Research Paper

Quantitative analysis of morphological and functional features in Meibography for Meibomian Gland Dysfunction: Diagnosis and Grading

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ABSTRACT

Background: To explore the performance of quantitative morphological and functional analysis in meibography images by an automatic meibomian glands (MGs) analyser in diagnosis and grading Meibomian Gland Dysfunction (MGD).

Methods: A cross-sectional study collected 256 subjects with symptoms related to dry eye and 56 healthy volunteers who underwent complete ocular surface examination was conducted between January 1, 2019, and December 31, 2020. The 256 symptomatic subjects were classified into MGD group (n = 195) and symptomatic non-MGD group (n = 61). An automatic MGs analyser was used to obtained multi-parametric measurements in meibography images including the MGs area ratio (GA), MGs diameter deformation index (DI), MGs tortuosity index (TI), and MGs signal index (SI). Adjusted odds ratios (ORs) of the multi-parametric measurements of MGs for MGD, and the area under the receiver operating characteristic (AUC-ROC) curves of multi-parametric measurements for MGD diagnosing and grading were conducted.

Findings: When consider age, sex, ocular surface condition together, the estimated ORs for DI was 1.62 (95% CI, 1.29-2.56), low-level SI was 24.34 (95% CI, 2.73-217.3), TI was 0.76(95% CI, 0.54-0.90), and GA was 0.86 (95% CI, 0.74-0.92) for MGD. The combination of DI-TI-GA-SI showed an AUC = 0.82 (<P < 0.001) for discriminating MGD from symptomatic subjects. The DI had a higher AUC in identifying early-stage MGD (grade 1-2), while TI and GA had higher AUCs in moderate and advanced stages (grade 3-5). Merging DI-TI-GA showed the highest AUCs in distinguishing MGD severities.

Interpretation: The MGs area ratio, diameter deformation, tortuosity and signal intensity could be considered promising biomarkers for MGD diagnosis and objective grading.

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1. Introduction

Meibomian Gland Dysfunction (MGD) is one of the most common ocular surface diseases extremely prevalent worldwide, especially in Asian populations (from 46.2% to 69.3%) [1,2]. As the most common trigger of dry eye diseases (DED) [2], MGD has also been reported to be associated with lots of other ocular diseases, i.e., allergic conjunctivitis, ocular rosacea, Sjögren syndrome [3-5], and systemic factors such as androgen deficiency, dyslipidemia, and aging [6-8], that degrades patients’ ocular comfort, visual function, and quality of life [9,10,11]. A high accuracy objective diagnosis strategy for MGD could provide significant benefits. However, existent diagnostic tests based on the observation of abnormal anatomy and physiology of the MGs opening and lid secretions [12] mostly rely on the experience of expert clinicians, making quantification difficult and more likely to be affected by interobserver variability. The variation of the results may also affect by involvement of gland selection errors, different pressure applied manually by the operators [13]. It’s also difficult for some patients to cooperate with the squeeze examination with obvious inflammatory state of eye lids and too sensitive to invasive tests [10]. Moreover, DED and MGD have high similarities in symptoms...
Research in context

Evidence before this study

We searched PubMed for articles published between January 1, 2000 and May 1, 2021 using the terms: ("meibomian gland dysfunction" OR "meibomian" OR "meibography") AND ("diagnosis" OR "evaluation" OR "severity" OR "grading"). Most previous studies on evaluation of meibomian glands (MGs) were relatively subjective and more likely to be affected by interobserver variability, involving invasive operations, and including gland selection errors. It still lacks effective and detailed quantitative analysis tools, and this difficulty poses a challenge for the development of diagnosis, grading, and treatment of meibomian gland dysfunction (MGD).

Added value of this study

Our study developed an automated multiparametric quantitative analyst of MGs in meibography images, and found that increased cross-sectionally uneven gland dilation, decreased gland area ratio, low axial distortion and low signal intensity of MGs showed high risks for the presence of MGD. A combination of all these features has good differentiation power in identifying MGD patients from symptomatic subjects (AUC = 0.82), and the merge of morphological features showed excellent accuracy in distinguishing MGD severities. These variables are quickly assessed by the non-invasive meibography through the self-developed automated algorithm.

Implications of all the available evidence

Since MGD is one of the most common ocular surface diseases degrades patients’ quality of life, it often overlaps with dry eye in pathological progress and requires specific meibomian gland-oriented therapies, our automated quantitative analysis of morphological and functional features of MGs have good differentialization power in identifying MGD patients from symptomatic subjects and excellent accuracy in distinguishing MGD severities, which will optimize the clinical management of the disease.

and often overlap in pathological progress [14,15], which leads to the negligence of distinguishing between the two in clinical screening [8]. While MGD requires specific meibomian gland-oriented therapies [16], a non-invasive and objective meibomian gland assessment method to quickly distinguish MGD from people with dry eye symptoms will optimize diagnosis and treatment algorithms.

Noncontact meibography techniques provide ophthalmologists a direct way to visualize the meibomian glands with an inverted eyelid in vivo [17]. Based on meibography images, meibograde was developed as a semiquantitative tool to approximately assess gland loss in vivo [17,18], which has been demonstrated to be associated with dry eye [19,20], aging [17], contact lens wearing [3], ocular allergies [21], and severe inflammatory ocular surface diseases [4,5]. Nevertheless, patients with such detectable meibomian gland loss are more likely to be in an advanced stage with seriously impaired meibomian gland function [22]. In fact, detailed morphological features such as gland shortening, distortion, hook, and tortuosity can also be observed in meibography images, some of which are evidence of MGs with worsened expressibility and meibograde [23]. Scholars believe that these gland irregularities may represent an early stage of MGD [9,12,22,24]. Due to the lack of automated quantitative assessment methods, clinical studies on whether and how these diverse detailed morphological features are related to the pathophysiological states of MGD are limited. In addition to morphological changes, observations from meibography images have revealed that changes in the secretion and quality of meibum inside meibomian glands could alter the signal intensity of the glands [25]. Exploring the information the signal intensity of MGs in meibography images brings [26] may provide a non-invasive functional indicator to determine whether MGs are in a high- or low-delivery state.

Recently, our team has developed an objective multi-parametric meibomian glands analyser to quantitatively analyse the meibomian glands using infrared meibography images [27]. This analyser can automatically segment and quantitatively analyse the morphological details (gland area ratio, tortuosity, and deformation) and potential functional information from the signal index values of all exposed meibomian glands. The automatic segmentation function of the analyser has been proven to have high similarity compared with manual segmentation by professional ophthalmologists.

The present study aimed to explore the performance of quantitative morphological and functional parameters in meibography images by our self-developed automatic MGs analyser in distinguish MGD patients from people with DED symptoms and MGD grading.

2. Methods

2.1. Data Collection

This cross-sectional, observational study included 256 symptomatic subjects and 56 healthy volunteers. Between January 1, 2019, and December 31, 2020, two-hundred-fifty-six subjects with ocular symptoms related to dry eye disease (DED) were collected from the patient pool of the Zhongshan ophthalmic center (ZOC) Dry Eye Clinic in Guangzhou, China. Fifty-six healthy volunteers without any systemic diseases or pre-existing ocular conditions or symptoms were recruited at the same period from the general population through the ZOC as a normal control group. All participants were required to include information of the Ocular Surface Disease Index (OSDI) symptom questionnaire, ocular surface staining, non-invasive tear-film break-up time (NIBUT), Schirmer I test, meibography, and meibum expressibility and quality. All procedures were conducted following the Declaration of Helsinki (1983) and were approved by the Institutional Review Board of Zhongshan Ophthalmic Centre, Sun Yat-sen University, China (protocol number: 2019KYPI110). Informed consent was obtained from each of the participants prior to data collection. This study followed STROBE guidelines strictly.

2.2. MGD group and Symptomatic non-MGD group Ascertainment

The diagnosis of MGD was based on an OSDI ≥13 and the presence of an altered quality of expressed secretions and/or decreased or absent expression as follows: (1) score >1 for either meibum quality or expressibility or (2) score = 1 for both meibum expressibility and meibum quality [12]. The symptomatic subjects who did not meet the diagnosis criteria mentioned above were defined as symptomatic non-MGD group. One-hundred-ninety-six subjects were diagnosed among 256 symptomatic subjects. The remaining 61 subjects without MGs disorders were defined as the symptomatic non-MGD group, including 33 [54.1%] DED cases. Diagnosis of DED in symptomatic non-MGD group was based on NIBUT, Schirmer I test and ocular surface staining; (1) NIBUT < 10s or (2) Schirmer I test (without anesthesia) ≤10mm in 5min, and a corneal fluorescein staining score of 4 or more based on the corneal staining scale [12]. The MGD patients were further evaluated with regard to the MGD severity level and scored with 1–5 according to the 2011 MGD workshop [12].

2.3. Clinical assessments

The symptom questionnaire OSDI was performed measuring the occurrence frequency of ocular symptoms, environmental triggers...
and vision related quality of life [28]. Corneal fluorescein staining score is conducted as a standard objective measure to visualize the extent of ocular surface damage [28,29]. Schirmer I test (without anaesthesia) measured the tear secretion to confirm aqueous deficiency [28]. NIBUT and infrared photography of the upper meibomian glands measured by Keratograph 5M (Oculus, Wetzlar, Germany). NIBUT is the interval of time that elapses between a complete blink and the appearance of the first break in the tear film measured by non-invasive methods to evaluate tear film stability [18,28]. Meibum expressibility was scored from the lower central five glands as follows: 0, all five glands; 1, three to four glands; 2, one to two glands; and 3, zero glands [30]. The quality of the expressed meibum was scored from 0-3: 0 = clear fluid; 1 = cloudy fluid; 2 = cloudy, particulate fluid; and 3 = opaque, toothpaste-like meibum [31]. The meibomian gland yield secretion score (MGYSS) was collected from five glands in the central, temporal, and nasal eyelids, for a total of fifteen glands in the lower eyelids. For each of these glands, the secretion was scored as follows: 0: no secretion; 1: inspissated/toothpaste consistency; 2: cloudy liquid secretion; and 3: clear liquid secretion [32]. The scores were then summed across the 15 glands for a single MGYSS, with a range from 0 to 45 [13]. The above assessments were collected from each participant.

One eye was randomly selected for analysis from each non-MGD subject and binocular MGD patient with the same severity. When the severity of MGD was different in both eyes for MGD subjects, the more serious eye was selected.

2.4. Multi-parametric automated meibomian gland analyser

To obtain the quantitative morphological and functional parameters using the acquired infrared meibography images, all images were processed and analysed with our self-developed automated MGs analyser. For details of the principle and process of this customized software, refer to our previous study [27]. In brief, the meibography images were converted to greyscale, exported and saved as bitmaps, after which they underwent an automated segmentation of the everted tarsal conjunctiva area as the region of interest (ROI), followed by the segmentation and identification of all glands within the ROI (Fig. 1). Then the segmented results were quantitatively analysed with pre-defined morphological and functional parameter calculations, exporting multi-parametric results including the gland area ratio (GA), gland diameter deformation index (DI), gland tortuosity index (TI), and gland signal index (SI). In detail, the gland area ratio is defined as the percentage of detected pixels in the glands over the segmented ROI area; the gland diameter DI addresses the diameter variations of a gland, such as uneven dilation and discontinuous atrophy; the gland TI is used to quantify the degree of curving and hairpin-loop-like winding changes of the glands; and the gland SI is the average image greyscale value of the segmented intact glands divided by the average image greyscale value of the non-gland area.

2.5. Statistical analysis

Comparisons of measurements among the three groups were assessed by analysis of variance (ANOVA) for normally distributed continuous variables and chi-squared test for categorical variables. The SI was transformed into categorical variables corresponded to clinical observation of the low, moderate, and high signal intensity of glands as low-level SI (SI < 4.5), medium-level SI (4.5 ≤ SI ≤ 6.5) and high-level SI (SI < 6.5). The cut-off values were set between two subgroups (10 cases as a unit) with large differences in prevalence of MGD in ascending range of SI. Bivariate correlations were performed using Spearman correlations coefficients between multiple measurements of MGs and MGYSS in all subjects with completed ocular

| Characteristics | Mean GA | Mean DI (SD) | Mean TI (SD) | Mean SI (range) |
|-----------------|---------|--------------|--------------|----------------|
| A1              | 44.23   | 7.98 (4.45)  | 11.92 (4.01) | 5.84 (4.5~6.5) |
| A2              | 39.69   | 15.86 (9.82) | 16.85 (17.64)| 6.98 (> 6.5)   |
| A3              | 41.82   | 10.99 (4.48) | 12.91 (7.59) | 6.09 (4.5~6.5) |
| A4              | 21.09   | 6.55 (4.10)  | 7.27 (3.78)  | 4.47 (< 4.5)   |

Fig. 1. (A) the appearance of the lid margins and meibomian orifices, (B) the Meibography images with automatically segmented boundaries of the everted tarsal conjunctiva area as the region of interest (ROI), and (C) the segmented meibomian glands of representative subjects. The sequence from 1 to 4 is based on the current stage of MGs pathological progression from healthy to severe. Examples of the targeted individual features of tortuosity (yellow arrow), dilation & distortion (red arrow), and atrophy or drop out (white arrow) are marked with arrows. The acquired multi-parametric analysis results of the four representative Meibography images are shown in the table below the figure. DI: diameter deformation index; TI: tortuosity index; SI: signal index; SD: standard deviation; GA: gland area ratio, which is the ratio of all meibomian gland area to the total analysis area.
surface examination mentioned above (n = 312). Binary logistic regression was conducted to identify the independent correlations of multiple measurements of MGs associated with MGD diagnosis. Variables were considered for adjustment in the binary logistic regression model if they displayed a P value of less than 0.05 in the tests of between-group comparison. Sex and age were also considered for adjustment in the model.

By dividing all the symptomatic subjects (n = 256) into a training set and a test set with a ratio of 80%: 20%, the detection probability of the multiple measurements of MGs to discriminate MGD from symptomatic subjects was evaluated by performing regression model with the training set and calculating the area under the curve of a receiver operating characteristic curve (AUC-ROC) in the test set. Kendall's tau-b correlation was performed in symptomatic subjects between multiple measurements of MGs and severities to screen potential grading indicators. The prediction value of each potential grading variable for each grade (grade 1–5) was evaluated with AUC-ROCs. The grading performance of the combined features were also evaluated by testing AUC-ROCs from training regression models by subdividing each grade into training and test sets (80%: 20%). The statistical analyses above were performed in SPSS 20.0 (SPSS, Chicago, IL). P < 0.05 were considered statistically significant. P-values were adjusted by Bonferroni correction in multiple comparisons.

2.6. Role of funding sources

All sources of funding had no role in study design, data collection, data analysis, data interpretation or writing of this manuscript, or the decision to submit it for publication. The corresponding authors had full access to all the data in the study and had final responsibility for the decision to submit for publication.

3. Results

3.1. The clinical measurements, morphological and functional analysis of MGs among groups

Typical cases with the appearance of lid margins, segmented meibomian glands, and the acquired multi-parametric analysis results of the glands are shown in Fig. 1. Data for the clinical measurements, the morphological and functional measurements of MGs of all eligible participants (n = 312) from the MGD group (n = 195), symptomatic non-MGD group (n = 61) and normal group (n = 56) and the P values from pairwise comparisons of the values among the 3 groups are presented in tables 1 and 2, respectively. Age and sex showed no significant difference within each pair of subject groups. Adverse changes in the OSDI, NI-BUT, meibum expressibility, MGYSS, corneal fluorescein staining (CFS) score, and Schirmer’s I test were increased significantly in the MGD group compared to the normal group and the symptomatic non-MGD group. The GA (41.07%) and TI (13.44) were found in the MGD group lower than in the symptomatic non-MGD group (GA 47.74%, TI 14.90) and normal group (GA 47.25%, TI 14.66) (P < 0.05, respectively). The DI was 12.84 in the MGD group, significantly higher than in the symptomatic non-MGD group (DI 9.62) and normal group (DI 10.83) (P < 0.05, respectively). The normal group showed the highest frequency of a medium-level SI (76.79%). The MGD groups showed the highest frequency of both a low-level SI (25.13%) and a high-level-SI (26.15%) among three groups. (tables 1 and 2).

| Parameter | Symptomatic groups | Normal Control | P of trend value |
|-----------|--------------------|----------------|-----------------|
| Sex, no. (%) | Male | 77 (60.51) | 118 (39.49) | 0.05 |
| Sex, no. (%) | Female | 118 (39.49) | 77 (60.51) | 0.01 |
| Severity, no. (%) | Grade 0 | 0 (0) | 61 (100) | 0.001 |
| Severity, no. (%) | Grade 1 | 47 (24.10) | 47 (24.10) | 0.001 |
| Severity, no. (%) | Grade 2 | 64 (32.82) | 64 (32.82) | 0.001 |
| Severity, no. (%) | Grade 3 | 47 (24.10) | 47 (24.10) | 0.001 |
| Severity, no. (%) | Grade 4 | 28 (14.36) | 28 (14.36) | 0.001 |
| Severity, no. (%) | Grade 5 | 9 (4.72) | 9 (4.72) | 0.001 |
| Expressibleness, no. (%) | 0 | 9 (4.26) | 51 (83.61) | 0.001 |
| Expressibleness, no. (%) | 1 | 89 (45.64) | 10 (16.39) | 0.001 |
| Expressibleness, no. (%) | 2 | 63 (32.31) | 0 | 0.001 |
| Expressibleness, no. (%) | 3 | 34 (17.44) | 0 | 0.001 |
| | OSDI, mean (SD) | 25.87 (10.56) | 17.05 (6.55) | 3.89 (2.77) | 0.001 |
| | NI-BUT, mean (SD), s | 8.80 (4.81) | 12.95 (5.14) | 16.25 (4.92) | 0.001 |
| | MGYSS, mean (SD) | 24.81 (7.12) | 40.54 (6.30) | 41.59 (3.93) | 0.001 |
| | CFS score, mean (SD) | 0.87 (0.88) | 0.05 (0.22) | 0.001 |
| | STI, mean (SD), mm | 12.25 (6.24) | 16.48 (6.05) | 16.39 (6.65) | 0.001 |
| Morphological parameters | DI, mean (SD) | 12.84 (5.77) | 9.62 (2.76) | 10.83 (2.87) | 0.001 |
| Morphological parameters | TI, mean (SD) | 13.44 (2.77) | 14.90 (3.22) | 14.66 (2.46) | 0.001 |
| Functional parameter:SI range, no (%) | SI:≤4 | 49 (25.13) | 6 (9.84) | 3 (5.36) | 0.001 |
| Functional parameter:SI range, no (%) | 4.5:≤SI:6.5 | 95 (48.72) | 42 (68.85) | 43 (76.79) | 0.001 |

Abbreviations: MG: meibomian gland; MGD: meibomian gland dysfunction; SD: standard deviation; OSDI: ocular surface disease index; NI-BUT: non-invasive tear film break-up; MGYSS: meibomian gland yield secretion score; CFS: corneal fluorescein staining score; STI: Schirmer test I; DI: diameter deformation index; TI: tortuosity index; GA: gland area ratio, which is the ratio of all meibomian gland area to the total analysis area; SI: signal index of the glands.
3.2. Evaluation of the diagnosis performance of the multiple measurements of MGs in MGD

Table 3 demonstrates results from the binary logistic regression models concerning how the DI, TI, GA, and SI of MGs discriminated MGD from symptomatic non-MGD subjects (n = 256), controlling for age, sex, MGYSS, NIBUT, FSC score and Schirmer test I. With the results reported as estimated odds ratios (95% CI), the increase of the DI (1.62 [1.29-2.56]) and the number of subjects with an SI under 4.5 (24.34 [2.73-217.30]) demonstrated significant positive associations with the presence of MGD. An increased TI (0.76 [0.54-0.96]) and GA (0.86 [0.74-0.92]) showed negative associations with the presence of MGD. The accuracy of identifying whether a symptomatic individual had MGD based on the DI, TI, GA, and SI was evaluated to calculate the AUC-ROC and showed that AUC = 0.82 (P < 0.001) in the test set (Fig. 2).

3.3. Evaluation of the clinical associations and grading performances of multiple measurements of MGs

Significant associations were found between the MG morphological parameters (DI, TI, and GA) and MGYSS (all P < 0.05, Fig. 3). The Spearman coefficients for the correlations between morphological parameters (GA, TI, and DI) and MGYSS were 0.40, 0.22 and -0.19. In symptomatic subjects, the DI, TI, and GA were significantly associated with MGD severities. Kendall’s tau-b correlation coefficients were 0.15 (P = 0.001), -0.25 (P < 0.001), and -0.40 (P < 0.001), respectively, excluding the SI (P = 0.25). The ROC curve analysis for the single and combined morphological parameter with respect to each level of MGD severity can also be seen in table 4. The performance of single variables showed that DI had a highest AUC-ROC in identifying early-stage MGD (grade 1, 0.76, P < 0.001; grade 2, 0.74, P < 0.001). GA had highest AUC-ROCs in moderate and advanced stages (grade 3, 0.83, P < 0.001; grade 4, 0.92, P < 0.001; grade 5, 1.00, P < 0.001), followed by TI (grade 3, 0.70, P = 0.001; grade 4, 0.77, P < 0.001; grade 5, 0.92, P < 0.001). The combined DI-TI-GA showed the highest AUC-ROCs compared to single variables in each grade in the test sets (grade 1, 0.79, P = 0.03; grade 2, 0.80, P = 0.01; grade 3, 0.87, P = 0.004; grade 4, 0.94, P = 0.003; grade 5, 1.00, P = 0.03) (table 4). The specific trends of the 3 morphological parameters across different severities can be visualized in figure 4.

Table 2

| Parameter               | MGD vs. Symptomatic non-MGD | MGD vs. Normal Control | Symptomatic non-MGD vs. Normal Control |
|-------------------------|------------------------------|------------------------|---------------------------------------|
| OSDI                    | -0.001                       | -0.001                 | -0.001                                |
| NIBUT (s)               | -0.001                       | -0.001                 | 0.001                                 |
| Meibum expressibility (0-3)| -0.001                      | -0.001                 | -0.99                                 |
| MGYSS                   | -0.001                       | -0.001                 | -0.99                                 |
| CFS score               | -0.001                       | -0.001                 | -0.99                                 |
| STI (mm)                | 0.04                         | -0.001                 | 0.49                                  |
| Morphological parameters|                             |                        |                                       |
| DI                      | -0.001                       | 0.02                   | 0.54                                  |
| TI                      | 0.001                        | 0.01                   | >0.99                                 |
| GA (%)                  | -0.001                       | -0.001                 | >0.99                                 |
| Functional parameter: SI|                             | -0.001                 | 0.55                                  |

Abbreviations: MGs: meibomian glands; MGD: meibomian gland dysfunction; SD: standard deviation; OSDI: ocular surface disease index; NIBUT: non-invasive tear film break-up; MGYSS: meibomian gland yield secretion score; CFS: corneal fluorescent staining score; ST I: Schirmer test I; DI: diameter deformation index; TI: tortuosity index; GA: gland area ratio; SI: signal index of the glands.

Table 3

| Variable | OR* | 95% CI | Lower | Upper |
|----------|-----|--------|-------|-------|
| DI       | 1.62| 1.29   | 2.56  |
| TI       | 0.76| 0.54   | 0.90  |
| GA (%)   | 0.86| 0.74   | 0.92  |
| STI      | 0.99| 0.96   | 0.99  |

*: OR adjusted for sex, age, MGYSS, NIBUT, FSC score and Schirmer test I.

Abbreviations: DI: diameter deformation index; TI: tortuosity index; GA: gland area ratio; SI: signal index of the glands.

4. Discussion

In the present study, we reported a comprehensive analysis of MGs based on an automated algorithm [27] using meibography images by quantifying the gland area ratio as GA, the gland irregular shapes as DI and axial distortions as TI in morphological analysis. Quantification of the optical density of the glands relative to the background as the SI produced a potential functional index of MGs. Our results demonstrated that the multi-measurements of MGs (DI, TI, GA and SI) from the regression model adjustment for sex, age and other pathologic changes on the ocular surface that may affect the presentation of disease [17,33] were associated significantly with the presence of MGD. The combination of DI-TI-GA-SI showed an excellent diagnosis performance in identification of MGD patients from people with suspected DED, which will benefit to fast screen for MGD in symptomatic population. The assessments of the morphological multi-parameters of MGs (DI, TI, GA and SI) have good consistency with the evaluation of meibum quality and expressibility (MGYSS), reflected typical objective changes in different MGD pathological stages. Merging the morphological parameters (DI-TI-GA) could
complement each other to increase the accuracy of to distinguish MGD severities. The automatic quantitative multi-measurements of detailed morphological and functional information from meibography images provide a non-invasive, fast, and easy processing method for MGD diagnosis and grading.

The GA was calculated as the whole MGs area ratio by eliminating the gaps between glands, which can reflect additional information of gland thickening, thinning and abnormal gaps compared to MGs drop out quantified as meibograde [22,34]. The thinned glands and large gaps had been observed and determined to represent incomplete gland atrophy in the DREAM study [35]. While MGs drop out was reported to represent the final stage of MGD [22], our study demonstrated that a decreased GA was independently associated with the presence of MGD from level 2 to level 5, revealed a thinned glands and large gaps were also involved in the progression of the disease. Conversely, for the lowest MGD severity (level 1), the GA was slightly increased, which may be a result of MG dilation and thickening. Increased MGs thickness was also observed to be inversely correlated with meibum expression and considered to be a compensatory response to gland loss and increased meibum demand [36]. Dilation of secretory acini could be part of the same process, as a result of the expansion of accumulated and inspissated debris within the acini observed in previous studies [23,37,38].

A typical dilation pattern was proven by another morphological parameter—the diameter deformation index (DI), which addresses the degree of diameter variations to present the uneven thickening or thinning of MGs. A high DI was observed in the MGs of early pathological stages (level 1-3) and declined at levels 4 and 5. ROC-AUCs analysis also showed that the DI had better predictive accuracy in the earliest stage, indicating that the increased DI may be a sensitive marker of early MGD before visible glands loss. These findings are consistent with previous objective observations that gland dilation is an early visual pathogenic finding and associated with progressive loss of MGs [22]. The pattern of uneven thickening or thinning can result from progressive pressure due to partial obstruction of the ductal system and an inflammatory process of the secretory acinus [22,39].

It has been confirmed that curving and hairpin-loop-like tortuosity as another morphology pattern occur both in progressive distortion of the ductal system [39,40] and asymptomatic people [41]. Whether the tortuosity of MGs could translate into a pathological change [42] or only a congenital pattern [41] is still controversial. In this study, the TI showed no significant change compared to symptomatic non-MGD and normal group in the early stage of MGD (level 1-2) but showed similarities with the GA that decreased significantly in moderate and severe MGD with increased severities prediction accuracy. Therefore, axial distortion is more likely to be a kind of congenital morphology pattern and not sensitive enough to be used as a single indicator for MGD diagnosis in the early stage, and a decline TI with a reduction of the GA can represent overall gland atrophy, which is different from gland dropout or shortened glands, as a complement sign of severe loss of acinar cells in late pathologic stages.

Existing functional examinations of the MGs such as the meibum quality and expressibility, which have limitations of only assessing several glands and are impossible to conduct when the gland is completely obstructed. Since changes in the consistency and colour of the meibum have been observed under pathological conditions [39], we defined the gland signal index (SI) to provide a non-contact biomarker that may offer functional information by quantifying the optical density of the meibum within the ductal system. We demonstrated that subjects with an extreme SI, who were defined as those with low levels (< 4.5) and high levels (> 6.5), were found to

![Fig. 3. Scatterplots for the correlations between average morphological parameters of MGs and MGYSS. The GA (%) and DI were significantly correlated with MGYSS (P < 0.001).](image)

The Spearman coefficients were rGA: 0.40, rTI: 0.22 and rDI: -0.19. MGYSS: meibomian gland yield secretion score; DI: diameter deformation index; TI: tortuosity index; GA: gland area ratio, the ratio of all meibomian gland area to total analysis area.

![Fig. 4. Boxplots for the comparison of morphological parameters across different severity levels. The DI, TI, and GA (%) across levels 0–5 of MGD severity are shown. Significant increases in the DI are shown at levels 1–3, while significant decreasing trends are shown in the TI at levels 3–5. The GA (%) increased slightly at level 1 and then gradually decreased at levels 2–5.](image)

Table 4 Receiver operating characteristic curve analysis for the combination and each morphological parameter with respect to each grade of MGD severity

| Severe Grade | DI AUC | P | TI AUC | P | GA (%) AUC | P | DI-TI-GA AUC | P |
|--------------|--------|---|--------|---|------------|---|-------------|---|
| Grade 1      | 0.75*  | <0.001| 0.57  | 0.19| 0.52       | 0.73| 0.79*        | 0.03|
| Grade 2      | 0.74*  | <0.001| 0.60  | 0.04| 0.67*      | 0.001| 0.80*        | 0.01|
| Grade 3      | 0.67*  | 0.003| 0.70  | 0.001| 0.83*      | <0.001| 0.87*        | 0.004|
| Grade 4      | 0.68   | 0.08 | 0.77  | <0.001| 0.92*      | <0.001| 0.94*        | 0.003|
| Grade 5      | 0.68   | 0.08 | 0.90  | <0.001| 1.00*      | <0.001| 1.00*        | 0.03|

*: P-value < 0.05.

Abbreviations: MGD: meibomian gland dysfunction; DI: diameter deformation index; TI: tortuosity index; GA: gland area ratio.
constitute a higher proportion of the MGD group than the symptom-
atic non-MGD group and normal group. Moreover, low-level SI MGs
harbour a 24.3-fold increased risk for the presence of MGD
compared to patients with a medium-level SI (4.5–6.5) of MGs (OR:
24.34, 95% CI 2.73 – 217.30). Although the large confidence interval
indicate sort of uncertainty, potentially due to the large dispersion of
SI itself and the conversion of continuous SI values into categorical
variables, the large impact of the low-level SI in MGD deserves clin-
cal attention. The low-level SI represents dim glands with low optical
signals, which is in agreement with the study by Daniel et al [35],
who observed a large number of pale glands were associated with
eyelids expressed none or thick paste like consistency sebum. We
concluded that the low-level SI MGs are sick glands with low-deliv-
ery state progressing towards atrophy. The high-level SI could be
related to the excessive accumulation of meibum in MGs caused by
obstruction or typical lipid compositions [43]. Thus, SI may provide
an overall assessment of glands function to assist MGD classification.
In this study, only the upper eyelids were included since the
upper lids were demonstrated to have more visible MG features and
a stronger correlation with clinical signs than lower lids [35]. More-
over, meibography images of the lower eyelid are more likely to
show unevenly focused and uncompleted exposed meibomian
glands, which are changing in automatic segmentation and analy-
sis. To further clarify the clinical potential and pathologic information
provided by morphological and functional measurements of MGs, a
prospective cohort study to track longitudinal changes in meibomian
glands from disease or intervention will also be necessary in the
future. In addition to DED and MGD, the morphology and function of
MGs are also observed to related to many ocular and systemic abnor-
malities, i.e., allergic conjunctivitis, ocular rosacea, dyslipidemia, dia-
betes, and aging., etc. [3-5,6-8,17], the automatic and quantitative
methods established in this study could have extended application
prospects in future studies regarding the roles of MGs in other related
ocular and systematic conditions.
In conclusion, with the automated multiparametric quantitative
analysis of MGs in meibography images, our study found that
morphological features including an increased DI (cross-sectionally
uneven gland dilation) and a slightly increased GA (thickened and
dilated glands) are common in the early stage of MGD, while a
decreased GA (greatly influenced by incomplete atrophy) together
with a decreased TI (decreased axial distortion) are signs of severe
MGD, and revealed that glands with a low-level SI showed a high risk
for the presence of MGD. A combination of DI-TI-GA-SI has good dif-
ferentiation power in identifying MGD patients from symptomatic
subjects, and the merge of morphological parameters DI-TI-GA
showed excellent accuracy in distinguishing MGD severities.

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Contributors
JY and PX devised the project, the main conceptual ideas and
proof outline. YQ-D contributed to the design and implementation of
the research and the analysis of the results and to the writing of the
manuscript. QW supervised the findings of this work and contributed
to the writing of the manuscript. ZZ-L & PX developed and verified
the automatic Meibomian glands analyser, and aided in interpreting
the results and worked on the manuscript. SQ-L, JZ, BW-W, and LL-P
performed the measurements and performed the analysis of data. All
authors discussed the results and commented on the manuscript. JY,
PX and YQ-D accessed and were responsible for the raw data associ-
ated with the study.

Data sharing statement
All data will be available upon reasonable request to the corre-
sponding author, and it will be shared according to the standards of
ethical policies regulating data sharing of human subjects.

Declaration of Competing Interest
We declare no conflicts of interests.

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