Laser scanning in vivo confocal microscopy (IVCM) for evaluating human corneal sub-basal nerve plexus parameters: protocol for a systematic review

Manikkuwadura Eranda Harshan De Silva,1 Alexis Ceecee Zhang,1 Amalia Karahalios,2,3 Holly Rose Chinnery,1 Laura Elizabeth Downie1

ABSTRACT

Introduction Laser scanning in vivo confocal microscopy (IVCM) enables non-invasive, high-resolution imaging of the cornea. In recent years, there has been a vast increase in researchers using laser scanning IVCM to image and quantify corneal nerve parameters. However, a range of methodological approaches have been adopted. The primary aim of this systematic review is to critically appraise the reported method(s) of primary research studies that have used laser scanning IVCM to quantify corneal sub-basal nerve plexus (SBNP) parameters in humans, and to examine corneal nerve parameters in healthy individuals.

Methods and analysis A systematic review of primary studies that have used laser scanning IVCM to quantify SBNP parameters in humans will be conducted. Comprehensive electronic searches will be performed in Ovid MedLine, Embase and the Cochrane Library. Two reviewers will independently assess titles and abstracts, and exclude studies not meeting the inclusion criteria. For studies judged eligible or potentially eligible, full texts will be independently assessed by two reviewers to determine eligibility. A third reviewer will resolve any discrepancies in judgement. Risk of bias will be assessed using a custom tool, covering five methodological domains: participant selection, method of image capture, method of image analysis, data reporting and other sources of bias. A systematic narrative synthesis of findings will be provided. A multilevel random-effects meta-analysis will be performed for corneal nerve parameters derived from healthy participants. This review will be reported according to the Preferred Reporting Items for Systematic Reviews and Meta-Analyses statement.

Ethics and dissemination As this review considers unpublished studies or those published in a language other than English.

Strengths and limitations of this study

► This will be the first systematic review to consider the level of methodological rigour applied when using laser scanning in vivo confocal microscopy (IVCM) for clinical research.
► This systematic review will consider all primary research studies, irrespective of the study design, that have used laser scanning IVCM to quantify corneal nerve parameters in human participants.
► This systematic review protocol is reported in accordance with the Preferred Reporting Items for Systematic review and Meta-Analyses Protocols (PRISMA-P).
► We have developed a purpose-specific risk of bias tool for assessing IVCM methodological quality, which we consider will be a valuable guide for researchers using this technique, to consider potential sources of bias when developing IVCM protocols.
► The review will not include unpublished studies or those published in a language other than English.

INTRODUCTION

In vivo confocal microscopy (IVCM) is a non-invasive imaging method for visualising the structure of the living human cornea. IVCM provides high-resolution, morphological detail of the corneal architecture and can be applied to assess corneal parameters, in particular a range of metrics relating to corneal nerve integrity (eg, density and branching characteristics). Several types of IVCM instruments are commercially available, including tandem, slit and laser scanning devices.1 Laser scanning IVCM, which uses a red wavelength diode laser source that poses no ocular safety hazard,2 is currently considered the gold standard device for clinical research. This technology provides a greater depth of focus, enhanced contrast and improved resolution compared with the alternative devices.1

While early studies using laser scanning IVCM to examine corneal health were mostly


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qualitative in nature (eg, for diagnosing corneal infection), the technique is now used to determine a range of quantitative clinical measures (eg, corneal nerve density). Accurate quantification of corneal nerve parameters is important clinically for monitoring the potential effect of therapeutic interventions on corneal health, detecting corneal neuropathy and acting as a surrogate biomarker for early-stage diabetic peripheral neuropathy.\(^3\)\(^4\)\(^6\)\(^7\) Although a general method for examining the cornea and analysing corneal nerve parameters using laser scanning IVCM has been described,\(^5\) there is currently no gold standard protocol for using laser scanning IVCM for corneal nerve analysis available in the literature. As a result, a range of different approaches have been adopted.\(^3\)\(^4\)\(^6\)\(^7\)

Various factors, in particular the methods used for image capture and analysis, may introduce bias and thereby affect the accuracy of quantitative measures, when using IVCM to investigate corneal nerve parameters.\(^8\) For example, as corneal nerve density varies with eccentricity (ie, greater in the central vs peripheral cornea),\(^9\) consideration should be given to the region of cornea imaged. In addition, factors such as the microscope field of view, depth of corneal imaging, image quality and postcapture image enhancements may influence the visibility and/or clarity of nerves within the image field, thereby potentially impacting on quantitative measurements.\(^8\) The number of images analysed, per individual, also affects the confidence of quantitative estimates; it has been shown that at least eight images, with less than 20% image overlap between each image, should be analysed to obtain reliable estimates of corneal nerve density.\(^10\) To avoid potential performance biases in studies involving different participant groups and/or clinical intervention studies, the confocal microscope operator and outcome assessor should be masked to the participant’s group allocation. In addition, the method for quantifying the sub-basal nerve parameters should be fully described by researchers, with preference given to the use of a validated, fully automated processing method (eg, ACCMetrics\(^11\)), to circumvent the potential bias induced by subjective judgement.

There has not been any previous research undertaken to consider the level of methodological rigour applied when using laser scanning IVCM for clinical research. As researchers who are experienced with performing the technique, we have developed a purpose-specific risk of bias tool covering five key methodological domains that we consider important for minimising bias when using laser scanning IVCM. The five domains are participant selection, method of image capture, method of image analysis, data reporting and other sources of bias (eg, industry funding). We foresee the use of this purpose-specific risk of bias tool as a valuable guide for researchers, to consider potential sources of bias when developing their IVCM protocols. In this respect, the present paper has the capacity to contribute to significantly improving the quality of future research in the field.

The major aim of this systematic review is to critically appraise (ie, assess the risk of bias in) the reported method(s) of primary research studies that have used laser scanning IVCM to quantify corneal sub-basal nerve parameters in human participants. We will also determine key differences in methodology between studies and identify the specific methodological domains that are least well performed and/or reported (ie, are judged as having the highest risk of bias) in the literature, as a basis for informing laser scanning IVCM methods and their robust reporting, in future clinical studies. We predict that there will be considerable variation in the image-capturing methodologies used by different investigators and between the studies, which may lead to potential biases and affect the reliability of reported data. For example, studies may have used an insufficient number of corneal images as a representative measure to quantify nerve density, potentially leading to sampling bias. Finally, a meta-analysis will be conducted on studies assessing corneal nerve fibre parameters in healthy individuals. As a result, this will help to establish a more precise estimate of corneal nerve parameters for future research to use as a reference for identifying corneal nerve pathology.

Objectives

The primary objective of this systematic review is to critically appraise (ie, assess the risk of bias in) the reported method(s) of primary research studies that have used laser scanning IVCM to quantify corneal sub-basal nerve parameters in human participants. The secondary objectives are:

1. to identify the methodological domains that are least well performed and/or reported (ie, are judged as having the highest risk of bias) in the included studies, as a basis for informing laser scanning IVCM methods and their robust reporting, in future clinical studies. As shown in table 1, the five main methodological domains that will be assessed are participant selection, method of image capture, method of image analysis, data reporting and other sources of bias;
2. to determine normative values for corneal sub-basal nerve plexus parameters by pooling the estimates from available studies.

METHODS AND ANALYSIS

The proposed systematic review and meta-analysis will be undertaken using the approach recommended by the Preferred Reporting Items for Systematic Reviews and Meta Analyses (PRISMA) statement.\(^12\)

Eligibility criteria

All studies published in English will be included, from the date of database inception until 17 May 2017. In cases where multiple publications of the same data exist, the study reporting on the largest number of human participants will be included. Also, studies will be selected according to the following criteria:

- publication in a peer-reviewed journal;
- human participants;
- studies designed to measure corneal sub-basal nerve parameters using laser scanning IVCM;
- studies in which corneal nerve parameters were assessed using laser scanning IVCM; and
- studies in which at least one primary outcome parameter was a corneal nerve parameter, for which normative values have been established.

In addition, studies will be selected according to the following eligibility criteria:

- use of IVCM as the primary imaging technique;
- studies in which the methodology has been described;
- studies fulfilling the study inclusion criteria for corneal sub-basal nerve microscopy;
- studies where multiple publications of the same data exist, the study reporting on the largest number of human participants will be included;
- studies in which at least one primary outcome parameter was a corneal nerve parameter, for which normative values have been established.

Critical appraisal criteria

For each study, the following critical appraisal criteria will be used to appraise the risk of bias:

- participant selection;
- method of image capture;
- method of image analysis;
- data reporting; and
- other sources of bias.

A purpose-specific risk of bias tool covering five key methodological domains that are least well performed and/or reported (ie, are judged as having the highest risk of bias) in the literature, as a basis for informing laser scanning IVCM methods and their robust reporting, in future clinical studies. We predict that there will be considerable variation in the image-capturing methodologies used by different investigators and between the studies, which may lead to potential biases and affect the reliability of reported data. For example, studies may have used an insufficient number of corneal images as a representative measure to quantify nerve density, potentially leading to sampling bias. Finally, a meta-analysis will be conducted on studies assessing corneal nerve fibre parameters in healthy individuals. As a result, this will help to establish a more precise estimate of corneal nerve parameters for future research to use as a reference for identifying corneal nerve pathology.
Table 1  Risk of bias table for assessing the methodological quality of studies using laser scanning in vivo confocal microscopy (IVCM) for evaluating human corneal sub-basal nerve plexus parameters

<table>
<thead>
<tr>
<th>Item</th>
<th>Risk of bias</th>
</tr>
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<tbody>
<tr>
<td></td>
<td>Low</td>
</tr>
<tr>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Participant selection</strong></td>
<td></td>
</tr>
<tr>
<td>Participant inclusion and exclusion criteria (selection bias)</td>
<td>Eligibility criteria are stated, with clear specifications to define participant status (eg, HbA1c level for participants with diabetes)</td>
</tr>
<tr>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Method of image capture</strong></td>
<td></td>
</tr>
<tr>
<td>Masking of confocal operator (performance bias)</td>
<td>Clearly states that the confocal operator was masked to participant/group allocation OR that all personnel were masked throughout the study OR not applicable (if a single study population was examined)</td>
</tr>
<tr>
<td></td>
<td></td>
</tr>
<tr>
<td>Speciﬁcation of participant ﬁxation target</td>
<td>States that a consistent ﬁxation target was used for all participants</td>
</tr>
<tr>
<td></td>
<td></td>
</tr>
<tr>
<td>Location of cornea imaged</td>
<td>Quantitative description of location of cornea imaged (eg, within a 2 mm radius of the corneal apex)</td>
</tr>
<tr>
<td></td>
<td></td>
</tr>
<tr>
<td>Speciﬁcation of corneal depth</td>
<td>Quantitative description of the corneal depth imaged is included (eg, imaged at a depth range of 10–15 μm below the basal epithelium or a method is used to project nerves imaged different depths onto a single plane)</td>
</tr>
<tr>
<td></td>
<td></td>
</tr>
<tr>
<td>Illumination setting on confocal microscope</td>
<td>States that images were acquired using fixed illumination intensity for all participants</td>
</tr>
<tr>
<td></td>
<td></td>
</tr>
<tr>
<td>Selection of eye (left or right or both)</td>
<td>Clearly speciﬁes which eye was assessed, with a sound method of selection (eg, random, all right eyes, average of right and left eyes)</td>
</tr>
<tr>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Method of image analysis</strong></td>
<td></td>
</tr>
<tr>
<td>Masking of outcome assessor (performance bias)</td>
<td>Clearly states that the assessor of the corneal images was masked to the participant/group allocation, or this bias domain is not applicable (if a single study population was studied)</td>
</tr>
<tr>
<td></td>
<td></td>
</tr>
<tr>
<td>Image selection—quality (sampling bias)</td>
<td>Clearly states that image quality was assessed AND that images where the imaging depth was inconsistent or the image was blurred were removed from the analysis sample AND representative images are provided within the manuscript that conﬁrm the images meet these criteria</td>
</tr>
<tr>
<td></td>
<td></td>
</tr>
<tr>
<td>Image selection—number and sampling (sampling bias)</td>
<td>Number of images analysed is clearly stated, and at least eight images were analysed per region with &lt;20% overlap between images</td>
</tr>
<tr>
<td></td>
<td></td>
</tr>
<tr>
<td>Image selection—method of randomisation and sequence generation (sampling bias)</td>
<td>Images described to be randomly selected for analysis and method for random selection is reported (eg, computer generated list, random table, other method of generating random list)</td>
</tr>
</tbody>
</table>

Continued
<table>
<thead>
<tr>
<th>Item</th>
<th>Risk of bias</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Low</td>
</tr>
<tr>
<td></td>
<td>High</td>
</tr>
<tr>
<td></td>
<td>Unclear</td>
</tr>
<tr>
<td></td>
<td>Images were analysed consecutively (per participant) using a manual method (eg, subjective quantification) and this is considered relevant to the outcome(s)</td>
</tr>
<tr>
<td>Image processing—order of analysis</td>
<td>Order of image analysis was not reported, when a manual method was used for quantification and this is considered relevant to the outcome(s)</td>
</tr>
<tr>
<td>Postcapture image enhancements</td>
<td>No statement regarding whether postcapture image enhancements were performed</td>
</tr>
<tr>
<td></td>
<td></td>
</tr>
<tr>
<td>Definition of sub-basal nerve parameters</td>
<td>The nerve parameters being evaluated are stated but are not sufficiently well defined to allow full reproduction of the method</td>
</tr>
<tr>
<td></td>
<td></td>
</tr>
<tr>
<td>Method and repeatability of sub-basal nerve parameter quantification (intra-observer and interobserver variability)</td>
<td>Use of a validated, fully automated processing method (eg, ACCMetrics, or similar3,4)</td>
</tr>
<tr>
<td></td>
<td></td>
</tr>
<tr>
<td>Data reporting</td>
<td>Data relating to all quantified nerve parameters (as mentioned in the study methods) are reported in the results with both point measures and measures of variability</td>
</tr>
<tr>
<td>Thoroughness of reporting of nerve parameters—selective reporting of outcomes</td>
<td></td>
</tr>
<tr>
<td>Comprehensiveness of nerve parameter data (population level)—attrition bias</td>
<td>Missing data for &lt;20% of recruited participants, and if multiple study groups or a longitudinal study, then there is an equal degree of missing data in both groups at follow-up, with no obvious reason why absence of data is related to study group or time point respectively</td>
</tr>
<tr>
<td>Other</td>
<td>Other sources of bias</td>
</tr>
<tr>
<td>CNBD, corneal nerve branch density; CNFD, corneal nerve fibre density; CNFL, corneal nerve fibre length; HbA1c, glycated haemoglobin; RCT, randomised controlled trial.</td>
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</tbody>
</table>
Study designs
We will include all primary, empirical research studies that have used a laser scanning confocal microscope to perform corneal confocal microscopy on at least one human participant, where corneal sub-basal nerve plexus parameters were quantified. We will include studies from across the spectrum of clinical research questions defined by the National Health and Medical Research Council (eg, intervention, diagnostic test accuracy, aetiology, prognosis and screening intervention) and study designs (eg, randomised controlled trial (RCT), pseudo-RCT, non-RCT, cohort, case-control, cross-sectional, interrupted time series, case series, case study), to enable the comparison of methodological quality across study types.

Studies reporting only on aspects of corneal architecture other than sub-basal nerve parameters (eg, epithelial thickness, endothelial cell count/morphology, corneal haze, and so on) will be excluded. We will exclude review papers (including systematic reviews), conference abstracts and studies reporting methods for analysing laser scanning IVCM images, where human participants were not recruited. We will also exclude studies that have used alternative types of confocal microscopes for image capture (eg, tandem scanning and slit scanning), as the type of confocal microscope affects the quantitation of corneal sub-basal nerve parameters.

Participants
We will include all studies that report corneal sub-basal nerve plexus findings for at least one human participant. There will be no restriction on participant health status for the systematic review (although restrictions will apply for the meta-analysis, which will only include data from healthy adults); thus, included studies may involve healthy individuals, as well as those with ocular and/or systemic conditions.

Information sources
A comprehensive search to identify all relevant studies will be undertaken in the following electronic databases: Ovid MEDLINE(R) (Epub Ahead of Print, In-Process & Other Non-Indexed Citations, Ovid MEDLINE(R) Daily and Ovid MEDLINE(R) 1946 to May 2017), Ovid EMBASE (Embase Classic+Embase, 1947 to May 2017) and the Cochrane Library.

To ensure literature saturation, we will scan the reference lists of included studies, or relevant reviews, identified by our search. We will also search the authors’ personal bibliographic reference files to ensure that all relevant studies are captured. We will also circulate a list of the included articles to our review team.

Search strategies
The search strategies are provided in online supplementary material.

Study records

Data management
The systematic search will be carried out by the review team, using the previously defined search strategies, and guided by Items 9 and 10 of the Preferred Reporting Items for Systematic review and Meta-Analysis Protocols (PRISMA-P) statement. After performing the search strategies separately in each electronic database, the researchers will import the results from each search into an EndNote library. As the same article may be located in more than one database, duplicate entries will be identified and removed.

Study selection
We will use Covidence systematic review software, an online program that facilitates collaboration between reviewers for systematic reviews, for the study screening process. Two reviewers (MEHDS and ACZ) will independently assess the titles and abstracts of all unique studies, identified from the electronic search strategies, and exclude those that do not meet the inclusion criteria. For studies judged to be eligible or potentially eligible for inclusion, the full-text articles will be sourced and independently assessed by the two reviewers, to clarify their eligibility for inclusion. Any discrepancies in classification that arise during this process will be resolved by consensus between the two reviewers and a third reviewer. For studies that progress to the full-text screening stage, we will record the reason that studies were excluded.

If there are cases where it is unclear whether the inclusion criterion are met, we will attempt to contact the study corresponding author for clarification; if no response is received within four weeks of the request, or the requested information is not provided, the information within the full-text article will be used to decide on the eligibility of the study.

A diagram will be created to report the flow of studies through the systematic review.

Data collection
Relevant data, from eligible studies, will be independently extracted by two reviewers in Covidence, using a standardised data extraction form. Extracted data will be summarised in tables. If any data extraction discrepancies arise, these will be resolved by discussion and consensus among the review team.

Data Items
Extracted data from each included study will include:

i. article details: year of publication, journal of publication;
ii. study details: type of research question (ie, intervention, diagnostic-test accuracy, aetiology, prognosis, screening intervention), setting, location, study design (eg, RCT, pseudo-RCT, non-RCT, cohort, case-control, cross-sectional, interrupted time series, case series, case study), number of participants, health status of the participant population(s) (eg, healthy, (PRISMA-P) statement. After performing the search strategies separately in each electronic database, the researchers will import the results from each search into an EndNote library. As the same article may be located in more than one database, duplicate entries will be identified and removed.

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diabetes, and so on), participant characteristics (age, gender), population eligibility criteria (inclusion and exclusion criteria);

iii. methodological details: unit of analysis (one eye (right or left), both eyes or average of both eyes, as applicable), corneal sub-basal nerve parameters assessed (see ‘Outcomes’ section for further details), IVCM image capture field of view (ie, 300 or 400 µm), IVCM mode of image capture (eg, volume, sequence or section scan), whether a representative IVCM sub-basal nerve plexus image is provided in the paper (dichotomous);

iv. other details: source of funding statement (dichotomous: present or absent), actual source of funding (eg, industry or Government funding), conflict of interest statement (dichotomous: present or absent), conflict of interest type (eg, employee of company conducting study);

v. quantitative measures: data (ie, mean (SD) or median (IQR)) for the following four key central, corneal sub-basal nerve parameters: CNFL, CNFD, CNBD and CTBD, as defined in the ‘Outcomes’ section. Where data are provided for both eyes, we will also extract the correlation coefficient. If longitudinal data are reported, we will use baseline data in our analyses. As all of the sub-basal nerve plexus parameters are continuous outcomes, we will extract data on the means and SD for each parameter, or similar measures of central tendency and variability.

Outcomes
The primary outcome will be the methodological quality of included research studies that have used laser scanning IVCM to quantify corneal sub-basal nerve parameters in human participants.

The secondary outcomes are as follows:

i. Identification of the methodological domains that are least well performed and/or reported (ie, are judged as having the highest risk of bias) in the included studies, as a basis for informing laser scanning IVCM methods and their robust reporting, in future clinical studies.

ii. Meta-analysis of mean normative values (ie, from healthy individuals) for corneal sub-basal nerve plexus parameters, quantified from the central cornea (as defined by the study authors), and using these definitions for the analysis.

► Corneal nerve fibre length (CNFL) defined as the total length of all nerve fibres in the image capture frame (mm/mm²). If an alternative definition is used, such as limiting the quantification of fibres to those of a certain minimum length, these data will be excluded.

► Corneal nerve fibre density (CNFD) defined as the total number of main fibres divided by the area of the image frame (fibres/mm²).

► Corneal nerve branch density (CNBD) defined as the total number of main nerve branches, being branches that stem from a nerve fibre, divided by the area of the image frame (branches on main fibre/mm²).

► Corneal nerve total branch density (CTBD) defined as the total number of branches within the area of the image frame (total branches/mm²).

Assessment of risk of bias in included studies
To facilitate the assessment of methodological quality in each of the included studies, as per the objective of this review, we developed a purpose-specific, 18-item risk of bias tool (table 1) to assess internal validity, encompassing five main domains:

► participant selection (including selection bias)

► method of image capture (including performance bias and sampling bias)

► method of image analysis (including performance bias)

► data reporting (selective reporting of outcomes and attrition bias)

► other sources of bias (funding source and conflicts of interest).

The risk of bias tool was developed by the review team (MEHDS, ACZ, HRC, LED), who possess expertise in using IVCM for corneal nerve analyses, for this review and was framed using the Cochrane Assessing Risk of Bias in included studies (Chapter 8 in the Cochrane Handbook of Systematic Reviews of Interventions).

Two reviewers will judge the risk of each type of bias (18 items in total) in each included study as either (1) low risk, (2) unclear risk (due to either lack of information or uncertainty about the potential for bias) or (3) high risk. Review authors will resolve any disagreements in bias assessment by consensus with a third reviewer. Reviewers will not be masked to the journal of publication or the study author name when undertaking the risk of bias assessment.

Wherever possible, we will justify each risk of bias assessment with direct quotations from the study. If there are cases where further information is considered necessary to determine the risk of bias in a particular domain, we will attempt to contact the study corresponding author for this information. If no response is received within four weeks of the request, or the requested information is not provided, the information within the full-text article will be used to inform the risk of bias assessment.

Data synthesis
For outcomes related to methodological quality, a systematic narrative synthesis will be provided, with relevant information summarised in text and tables.

If there are at least three relevant studies, we will undertake meta-analyses of the quantitative data for the specified corneal sub-basal nerve plexus parameters (ie, mean (SD)). Data from male and female participants will be pooled, as studies have shown that gender has no significant effect on corneal sub-basal nerve plexus parameters.20 21
We will convert non-parametric data to means (SD), using an established approach. We will fit a multi-level random-effects model to pool the estimates. The multilevel model will take into account the correlation between estimates from the same study that are presented separately for each sex and/or estimates presented separately for the left and right eyes. Next, we will fit a meta-regression model to assess how much of the between-study variation is explained by the following characteristics: (1) participant age (as this factor is potentially important); (2) study design (eg, RCT, cohort (including pseudo-RCT and non-RCT) and other (including cross-sectional, case series/study)).

Statistical analyses will be carried out using the metafor package in R.

**Meta-bias(es)**

As there are no limitations on the potential study designs eligible for inclusion in this review, we expect that we will not be able to compare the outcomes reported in published reports with study protocols, unless the included study is a RCT, to assess for selective outcome reporting or selective analysis reporting. Furthermore, as our meta-analysis is being undertaken to determine values for normative parameters (ie, corneal sub-basal nerve plexus parameters), rather than the effect of an intervention, we do not expect meta-biases (such as publication bias, delayed publication, and so on) to be a significant factor for this analysis.

**Sensitivity analyses**

Provided there are a sufficient number of studies included in the review, sensitivity analyses will be performed for the CNFL outcome, to assess for the effect of excluding studies that (1) were appraised as having a high risk of bias in the domains of image selection—number and sampling, or method for quantifying sub-basal nerve parameters; (2) included contact lens wearers (ie, contact lens wear was not listed as an eligibility exclusion criterion); (3) were lower order levels of evidence (eg, case reports, case series, interrupted time series); and (4) are from the same corresponding senior author, in the event that at least 50% of the included papers are from the research laboratory of the same corresponding author.

**Confidence in cumulative evidence**

If appropriate, we will present a ‘Summary of Findings’ table for the quantitative outcomes. In this case, the quality and strength of the body of evidence will be assessed using an approach based on the Grading of Recommendations Assessment, Development and Evaluation.

**CONCLUSIONS**

In recent years, an increasing number of research studies have adopted non-invasive, laser scanning IVCM to quantify corneal sub-basal nerve plexus parameters. However, there has not been any research to formally consider the quality of the methods used in these investigations. This systematic review will provide insight into the quality of the methods reported in clinical studies using laser scanning IVCM to quantify corneal nerve parameters. The review will also identify specific methodological domains that are least well performed and/or reported (ie, are judged as having the highest risk of bias) in the literature, as a basis for informing laser scanning IVCM methods and their robust reporting, in future clinical studies. Furthermore, by researchers considering the elements of the purpose-specific risk of bias tool as a guide when developing their IVCM protocols, this review has the capacity to significantly improve the quality of future research in the field. By undertaking a meta-analysis, we will also determine mean normative values (ie, from healthy individuals) for central corneal sub-basal nerve plexus parameters. These data will be of significant value for future studies, as reference normative values, building on a previous pooled analysis of data derived from multiple laser scanning IVCM testing centres.

**REFERENCES**