*}Significant difference compared with controls.

[†]Significant difference compared with no neuropathy.

[‡]Significant difference compared with mild neuropathy.

ANCOVA, analysis of covariance; ANFL, average nerve fiber length; CCM, corneal confocal microscopy; CNBD, corneal nerve branch density; CNFD, corneal nerve fiber density; CNFL, corneal nerve fiber length; CPT, cold perception threshold; IWL, inferior whorl length; LSD, least significant difference; SNCV, sural nerve conduction velocity; VPT, vibration perception threshold; WPT, warm perception threshold.

BMJ Open Diab Res Care: first published as 10.1136/bmjdrc-2020-001801 on 21 December 2020. Downloaded from http://drc.bmj.com/ on January 1, 2021 by guest. Protected by copyright

Figure 1 Corneal confocal microscopy images of the central cornea (top row) and the inferior whorl (bottom row) in a healthy control (first column) and in patients with no (second column), mild (third column) and moderate to severe (fourth column) neuropathy, DPN, diabetic peripheral neuropathy; NDS, Neuropathy Disability Score.

p<0.0001) and moderate to severe (19.15±1.06, p<0.0001) neuropathy compared with controls (28.38±1.3).

Diagnostic utility of different measures of DPN

The ROC curves (table 3) for quantitative sensory testing, nerve conduction and CCM for the diagnosis of diabetic neuropathy (NDS >3) (figure 2) revealed comparable area under the curve (AUC) but variable sensitivities and specificities for VPT (AUC 0.79, sensitivity 55%, specificity 90%), WPT (AUC 0.67, sensitivity 50%, specificity 76%), CPT (AUC 0.64, sensitivity 80%, specificity 47%), SNCV (AUC 0.70, sensitivity 76%, specificity 54%), CNFD (AUC 0.71, sensitivity 58%, specificity 83%), CNBD (AUC 0.70, sensitivity 69%, specificity 65%), CNFL (AUC 0.68, sensitivity 64%, specificity 67%), IWL (AUC 0.72, sensitivity 70%, specificity 65%) and ANFL (AUC 0.72, sensitivity 71%, specificity 66%).

DISCUSSION

Recommendations for the diagnosis of diabetic neuropathy require the presence of symptoms and signs and abnormal nerve conduction studies or a measure of small fiber damage if nerve conduction study (NCS) is normal.24 25 Abnormal monofilament testing and VPT reflect large nerve fiber damage and predict an increased risk of diabetic foot ulceration, 26 while thermal sensory thresholds and intraepidermal nerve fiber density identify early small nerve fiber dysfunction²⁷ and degeneration, 28 respectively. CCM is a rapid, non-invasive ophthalmic technique for the quantification of small nerve fiber damage in DPN. 29 30 Studies have shown a relationship between central corneal nerve loss and the severity of diabetic neuropathy²⁰ 31-34 with good sensitivity and specificity for the diagnosis of DPN. 19 33 In a small early study using wide-field mapping in patients

Table 3 Diagnostic performance of quantitative sensory testing, nerve conduction and CCM in the diagnosis of diabetic peripheral neuropathy

Neuropathy measure	AUC	P value	Sensitivity (%)	Specificity (%)	Youden's cut-off point
VPT	0.79	<0.0001	55	90	17.37
WPT	0.67	<0.001	50	76	44.2
CPT	0.64	<0.002	80	47	27.3
SNCV	0.70	<0.01	76	54	45.9
CNFD	0.71	<0.0001	58	83	23.95
CNBD	0.70	< 0.0001	69	65	58.54
CNFL	0.68	<0.0001	64	67	21.6
IWL	0.72	<0.0001	70	65	24.4
ANFL	0.72	<0.0001	71	66	23.4

ANFL, average nerve fiber length; AUC, area under the curve; CCM, corneal confocal microscopy; CNBD, corneal nerve branch density; CNFD, corneal nerve fiber density; CNFL, corneal nerve fiber length; CPT, cold perception threshold; IWL, inferior whorl length; SNCV, sural nerve conduction velocity; VPT, vibration perception threshold; WPT, warm perception threshold.

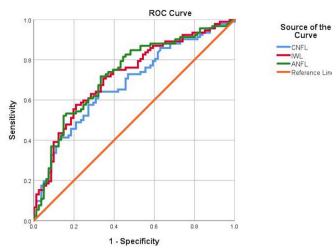


Figure 2 ROC curves for CNFL (mm/mm²), IWL (mm/mm²) and ANFL (mm/mm²). ANFL, average nerve fiber length; CNFL, corneal nerve fiber length; IWL, inferior whorl length; ROC, receiver operating characteristic.

with diabetes with and without neuropathy, we showed greater corneal nerve loss at the inferior whorl region. Subsequently, in a cohort of patients with and without DPN, we demonstrated that IWL assessment enhanced the diagnostic ability of CCM and it was more markedly reduced in patients with painful diabetic neuropathy. More recently, in a longitudinal study in patients with DPN, we have shown more rapid corneal nerve loss at the inferior whorl compared with the central cornea. Furthermore, IWL rather than CNFL reduction has been associated with increased glucose variability and time above range. The subset of the inferior whorl compared with the central corneal nervel of the inferior whorl compared with the central corneal section.

The current study demonstrates progressive loss of corneal nerves at the central and inferior whorl region with increasing severity of DPN.^{4 33} Moreover, in patients without neuropathy we show that IWL was reduced but CNFL was still preserved, consistent with an early distal axonopathy in DPN.³⁸ We also show progressive abnormality in the VPT, CPT, WPT and SNCV with increasing severity of DPN. In particular we show an early increase in the VPT of patients without DPN, which may reflect the early involvement of the Pacinian corpuscles.³⁹

For the diagnosis of DPN we show comparable AUCs for quantitative sensory testing, nerve conduction and CCM. However, the sensitivity and specificity of these tests vary and may reflect the limitations of NDS in identifying early DPN. Thus VPT had a relatively low sensitivity but high specificity, while SNCV and CPT, more precise measures of large myelinated fibers, had higher sensitivity but lower specificity. However, CNFD had comparable sensitivity and specificity with VPT, while the other measures of corneal nerve fiber loss, particularly IWL and ANFL, had better sensitivity and specificity in the diagnosis of DPN. Perkins et al^{b} assessed 998 patients with diabetes and showed comparable AUC and sensitivity and specificity in the diagnosis of DPN using the Toronto criteria. Pritchard et al¹³ have reported 90% sensitivity and 50% specificity for CNFL compared with 80% sensitivity and

60% specificity for IWL; however, they used fully automated software to analyze CCM images, which underestimates corneal nerve parameters compared with the manual analysis used in this study.

A limitation in the interpretation of our findings is that NDS is a relatively crude and subjective measure to diagnose and stratify the severity of DPN. Indeed, this may explain why the group designated as having no neuropathy already have abnormal VPT, CPT, SNCV and corneal nerve loss. Nevertheless, this is the largest study to date to demonstrate comparable utility of a range of CCM measures compared with quantitative sensory testing and nerve conduction studies, which are established measures of DPN.

Author affiliations

¹Division of Cardiovascular Sciences, Cardiac Centre, Faculty of Biology, Medicine and Health, The University of Manchester, Manchester, UK

²Research Division, Weill Cornell Medical College in Qatar, Doha, Qatar
³Department of Experimental Medicine, University of Rome La Sapienza, Roma, Lazio, Italy

⁴Medicine, Weill Cornell Medical College in Qatar, Doha, Qatar ⁵Department of Neurology, Maastricht University, Maastricht, Limburg, The Netherlands

⁶Department of Clinical Neurosciences, Università di Milano - Fondazione I.R.C.C.S. Istituto Neurologico Carlo Besta, Milan, Italy

⁷University Department of Medicine, Central Manchester University Hospitals NHS Foundation Trust, Manchester, UK

Acknowledgements The study was supported by the Manchester Biomedical Research Centre and the Greater Manchester Comprehensive Local Research Network. Dr Mitra Tavakoli undertook corneal confocal microscopy and Dr Hassan Fadavi undertook clinical assessment and quantitative sensory testing in a proportion of patients.

Contributors MF researched data, performed the statistical analysis and wrote the manuscript. AK researched data and wrote the manuscript. SA, INP, AW, LD, SD, GP, UA and AM researched data. CGF, GL and HS reviewed and revised the manuscript. RAM designed the study and reviewed and revised the manuscript. RAM is the guarantor of this work and had full access to the data of the study and takes responsibility for the integrity of the data and the accuracy of the data analysis. The authors alone are responsible for the content and writing of the paper.

Funding The research received funding from the European Union Seventh Framework Programme FP7/2007-2013 (n°602273), Diabetes UK (RD05/0003048), and Juvenile Diabetes Research Foundation (8-2008-362).

Competing interests None declared.

Patient consent for publication Not required.

Ethics approval The research adhered to the tenets of the Declaration of Helsinki and was approved by the NRES Committee North West - Greater Manchester East (REC reference: 14/NW/0093), North Manchester Research Ethics Committee (REC reference: 09/H1/006/38) and Central Manchester Local Research Ethics Committee (07/H1006/68).

Provenance and peer review Not commissioned; externally peer reviewed.

Data availability statement Data are available upon reasonable request. The data sets generated and analyzed during the study are available from the corresponding author on a reasonable request.

Open access This is an open access article distributed in accordance with the Creative Commons Attribution Non Commercial (CC BY-NC 4.0) license, which permits others to distribute, remix, adapt, build upon this work non-commercially, and license their derivative works on different terms, provided the original work is properly cited, appropriate credit is given, any changes made indicated, and the use is non-commercial. See: http://creativecommons.org/licenses/by-nc/4.0/.

ORCID iDs

Georgios Ponirakis http://orcid.org/0000-0002-6936-1248

Rayaz A Malik http://orcid.org/0000-0002-7188-8903

REFERENCES

- 1 Alam U, Jeziorska M, Petropoulos IN, et al. Diagnostic utility of corneal confocal microscopy and intra-epidermal nerve fibre density in diabetic neuropathy. PLoS One 2017;12:e0180175.
- 2 Chen X, Graham J, Dabbah MA, et al. Small nerve fiber quantification in the diagnosis of diabetic sensorimotor polyneuropathy: comparing corneal confocal microscopy with intraepidermal nerve fiber density. *Diabetes Care* 2015;38:1138–44.
- 3 Kalteniece A, Ferdousi M, Adam S, et al. Corneal confocal microscopy is a rapid reproducible ophthalmic technique for quantifying corneal nerve abnormalities. PLoS One 2017;12:e0183040.
- 4 Petropoulos IN, Alam U, Fadavi H, et al. Corneal nerve loss detected with corneal confocal microscopy is symmetrical and related to the severity of diabetic polyneuropathy. *Diabetes Care* 2013;36:3646–51.
- 5 Perkins BA, Lovblom LE, Bril V, et al. Corneal confocal microscopy for identification of diabetic sensorimotor polyneuropathy: a pooled multinational Consortium study. *Diabetologia* 2018;61:1856–61.
- 6 Pacaud D, Romanchuk KG, Tavakoli M, et al. The reliability and reproducibility of corneal confocal microscopy in children. *Invest Ophthalmol Vis Sci* 2015;56:5636–40.
- 7 Azmi S, Ferdousi M, Petropoulos IN, et al. Corneal confocal microscopy shows an improvement in small-fiber neuropathy in subjects with type 1 diabetes on continuous subcutaneous insulin infusion compared with multiple daily injection. *Diabetes Care* 2015;38:e3-4.
- 8 Azmi S, Ferdousi M, Petropoulos IN, et al. Corneal confocal microscopy identifies small-fiber neuropathy in subjects with impaired glucose tolerance who develop type 2 diabetes. *Diabetes Care* 2015;38:1502–8.
- 9 Pritchard N, Edwards K, Russell AW, et al. Corneal confocal microscopy predicts 4-year incident peripheral neuropathy in type 1 diabetes. *Diabetes Care* 2015;38:dc142114–5.
- 10 Petropoulos IN, Manzoor T, Morgan P, et al. Repeatability of in vivo corneal confocal microscopy to quantify corneal nerve morphology. Cornea 2013;32:e83–9.
- 11 Davidson EP, Coppey LJ, Kardon RH, et al. Differences and similarities in development of corneal nerve damage and peripheral neuropathy and in diet-induced obesity and type 2 diabetic rats. Invest Ophthalmol Vis Sci 2014;55:1222–30.
- 12 Cai D, Zhu M, Petroll WM, et al. The impact of type 1 diabetes mellitus on corneal epithelial nerve morphology and the corneal epithelium. Am J Pathol 2014;184:2662–70.
- 13 Pritchard N, Dehghani C, Edwards K, et al. Utility of assessing nerve morphology in central cornea versus Whorl area for diagnosing diabetic peripheral neuropathy. Cornea 2015;34:756–61.
- 14 Petropoulos IN, Ferdousi M, Marshall A, et al. The inferior Whorl for detecting diabetic peripheral neuropathy using corneal confocal microscopy. *Invest Ophthalmol Vis Sci* 2015;56:2498–504.
- Müller LJ, Marfurt CF, Kruse F, et al. Corneal nerves: structure, contents and function. Exp Eye Res 2003;76:521–42.
- 16 Kalteniece A, Ferdousi M, Petropoulos I, et al. Greater corneal nerve loss at the inferior whorl is related to the presence of diabetic neuropathy and painful diabetic neuropathy. Sci Rep 2018;8:3283.
- 17 Petropoulos IN, Alam U, Fadavi H, et al. Rapid automated diagnosis of diabetic peripheral neuropathy with in vivo corneal confocal microscopy. *Invest Ophthalmol Vis Sci* 2014;55:2071–8.
- 18 Papanas N, Ziegler D. Corneal confocal microscopy: recent progress in the evaluation of diabetic neuropathy. *J Diabetes Investig* 2015;6:381–9.

- 19 Ahmed A, Bril V, Orszag A, et al. Detection of diabetic sensorimotor polyneuropathy by corneal confocal microscopy in type 1 diabetes: a concurrent validity study. *Diabetes Care* 2012;35:821–8.
- 20 Ziegler D, Papanas N, Zhivov A, et al. Early detection of nerve fiber loss by corneal confocal microscopy and skin biopsy in recently diagnosed type 2 diabetes. *Diabetes* 2014;63:2454–63.
- 21 Ferdousi M, Kalteniece A, Petropoulos I, et al. Diabetic neuropathy is characterized by progressive corneal nerve fiber loss in the central and inferior Whorl regions. *Invest Ophthalmol Vis Sci* 2020;61:48.
- 22 Young MJ, Boulton AJ, MacLeod AF, et al. A multicentre study of the prevalence of diabetic peripheral neuropathy in the United Kingdom Hospital clinic population. *Diabetologia* 1993;36:150–4.
- 23 Tavakoli M, Malik RA. Corneal confocal microscopy: a novel non-invasive technique to quantify small fibre pathology in peripheral neuropathies. *J Vis Exp* 2011;47. doi:10.3791/2194. [Epub ahead of print: 03 Jan 2011].
- 24 Tesfaye S, Boulton AJM, Dyck PJ, et al. Diabetic neuropathies: update on definitions, diagnostic criteria, estimation of severity, and treatments. *Diabetes Care* 2010;33:2285–93.
- 25 Pop-Busui R, Boulton AJM, Feldman EL, et al. Diabetic neuropathy: a position statement by the American diabetes association. *Diabetes Care* 2017;40:136–54.
- 26 Abbott CA, Vileikyte L, Williamson S, et al. Multicenter study of the incidence of and predictive risk factors for diabetic neuropathic foot ulceration. *Diabetes Care* 1998;21:1071–5.
- 27 Themistocleous AC, Ramirez JD, Serra J, et al. The clinical approach to small fibre neuropathy and painful channelopathy. *Pract Neurol* 2014;14:368–79.
- 28 Javed S, Petropoulos IN, Tavakoli M, et al. Clinical and diagnostic features of small fiber damage in diabetic polyneuropathy. Handb Clin Neurol 2014;126:275–90.
- 29 Malik RA, Kallinikos P, Abbott CA, et al. Corneal confocal microscopy: a non-invasive surrogate of nerve fibre damage and repair in diabetic patients. *Diabetologia* 2003;46:683–8.
- 30 Petropoulos IN, Ponirakis G, Khan A, et al. Corneal confocal microscopy: ready for prime time. Clin Exp Optom 2020;103:265–77.
- 31 Quattrini C, Tavakoli M, Jeziorska M, et al. Surrogate markers of small fiber damage in human diabetic neuropathy. *Diabetes* 2007:56:2148–54
- 32 Kallinikos P, Berhanu M, O'Donnell C, et al. Corneal nerve tortuosity in diabetic patients with neuropathy. *Invest Ophthalmol Vis Sci* 2004;45:418–22.
- 33 Tavakoli M, Quattrini C, Abbott C, et al. Corneal confocal microscopy: a novel noninvasive test to diagnose and stratify the severity of human diabetic neuropathy. *Diabetes Care* 2010;33:1792–7.
- 34 Edwards K, Pritchard N, Vagenas D, et al. Utility of corneal confocal microscopy for assessing mild diabetic neuropathy: baseline findings of the landmark study. Clin Exp Optom 2012;95:348–54.
- 35 Edwards K, Pritchard N, Gosschalk K, et al. Wide-field assessment of the human corneal subbasal nerve plexus in diabetic neuropathy using a novel mapping technique. Cornea 2012;31:1078–82.
- 36 Kalteniece A, Ferdousi M, Azmi S, et al. Corneal confocal microscopy detects small nerve fibre damage in patients with painful diabetic neuropathy. Sci Rep 2020;10:3371.
- 37 Issar T, Tummanapalli SS, Kwai NCG, et al. Associations between acute glucose control and peripheral nerve structure and function in type 1 diabetes. *Diabet Med* 2020;37:1553–60.
- 38 Malik RA, Veves A, Tesfaye S, et al. Small fibre neuropathy: role in the diagnosis of diabetic sensorimotor polyneuropathy. *Diabetes Metab Res Rev* 2011;27:678–84.
- 39 Paré M, Albrecht PJ, Noto CJ, et al. Differential hypertrophy and atrophy among all types of cutaneous innervation in the glabrous skin of the monkey hand during aging and naturally occurring type 2 diabetes. *J Comp Neurol* 2007;501:543–67.