Histopathology and selective biomarker expression in human meibomian glands

Lixing W Reneker
Rebecca T Irlmeier
Ying-Bo Shui
Ying Liu
Andrew J W Huang

Follow this and additional works at: https://digitalcommons.wustl.edu/open_access_pubs
Histopathology and selective biomarker expression in human meibomian glands

Lixing W Reneker 1, Rebecca T Irlmeier, Ying-Bo Shui, Ying Liu, Andrew J W Huang 2

ABSTRACT

Background/aims Meibomian gland dysfunction (MGD) is the most common form of evaporative dry eye disease, but its pathogenesis is poorly understood. This study examined the histopathological features of meibomian gland (MG) tissue from cadaver donors to identify potential pathogenic processes that underlie MGD in humans.

Methods Histological analyses were performed on the MGs in the tarsal plates dissected from four cadaver donors, two young and two old adults, including a 36-year-old female (36F) and three males aged 30, 63 and 64 years (30M, 63M and 64M).

Results The MGs of 36F displayed normal anatomy and structure, whereas the MGs of 30M showed severe ductal obstruction with mild distortion. The obstruction was caused by increased cytokeratin levels in association with hyperproliferation, but not hyperkeratinisation. In two older males, moderate to severe MG atrophy was noted. Cell proliferation was significantly reduced in the MG acini of the two older donors as measured by Ki67 labelling index (6.0%±3.4% and 7.9%±2.8% in 63M and 64M, respectively) when compared with that of the two younger donors (23.2%±5.5% and 16.9%±4.8% in 30M and 36F, respectively) (p<0.001). The expression patterns of meibocyte differentiation biomarkers were similar in the older and younger donors.

Conclusion Our histopathological study, based on a small sample size, suggests potentially distinct pathogenic mechanisms in MGD. In the young male adult, hyperproliferation and aberrant differentiation of the central ductal epithelia may lead to the obstruction by overproduced cytokeratins. In contrast, in older adults, decreased cell proliferation in acinar basal epithelia could be a contributing factor leading to MG glandular atrophy.

INTRODUCTION

Meibomian glands (MGs), located in the upper and lower tarsal plates of the eyelids, are holocrine glands composed of secretory acini surrounding a central duct.1 Based on their anatomical locations, acinar epithelia cells are classified as basal (germinative or proliferative), differentiating, mature and superficial degenerated cells. Each acinus at its basal circumference comprises a single layer of proliferative epithelial cells, which are essential for acinar regeneration. When the basal cells start to differentiate, they move inward and mature into lipid-producing meibocytes. After reaching maturity, the meibocytes disintegrate as holocrine cells and release lipids (meibum) into the short collecting ductules that connect with the central secretory duct lined by the squamous epithelial cells. Meibum produced from the mature and decomposed meibocytes travels through the ductule to the central duct and is discharged through the central duct and onto the ocular surface via the MG orifice. The secreted meibum constitutes the outermost layer of the tear film, overlying the inner mucous layer and providing tear film stability.2 Normal meibum from healthy MGs forms a barrier against tear film evaporation and protects the ocular surface from microbial and environmental insults such as dust and pollen.

Meibomian gland dysfunction (or disease; MGD) is a term broadly used to encompass various functional abnormalities of MGs.3–5 Conventionally, MGD can be classified as MG ductal obstruction, MG hyposecretion and MG hypersecretion. MG ductal obstruction and hyposecretion are more common than MG hypersecretion, which is frequently associated with eyelid inflammation and occurs in rosacea, chalazion and allergy.6–8 Animal models of MGD have shown that ductal hyperkeratinisation can result in ductal obstruction, meibum stasis, cystic dilatation and, eventually, acinar atrophy and MG dropouts.9 Keratinised materials, however, are not normally present in the ducts of human MGs.10 Compromised meibum quality is another contributing factor to ductal obstruction.11 Studies in both animals and humans suggest that acinar epithelial abnormalities, such as diminished renewal of acinar basal cells and/or impaired meibocyte differentiation, contribute to the pathogenesis of age-related MGD.12–14 Androgen deficiency and medications, such as isotretinoin, are also thought to play a role in MGD.15–17

The high prevalence of MGD and its diverse pathology implicate that it is a multifactorial disease,18–19 but the histopathological changes and underlying molecular mechanisms of MGD have not been extensively elucidated, which makes it clinically challenging to diagnose and manage MGD effectively. To gain a better understanding of the pathogenesis of MGD, we performed histopathological examinations and biomarker expression analyses on MG tissues of four cadaver donors, including two young adults and two older adults. Our current study implies that fundamentally different pathogenic mechanisms underlie MGD. Despite the limited sample size in this study, our findings unveiled previously poorly described pathogenic mechanisms of MGD and shed light on potential avenues for the development of
mechanism-targeted diagnostic and therapeutic strategies for MGD.

**MATERIALS AND METHODS**

**Acquisition of donor tarsal tissues**

Fresh tarsal tissues from four cadaver donors were obtained from Mid-America Transplant (St. Louis, Missouri, USA). The donors were free of known systemic illness and included two young adults and two older adults: a 30-year-old male (30M), a 36-year-old female (36F), a 63-year-old male (63M) and a 64-year-old male (64M). The medical and ocular histories of the donors were deidentified prior to removal of the tarsal plates from the inner upper eyelids of the fresh cadavers. The removal of donor corneas and tarsal plates was performed by eye bank staff under the standard protocol for procurement of ocular tissues. The use of human tissues in research conformed to the ethical guidelines of the Institutional Review Board of Mid-America Transplant.

**Histology, immunofluorescence, immunohistochemistry and quantification of cell proliferation**

Human tarsal tissues were fixed overnight in 4% paraformaldehyde in phosphate-buffered saline (PBS) and then processed for embedding in paraffin. Tissue sections (5 µm) were treated with xylene; rehydrated in a decreased ethanol series; and subjected to heat-induced antigen retrieval for immunohistochemistry.20 For antigen retrieval of immunostaining, tissue sections were boiled in 10 mM citrate buffer (pH 9.0) for 1 hour to heat denature proteins and then washed with PBST (PBS plus 0.1% Tween-20) for 1 hour. Tissue sections were then incubated overnight at 4°C with primary antibodies diluted in the same buffer (Table 1). Slides were washed with PBS, incubated at room temperature for 1 hour with either fluorophore-conjugated or biotinylated secondary antibodies and then washed with PBS. For immunofluorescence, cell nuclei were stained with 4',6-diamidino-2-phenylindole (DAPI) and cell membranes were stained with triton X-100. Slides were mounted with Vectorshield (Vector Laboratories, Burlingame, California, USA) and colour was developed by using 3,3'-diaminobenzidine as a substrate.

**Acquisition of donor tarsal tissues**

Figure 1  Human MG in tarsal plate and histology in MG tissue from four cadaver donors. (A–D) Tarsal plates dissected from cadavers’ eyelids, designated as 36F (A), 30M (B), 64M (C) and 63M (D), were photographed directly before trimming off some of the orbicularis oculi muscle. Note: The thickness of the dissected tissue and setting of the photographic illumination gave rise to different appearance in tissue transparency. The two older donors (C, D) showed moderate to severe MG atrophy with glandular shortening (arrow in C) or dropout (arrow in D). (E–H) Histology (H&E staining). The circumference of MG acinus (a) in the 36F is made of a layer of basal epithelium that surrounds a cluster of lipid (meibum)-producing meibocytes (E). In 30M (F), thickened ductal epithelia (arrows in F) and hypercellularity were noted in the Cd. MGs of the two older donors (G and H) exhibited fibrotic areas with MG remnants (G’, arrowheads) and acinar atrophy (H, arrows). Scale bar denotes 100 µm. 36F, 36-year-old female; 30M, 30-year-old male; 63M, 64-year-old male; 64M, 64-year-old male; a, acinus; Cd, central duct; d, ductule; MG, meibomian gland.

**Table 1** Sources and dilution ratios of primary antibodies

| Protein | Company       | Product catalogue  | Dilution ratio |
|---------|---------------|--------------------|----------------|
| FASN    | Santa Cruz    | sc-48357           | 1:500          |
| Ki67    | Cell Signaling| 9129               | 1:500          |
| Krt1    | Santa Cruz    | sc-65999           | 1:1000         |
| Krt6a   | BioLegend     | 905701             | 1:5000         |
| Krt10   | Covance       | PRB-159P           | 1:1000         |
| Krt14   | Covance       | PRB-155P           | 1:1000         |
| Krt16   | Novusbio      | NBP2-45528         | 1:1000         |
| Krt17   | Novusbio      | NBP2-16089         | 1:1000         |
| p63     | Santa Cruz    | SC-25268           | 1:200          |
| PCNA    | Abcam Inc.    | ab29               | 1:1000         |
| PPARγ   | Cell Signaling| 2435               | 1:200          |

FASN, fatty acid synthase; PCNA, proliferating cell nuclear antigen; PPARγ, peroxisome proliferator activated receptor gamma.

**RESULTS**

**Histology of MGs in human cadaver donors**

The MGs in the tarsal plates dissected from cadaver eyelids were first examined for gross morphology (figure 1A–D). The MGs of 36F were uniformly distributed in the tarsal plate and acini were evenly sized and appeared normal (figure 1A). Mild ductal distortion was observed in the MGs of 30M (figure 1B). In the two older donors (63M and 64M), MG atrophy, including glandular shortening and dropouts (arrows in figure 1C,D), was readily visualised in tarsal tissues.

Histology revealed notable variations in the acinar and ductal structure of MGs among the donors (figure 1E–H). In 36F, the lipid-producing acini were connected to the central duct by short branching ductules (figure 1E). By comparison, in 30M the MG central ducts were thickened from epithelial hypercellularity and filled with cellular debris and nuclear fragments (figure 1F, arrows). Compared with the two younger donors (30M and 36F), the MG tissues of the two older donors (64M and 63M) displayed altered features, with obliterated (figure 1G) and shrunken acini (figure 1H) scattered among areas containing normal-appearing acini (figure 1G).

**Ductal obstruction without increased keratinisation biomarkers**

Krt10 and its type II pair Krt1 are considered to be keratinisation markers because they are expressed in terminally differentiated...
keratinocytes.21 To investigate whether hyperkeratinisation of the ductal epithelium may account for the ductal obstruction in 30M, the expression patterns of Krt10 and Krt1 in MGs of the four donors were examined. We first verified the respective specificity of anti-Krt10 and anti-Krt1 antibodies by confirming their positive staining patterns at the mucocutaneous junction (MCJ) of cadaver eyelids (online supplementary figure S1). Expression of both Krt10 and Krt1 was interrupted at the MCJ where the epidermal epithelium transitions into the conjunctival epithelium (indicated by open arrows in online supplementary figure S1). Krt10 and Krt1 expression patterns in MGs of 36F, 63M and 64M differed from those of 30M. In 36F, both Krt10 and Krt1 were present in the suprabasal layer of the collecting ductules and the central duct (arrows in figure 2A,C). Similar patterns were seen in the MGs of 63M and 64M (data not shown). Interestingly, in the central duct of 30M, a Krt10/Krt1-positive suprabasal layer (red arrow in figure 2B inset) was sandwiched between the basal layer (black arrow) and a yet uncharacterised Krt10/Krt1-negative superficial layer (figure 2D, open arrows). The Krt10/Krt1-positive middle layer in the central duct of 30M was not thicker than the corresponding layer in the central duct of the 36F. Furthermore, the debris blocking the central duct of 30M did not stain positive for Krt10 (figure 2B, arrowhead). Therefore, the Krt10/Krt1 expression pattern in 30M suggests that hyperkeratinisation is unlikely to be the direct cause of ductal obstructive MGD in this case.

The abnormal Krt10/Krt1 expression pattern in the central duct of 30M prompted us to investigate the expression of Krt14, a marker for basal cells in most epithelial tissues, including the skin.21 In MGs of both 30M and 36F, Krt14 was highly expressed in the acini and in the basal epithelium of the central duct (figure 2E,F). However, in 30M, Krt14 expression was found not only in the basal layer of the central duct but also in multiple suprabasal layers (figure 2F inset). The abnormal and expanded expression of Krt14 in the central ductal epithelia was confirmed on multiple MGs from 30M and also by using cross sections of MGs from this donor (data not shown). These results suggest abnormal proliferation and differentiation of the central ductal epithelia may cause ductal obstruction in this donor.

Ductal obstruction from overproduction of cytokeratins linked to hyperproliferation

Keratin 16 (Krt16) is typically expressed in stratified squamous epithelia and certain glandular tissues.20–22 Overexpression of Krt16 and its type II pair Krt6, along with Krt17 overexpression, is associated with keratinocyte hyperproliferation and considered to be hallmarks of psoriasis, a chronic inflammatory skin disease.23 24 To explore the role of these cytokeratins in MGD, we examined the expression of keratins 16, 6a and 17 in the MGs of 30M, 36F and 64M (figure 3).

In MGs of 36F and 64M, Krt16 was mostly expressed in the ductules adjacent to the central duct, moderately expressed in the acinar epithelia and certain glandular tissues.20–22 Overexpression of Krt16 and its type II pair Krt6, along with Krt17 overexpression, is associated with keratinocyte hyperproliferation and considered to be hallmarks of psoriasis, a chronic inflammatory skin disease.23 24 To explore the role of these cytokeratins in MGD, we examined the expression of keratins 16, 6a and 17 in the MGs of 30M, 36F and 64M (figure 3).

In 30M, Krt16 was distributed in acinar and ductal epithelia more ubiquitously than Krt16/Krt6a (figure 3G–I). The Krt17 level appeared to be increased in the thickened central ducts of 30M, as compared with Krt17 expression in the central ducts of 36F and 63M (compare figure 3H with figure 3G,I). Taken together, the findings suggest that overproduction of Krt16/Krt6a and Krt17 in the central ductal epithelia of the 30M is likely the underlying or a contributing cause of ductal obstruction in this donor.

Like Krt16 in psoriatic dermatitis, overexpression of Krt17 has been found in this disease.25 In our study of MGs, Krt17 was distributed in acinar and ductal epithelia more ubiquitously than Krt16/Krt6a (figure 3G–I). The Krt17 level appeared to be increased in the thickened central ducts of 30M, as compared with Krt17 expression in the central ducts of 36F and 63M (compare figure 3H with figure 3G,I). Taken together, the findings suggest that overproduction of Krt16/Krt6a and Krt17 in the central ductal epithelia of the 30M is likely the underlying or a contributing cause of ductal obstruction in this younger male donor, who probably had clinically undetected obstructive MGD.

Hyperproliferation and aberrant differentiation of central ductal epithelia

Because overexpression of Krt16 and Krt17 is associated with keratinocyte hyperproliferation, we examined the cell proliferation markers Ki67 and proliferating cell nuclear antigen (PCNA) in central ductal epithelia and compared the levels in 30M and 36F (figure 4A–D). Both Ki67- and PCNA-positive cells were markedly increased in the basal epithelia of the central duct of 30M when compared with the levels in the central duct of 36F. Furthermore, the Ki67 labelling index in the MG ductal basal epithelia of 30M

Figure 2 Expression of Krt10, Krt1 and Krt14 in MGs of the two young donors, one with normal-appearing MGs (36F) and the other with evidence of ductal obstruction (30M). (A,B) Krt10 was expressed in the suprabasal layers of ductules and central duct in 36F (A, red arrows). In the central duct of 30M, the Krt10-positive suprabasal layers (inset in B, red arrow) were ‘sandwiched’ between two Krt10-negative layers, presumably the basal layer (black arrow) and the superficial layer (open arrow). Ductal blockage (arrowhead in B) was caused by Krt10-negative proteins mixed with cell nuclear fragments. (C,D) Krt1 expression level in the MGs of 36F and 30M was high in the ductules and reduced in the suprabasal layer of the central duct (as indicated by arrows in C). Similar to the Krt10 pattern seen in MGs of 30M (B), the superficial layer of the central duct in this donor was negative for Krt1 (open arrows in D). (E,F) Krt14 was highly expressed in the MG acinar and ductule epithelia in both 36F (E) and 30M (F). Krt14 expression was limited to the basal epithelium in the central duct in 36F (arrows in E). The thickness of the Krt14-positive layer was notably increased in the central duct of 30M (F, black arrow in the inset). Scale bar denotes 100 μm. 36F, 36-year-old female; 30M, 30-year-old male; a, acinus; Cd, central duct; d, ductule; MG, meibomian gland.
Figure 3  Expression of Krt16, Krt6a and Krt17 in MGs of an older donor (64M) and of the two young donors, one with normal-appearing MGs (36F) and the other with evidence of ductal obstruction (30M). (A–C) Krt16 protein was mostly distributed in MG ductules and decreased in the MG central duct of 36F (A) and 64M (C) as compared with Krt16 expression in 30M (B). The central ductal cells in 30M showed Krt16 overexpression (B, montage and inset on the top right) and an orifice ‘plug’ (B, red arrow in bottom left inset). (D–F) Krt6a expression was similar to Krt16 expression in the MG ductules of 36F (D) and 64M (F). Krt6a overproduction was seen in the central duct of 30M (E). Krt17 was ubiquitously expressed in the MGs of 30M, 36F and 64M, with a higher level in the epithelial cells of ductules. Krt17 overexpression was noted in the central duct of 30M, but not in the central duct of 36F and 64M (H). Scale bar denotes 100 µm. 36F, 36-year-old female; 30M, 30-year-old male; 64M, 64-year old male; a, acinus; Cd, central duct; d, ductule; MG, meibomian gland.

Figure 4  Expression of cell proliferation markers in MGs of the donor with ductal obstruction (30M), the donor with normal-appearing MGs (36F) and one of the older donors (63M). (A, B) Ki67 expression. Ki67-positive cells were rarely identified in the basal epithelia of the central ducts in 36F (A). In contrast, many Ki67-positive cells were noted in the basal epithelia of 30M (B, black arrows). (C, D) PCNA expression was found in the nuclei of the acinar basal epithelium of 36F (C, arrow), whereas in 30M PCNA-positive cells were abundant in the epithelia of the central duct (D, black arrow). In 30M, cell nuclear fragments were retained in the superficial layer of the MG central duct (B and D, red arrows). (E) Ki67 labelling index in MG ductal basal epithelia. Cell proliferation rate in ductal basal epithelia was higher in 30M than in 36F and 63M (p<0.001) (figure 4E). The histopathology also showed that nuclear fragments were retained in the thickened suprabasal layer of the central duct of 30M (red arrows in figure 4B,D), suggesting the abnormal differentiation of the ductal epithelia. These results together suggest that the potential cause of obstructive MGD in 30M is epithelial hyperproliferation with aberrant differentiation, associated with overproduction of cytokeratins, leading to the accumulation of proteins and nuclear debris in the central duct.

Age-related decline of acinar cell renewal and gland atrophy

Transcription factor p63 (TP63), a member of p53 family, is expressed by proliferative stem cells or progenitor cells in several stratified epithelia. We found p63 expression in the basal epithelia of MG acini, the ductule and the central duct in MG tissue of each donor (figure 5A,B and data not shown for 30M and 64M), but p63 expression was reduced in some of the MG basal acini of 63M (figure 5B, red ‘a’). When cell proliferation marker Ki67 was colocalised with p63, Ki67-positive cells were also notably decreased in the acini of 63M, as compared with the number of Ki67-positive cells in the acini of 36F (compare figure 5B with figure 5A). The Ki67 labelling index was significantly lower in the older group (6.0%±3.4% and 7.9%±2.8% for 63M and 64M, respectively) than that in the younger group (23.2%±5.5% and 16.9%±4.8% for 30M and 36F, respectively) (p<0.001) (figure 5C), suggesting that this decrease of cell proliferation may compromise acinar cell renewal and haemostasis in older individuals, leading to age-related MG atrophy (figure 1D).

The main function of MGs is to produce and secrete lipids (meibum) that coat the outer tear film and prevent its evaporation. The expression patterns of the meibocyte differentiation markers, peroxisome proliferator activated receptor gamma and fatty acid synthase, were compared between the 36F and older (64M) donor, representing a young with normal-appearing
MGs and an older donor with age-related atrophy. The staining patterns for these differentiation biomarkers were similar in the MGs of 36F and 64M (figure 5D–G), suggesting that, in this older donor’s acini where cell proliferation was reduced (figure 5C), the expression of meibocyte differentiation markers appeared to be normal.

DISCUSSION

MGD is the major cause of evaporative dry eye disease and has diverse aetiologies. 3 10 19 29 Ductal obstruction and acinar or glandular atrophy are common features of MGD. To understand the pathogenesis of MGD, it is of paramount importance to investigate the histopathological changes in human MGs. This study describes the histological findings and the results of immunostaining for selective biomarkers in the MGs of cadaver tarsal plate tissues from two donors in the fourth decade of life (young age group) and two donors in the seventh decade of life (older age group). Despite of limited sample size, the data shed light on the notion that multiple pathogenic factors can contribute to the development of MGD.

Hyperkeratinisation of ductal epithelia has been shown in animal models of MGD to alter the structure and function of MGs by plugging the MG central ducts, causing ductal dilation and disruption of meibum excretion. 9 10 Ductal plugs comprising non-keratinised epithelial cells have also been observed in animal models of MGD. 12 30 In this current study of human MGs, the tissue debris blocking the central ducts of 30M did not display the typical characteristics of hyperkeratinisation (figures 2 and 3). In the MGs of this donor, the keratinisation biomarkers Krt10 and Krt1 were present in the middle layer, above the Krt14-expressing basal layer, but underneath the superficial layer of the central ductal epithelia. Based on this abnormal expression pattern of the keratinisation biomarkers (Krt10 and Krt1), an alternative mechanism other than hyperkeratinisation seems likely to be the cause of ductal obstruction in this young male adult.

We propose that hyperproliferation and aberrant differentiation of the central ductal epithelial cells led to MG ductal obstruction and plugging in 30M. This conclusion is supported by the presence of overproduction of Krt6a/Krt16/Krt17 (figure 3) and increased cell proliferation markers (figure 4). Besides ductal obstruction, excessive cytokeratins are also disruptive to the quality of meibum as well as the stability of the lipid layer in the tear film. Meibum undergoes a maturation process as it is secreted into the ductules, with protein removal and/or lipid accumulation prior to excretion onto the ocular surface and incorporation into the tear film. 31 It has been shown that the addition of purified cytokeratins to meibum lipid can affect the fluidity of the meibum. 32 Such an increase in the protein–lipid ratio correlates with increased meibum viscosity. 33 Thus, it is conceivable that overproduction and retention of Krt6a/Krt16/Krt17 in the central duct, as found in the MGs of 30M, could have a profound effect on meibum quality and fluidity and lead to MG obstruction and the eventual development of MGD.

MGD is often associated with certain skin diseases, such as acne rosacea, atopic dermatitis, seborrhoeic dermatitis and psoriasis. 8 It was reported that patients with psoriasis vulgaris have a high prevalence of obstructive MGD, characterised by ductal plugging and impaired meibum secretion. 20 In psoriatic skin, the Krt6/Krt16 pair and Krt17, along with cell proliferation maker K67, are considered to be biomarkers for keratinocyte hyperproliferation. 21 The pertinent medical record of the 30M prior to corneal and tarsal plate removal did not reveal any indication of psoriatic skin lesions. Krt6 and Krt16 are known as stress-activated keratins. 35 Our interesting findings in this donor’s MGs implicate a potential utility of Krt6a/Krt16 staining of meibum excreta as diagnostic biomarkers to ascertain the diagnosis of obstructive MGD.

It is well documented that the process of ageing is associated with a decrease in the number of MGs along with an increase in MG dropouts. 10 14 15 Postulated factors include age-mediated hormonal changes, stem cell attenuation and growth factor deficiency. Other studies have demonstrated reduced K67 nuclear staining in old murine and human MGs, 14 15 which is consistent with our current finding of a reduced number of K67-positive cells in the acinar basal epithelia in MGs of 64M and 63M (figure 5C). Furthermore, we found that p63, a transcription factor for epithelial morphogenesis and stemness, was also downregulated in some acini in the MGs of 63M (figure 5B). Taken together, these observations suggest that a deficiency in acinar cell proliferation and/or MG progenitor/stem cell renewal may be a major contributing factor for the development of age-related MGD. In searching for the endogenous regulators responsible for MG cell renewal and homeostasis, we recently found that fibroblast growth factor receptor 2 (FGFR2) is highly expressed in both murine and human MGs. Using an inducible conditional knockout mouse model, we have demonstrated that FGFR2 signalling plays an essential role in MG acinar...
proliferation and renewal.\textsuperscript{20} Nevertheless, the significance of FGFR2 and its ligands in human MG homeostasis awaits further investigations.

In summary, while our study is limited by the small sample size due to the poor availability of donor eyelids, the histopathological and biomarker analyses of MG tissues from a young donor with normal-appearing MGs, a young donor with ducal obstruction and two donors in their early 60s shed insights into the diverse underlying pathogenic mechanisms of MGD. The findings of our study suggest that ductal epithelial hyperproliferation and abnormal differentiation, without notable hyperkeratinisation, can lead to MG obstruction, as in the case of 30M, whereas a decline in acinar cell proliferation and renewal is more likely the underlying cause of age-related MG atrophy, as in the two older donors. The findings also suggest that overexpression of hyperproliferative keratins in meibum may serve as potential biomarkers to diagnose specific type of obstructive MGD.

Contributors LWR and AJWH were involved in study conceptualisation, supervision, data analysis, manuscript drafting and editing. All authors were involved in data collection and interpretation.

Funding This study was supported by the National Eye Institute of the National Institutes of Health under award number EY24221.

Competing interests None declared.

Patient consent for publication Not required.

Provenance and peer review Not commissioned; externally peer reviewed.

Data availability statement Data are available in a public, open access repository. All data relevant to the study are included in the article or uploaded as supplementary information.

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 ID Lixing W Reneker http://orcid.org/0000-0002-5622-1326

REFERENCES

1. Knop E, Knop N, Millar T, et al. The International workshop on meibomian gland dysfunction: report of the Subcommittee on anatomy, physiology, and pathophysiology of the meibomian gland. Invest Ophthalmol Vis Sci 2011;52:1938–78.

2. Butovich IA. Meibomian glands, meibum, and meibogenesis. Exp Eye Res 2017;163:2–16.

3. Baudouin C, Messmer EM, Aragona P, et al. Revisiting the vicious circle of dry eye disease: a focus on the pathophysiology of meibomian gland dysfunction. Br J Ophthalmol 2016;100:300–6.

4. Nelson JD, Shimazaki J, Benitez-del-Castillo JM, et al. The International workshop on meibomian gland dysfunction: report of the definition and classification Subcommittee. Invest Ophthalmol Vis Sci 2011;52:1930–7.

5. Nien CJ, Paugh JR, Massel S, et al. Age-related changes in the meibomian gland. Exp Eye Res 2009;89:1021–7.

6. Mizoguchi S, Iwashi H, Arita R, et al. Ocular surface inflammation impairs structure and function of meibomian gland. Exp Eye Res 2016;173:78–84.

7. Reyes NJ, Yu C, Mathew R, et al. Neutrophils cause obstruction of eyelid sebaceous glands in inflammatory eye disease in mice. Sci Transl Med 2018;10:aas9164.

8. Randon M, Liang H, Abbas R, et al. [A new classification for meibomian gland diseases with in vivo confocal microscopy]. J Fr Ophthalmol 2016;39:239–47.

9. Lambert R, Smith RE. Hyperkeratinisation in a rabbit model of meibomian gland dysfunction. Am J Ophthalmol 1988;105:703–5.

10. Obata H. Anatomy and histopathology of human meibomian gland. Cornea 2002;21(7 Suppl):S70–4.

11. Nakayama N, Kawashima M, Kaido M, et al. Analysis of Meibum before and after intrastral meibomian gland probing in eyes with obstructive meibomian gland dysfunction. Cornea 2015;34:1206–8.

12. Partift GI, Xie Y, Geyfman M, et al. Absence of ductal hyperkeratinization in mouse age-related meibomian gland dysfunction (ARMGD). Aging 2013;5:825–34.

13. Jester JV, Partift GI, Brown DJ. Meibomian gland dysfunction: hyperkeratinization or atrophy? BMC Ophthalmol 2015;15(Suppl 1):156.

14. Nien CJ et al. Effects of age and dysfunction on human meibomian glands. Arch Ophthalmal 2011;129:462–9.

15. Azcarate PM, Venincasa VO, Feuer W, et al. Androgen deficiency and dry eye syndrome in the aging male. Invest Ophthalmol Vis Sci 2014;55:5046–53.

16. Ding J, Kam WR, Dieckow J, et al. The influence of 13-cis retinoic acid on human meibomian gland epithelial cells. Invest Ophthalmol Vis Sci 2013;54:4341–50.

17. Moy A, McNamara NA, Lin MC. Effects of isotretinoin on meibomian glands. Optom Vis Sci 2015;92:925–30.

18. Schaumberg DA, Nichols JJ, Papas EB, et al. The international workshop on meibomian gland dysfunction: report of the Sub委员会 on the epidemiology of, and associated risk factors for, MGD. Invest Ophthalmol Vis Sci 2011;52:1994–2005.

19. Tomlinson A, Bron AJ, Koth DR, et al. The International workshop on meibomian gland dysfunction: report of the diagnosis Subcommittee. Invest Ophthalmol Vis Sci 2011;52:2006–49.

20. Reneker LW, Wang L, Inrmeier RT, et al. Fibroblast Growth Factor Receptor 2 (FGFR2) is required for meibomian gland homeostasis in the adult mouse. Invest Ophthalmol Vis Sci. 2017;58:2638–46.

21. Tovsola DM, Boor P, Alain C, et al. Keratins in health and disease. Curr Opin Cell Biol 2013;25:73–81.

22. Lessard JC, Pina-Paz S, Rotty JD, et al. Keratin 16 regulates innate immunity in response to epidermal barrier breach. Proc Natl Acad Sci USA 2013;110:19357–42.

23. Ogawa E, Sato Y, Minagawa A, et al. Pathogenesis of psoriasis and development of treatment. J Dermatol 2018;45:264–72.

24. Yang L, Fan X, Cui T, et al. Nrf2 promotes keratinocyte proliferation in psoriasis through up-regulation of keratin 6, keratin 16, and keratin 17. J Invest Dermatol 2017;137:2168–76.

25. Yang L, Jin L, Ke Y, et al. E3 ligase TRIM21 ubiquitylates and stabilizes keratin 17 to induce STAT3 activation in psoriasis. J Invest Dermatol 2018;138:2568–77.

26. Pasa R, Yang A, McKeon F, et al. Association of p63 with proliferative potential in normal and neoplastic human keratinocytes. J Invest Dermatol 1999;113:1099–105.

27. Yoh K, Pyres R. Pathway regulation of p63, a director of epithelial cell fate. Front Endocrinol 2015;6:51.

28. Green-Church KB, Butovich I, Willcox M, et al. The International workshop on meibomian gland dysfunction: report of the Subcommittee on tear film lipids and Lipid–Protein interactions in health and disease. Invest Ophthalmol Vis Sci. 2011;52:1979–93.

29. Bron AJ, de Paiva CS, Chauhan SK, et al. TFOS DEWS II pathophysiology report. Ocul Surf 2017;15:438–510.

30. Ong B-L, Hodson SA, Wigham T, et al. Evidence for keratin proteins in normal and abnormal human meibomian fluids. Curr Eye Res 1991;10:1113–9.

31. Suhailim JL, Partift GI, Xie Y, et al. Effect of desiccating stress on mouse meibomian gland function. Ocul Surf 2014;12:59–68.

32. Palmiasinapan CK, Schröder BM, Balarin L, et al. Effects of keratin and lung surfactant proteins on the surface activity of meibomian lipids. Invest Ophthalmol Vis Sci. 2013;54:2571–81.

33. Ashraf Z, Pasha U, Greenstone V, et al. Quantification of human sebum on skin and human meibum on the eye lid margin using Sebutape, spectroscopy and chemical analysis. Curr Eye Res 2011;36:553–62.

34. Zengin N, Tol H, Balevi Sükür, Balevi S, et al. Tear film and meibomian gland functions in psoriasis. Acta Ophthalmol Scand 1996;74:358–60.

35. Fu DJ, Thomson C, Lunny DP, et al. Keratin 9 is required for the structural integrity and terminal differentiation of the palmpntlar epidermis. J Invest Dermatol 2014;134:754–63.