

An Overview of Current Glaucomatous Trabecular Meshwork Models

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An overview of current glaucomatous trabecular meshwork models 24

25 Abstract: The trabecular meshwork (TM) is a complex tissue that regulates aqueous humor 26 outflow from the eye. Dysfunction of the TM is a major contributor to the pathogenesis of 27 open-angle glaucoma, a leading cause of irreversible blindness worldwide. The TM is a porous 28 structure composed of trabecular meshwork cells (TMC) within a multi-layered extracellular 29 matrix (ECM). Although dysregulation of the outflow throughout the TM represents the first 30 step in the disease process, the underlying mechanisms of TM degeneration associate cell loss 31 and accumulation of ECM, but remain incompletely understood, and drugs targeting the TM 32 are limited. Therefore, experimental models of glaucomatous trabeculopathy are necessary for preclinical screening, to advance research on this disease's pathophysiology, and to develop 33 34 new therapeutic strategies targeting the TM. Traditional animal models have been used 35 extensively, albeit with inherent limitations, including ethical concerns and limited 36 translatability to humans. Consequently, there has been an increasing focus on developing 37 alternative in vitro models to study the TM. Recent advancements in three-dimensional cell 38 culture and tissue engineering are still in their early stages and do not yet fully reflect the 39 complexity of the outflow pathway. However, they have shown promise in reducing reliance 40 on animal experimentation in certain aspects of glaucoma research. This review provides an 41 overview of the existing alternative models for studying TM and their potential for advancing 42 research on the pathophysiology of open-angle glaucoma and developing new therapeutic 43 strategies.

44 Keywords: glaucoma model, trabecular meshwork, 3D culture, in vitro model, tissue 45 engineering, outflow

46

Introduction 47

Glaucoma is a blinding optic neuropathy affecting over 70 million people worldwide¹. Its 48 most important risk factor is elevated intraocular pressure (IOP). The trabecular meshwork 49 (TM) plays a key role in the pathophysiology of glaucoma. This filter is located within the 50 iridocorneal angle and constitutes the main outflow pathway for the aqueous humor. It is a 51 52 fenestrated triangle-form structure in which trabecular meshwork cells (TMC) populate a multi-layered extracellular matrix. The TM tissue is not a rigid structure, but a highly 53 dynamic, avascular filtration system that has a multitude of roles, including filtering aqueous 54 55 humor of waste material, sensing and regulating IOP through the mechanical stretch, and altering ECM composition and deposition ²⁻⁴. The TM contributes to the regulation of IOP by 56 regulating the outflow of aqueous humor from the eye's anterior chamber, primarily through 57 the juxtacanalicular tissue and the endothelium of Schlemm's canal (SC)^{2,5}. The 58 59 juxtacanalicular tissue, located adjacent to the inner wall of SC, plays a crucial role in the 60 regulation of outflow resistance. It consists of specialized cells and extracellular matrix (ECM) components that modulate the flow of aqueous humor ⁶. Dysfunction of the TM and the inner 61 62 wall of SC is the cause of IOP elevation and represents the *primum movens* of primary openangle glaucoma (POAG) but the precise mechanisms at the origin are still unclear. It is 63 known that abnormal TMC function and excess of ECM deposit contribute to TM stiffening 64 in POAG⁷. Alterations in the structure and composition of the TM ECM generate aqueous 65 humor outflow resistance. Cell-ECM interactions within the TM and SC play a role in 66 67 regulating outflow resistance. Cellular responses to ECM components and their remodeling can influence the contractility and overall functionality of the tissue^{6,8,9}. The endothelial cells 68 lining the inner wall of SC exhibit contractile properties. Contraction and relaxation of these 69 cells regulate the diameter of the canal and thereby influence outflow resistance^{10,11}. Tight 70 junctions between endothelial cells of SC also contribute to the formation of a barrier that 71 controls the paracellular movement of aqueous humor¹². A second 'unconventional' outflow 72

pathways exists in the human eye, but it only accounts for less than 10% of the total outflow,
and thus does not significantly contribute to the regulation of IOP in the normal eye¹³.

The development of robust models of physiologic and pathologic TM is necessary to study the different mechanisms at the origin of this deregulation but also to develop and test treatments targeting TM, which are currently rare. Several experimental models have been described with advantages and disadvantages, from animal models or perfusion-cultured anterior segments to cell culture models.

To study the pathophysiology of the TM or its changes under the effects of treatments, 80 81 all models that artificially increase the IOP by blocking or sclerosing the outflow pathways, for example using laser photocoagulation, episcleral vein cauterization, injection of 82 83 microbeads or hyaluronic acid into the anterior chamber or hypertonic saline injection into the 84 episcleral veins are not appropriate as they do not mimic the natural course of glaucomatous dysfunction to the aqueous outflow pathway¹⁴. Indeed, a distinction must be made between 85 animal models of ocular hypertension made to study RGC loss and those specifically 86 87 developed for the study of TM.

88

Models of glaucoma exist in mice, rats, dogs, cats, and primates¹⁵. Each has its challenges, 89 including the availability of genetic resources and difficulty of genetic manipulation, ethical 90 considerations, cost, and maintenance. Translational research in glaucoma is faced with 91 numerous challenges. One significant factor is the divergence in eye anatomy between 92 93 animals and humans. Furthermore, the etiology of the disease differs between glaucoma animal models and human patients. Additionally, there are notable disparities in study design 94 95 and statistical analysis methods employed in preclinical and clinical investigations. Moreover, significant differences exist in terms of dosage, scale, timing of intervention, methodologies, 96

endpoints, age groups studied, and the presence or absence of IOP-lowering treatment, which 97 further complicates the comparison between preclinical and clinical studies in glaucoma¹⁶. 98 99 The animal whose TM anatomy is the closest to humans is the primate, but their use is limited because of their expensive price, the inaccessibility of transgenic strains and the rise of ethical 100 101 concern. Most preclinical studies in the field of glaucoma used mice as their outflow tract 102 anatomy is comparable to humans. It is described as follows: TM, juxtacanalicular tissue, Schlemm canal, collector channels, and episcleral veins¹⁵. However, the main anatomical 103 104 difference is that the human TM structure consists of 9 to 18 trabecular beams whereas in the 105 mouse TM, there are only 2 to 5 layers of lamellae. This difference in the structure of the 106 animal TM as well as the regulation of aqueous humor excretion constitutes a limitation to use 107 of animal models. The results obtained with animal models are therefore not always 108 transposable to humans although the majority of IOP lowering glaucoma medications in human are also effective in mouse eyes¹⁶. Moreover, the lack of cooperation of animals and 109 110 the use of anesthesia to perform experimentation affects the results. The IOP values vary by 111 animal breed, method of sedation, and measure. In general, excessive restraint, inadequate 112 positioning, or lack of experience in the use of equipment can increase IOP. Last but not least, they are questioned from an ethical point of view. Recently, a consensus recommendation for 113 114 mouse models to study the aqueous humor outflow was published in 2022 to set standard practice in this field ¹⁷. The most useful models are genetic models that allow OHT by 115 transduction of the TM with glaucoma-related genes¹⁸ (e.g., $MYOC^{19,20}$, $TGF\beta^{21-23}$, $GREM1^{24}$, 116 $CTGF^{25}$, $DKK1^{26}$, $SFRP1^{27}$, $CD44^{28}$). To meet the expectations of this consensus 117 recommendation animal models used to study outflow physiology must¹⁷: 118

- 119 have an open iridocorneal angle
- 120 present decreased outflow facility
- 121 present elevated IOP

include morphologic descriptions of the conventional outflow tract (TM, Schlemm's canal, collector channels, and intrascleral and episcleral regions) preferably by
 electronic microscopy to be able to analyze the organization of the ECM

125 - and assessment of TMC numbers.

The Myoc^{Y437H} mouse, responsible for a juvenile glaucoma, is the best-known and most used²⁰. The genetically modified mice express high levels of mutant myocilin in the TM and sclera resulting in a decline of AH outflow facility and a secondary increase of IOP. Morphological changes can be observed in the TM: decreased intertrabecular spaces and a progressive loss of TMC. However, the level of IOP can vary based on the genetic background of the mouse strain selected for inducing the MYOCY437H mutation²⁹.

132 In recent years, progress in three-dimensional cell cultures and tissue engineering 133 fabrication has offered promising approaches to reduce the use of animal models, partly 134 encouraged by the three Rs rule (Reduce the number of animals used; Replace the living 135 animal with alternative experimental techniques; and Refine the techniques to minimize animal suffering) ^{30,31}. While conventional 2D cell culture models represent an attractive 136 137 alternative to animal models for analyzing the TM and allow for more accurate assessment at 138 the cellular scale, they are limited by the absence of differentiation, polarization, or 139 relationship with an extracellular matrix, making them insufficient in reflecting the actual porous architecture of the TM ^{32,33}. However, this deficient architecture is precisely the cause 140 of its dysfunction ³⁴. *Ex vivo* models such as organ perfusion chambers, whole tissue explants, 141 142 and decellularized tissues are commonly used natural sources; however, their limited availability restricts their use in perfusion studies and drug testing^{35,36}. This is why a three-143 144 dimensional (3D) cellular model of the TM would be an interesting tool to advance research 145 on this pathology by considering the biomechanics, which is a key element in the pathophysiology of glaucoma⁴. 3D cell culture would enable the recreation of the 146

microenvironment encountered *in vivo* and provide cells with an environment allowing them 147 to interact with each other³³. This, in turn, would lead to a better understanding of the 148 physiological functioning of the TM, its behavior under conditions of stress or toxicity, and 149 the effect of treatments³⁷. Additionally, *in vitro* models provide a more precise analysis of cell 150 behavior and molecular mechanisms involved in the pathology than do animal experiments³⁸. 151 152 This approach allows to investigate specific biological phenomena in isolated cells, eliminating potential confounding factors present in whole organisms. However, it is crucial 153 154 to acknowledge that while in vitro systems provide controlled environments, they cannot fully 155 replicate the complex conditions and interactions that occur within a living organism. Technological advances in TM in vitro models can help fill the gap in considering the 156 157 mechanistic modifications involved in glaucoma, such as changes in porosity resulting from alterations in morphology and the mechanical properties of the interaction between TMC and 158 159 ECM.

In this literature review, we will provide an overview of existing alternative TM models to animal experimentation. We will detail the different cell types used, culture modes, and means to obtain a pathological model. Finally, we will focus on the potential applications of these different models.

164

165 Available cell types

166

167 *Primary cell cultures:*

168 TM is composed of three regions, from the anterior chamber to the Schlemm canal: the uveal 169 meshwork, the corneoscleral meshwork, and the juxtacanalicular tissue, the location of 170 greatest resistance to AH outflow. TMC have a different organization in these three regions, but it is complex to isolate cells from only one of these three regions. Thus, TMC cultures are
generally a mixture of cells from all three regions ³².

173 Cultures of primary human TMC (HTMC) are sampled from donor eye tissue 174 commonly from a corneal rim discarded from corneal transplant³⁹. Whole globe or anterior 175 segment from normal subjects, developing fetuses, or patients with glaucoma can also be 176 purchased.

177 Cultures of TMC from patients with glaucoma are more difficult due to the accelerated
 178 loss of these cells. Nevertheless, they retain their glaucomatous characteristics after culture⁴⁰.

TMC change their morphology after 6 to 8 passages, thus it is recommended to use TMC from human eyes before the 7th passages. Methods used to validate that cells are TMC is required for publications including at least responsiveness of myocilin expression by cells to dexamethasone ³².TMC cultures can also come from animal eyes with the limitations that this brings.

184

185 Immortalized human TM cells

Immortalized TMC lines can be generated with TMC transfected with an original defective
mutant of the SV40 virus^{41,42}. However, during the immortalization process, some properties
of the TMC can be lost, for example the myocilin expression. It is recommended that cell line
findings be replicated with non-immortalized TMC³².

190

191 Induced pluripotent stem cell-derived trabecular meshwork (iPSC-TM)

192 The reproducibility of primary cell cultures is a challenge and immortalized cell lines are 193 considered poorly relevant to TMC physiology and disease patterns. The experimental 194 transplantation of iPSC-derived TM (iPSC-TM) highlighted the huge therapeutic potential of 195 these new human cell models, offering perspectives for toxicological or therapeutic 196 evaluations^{43,44}. Moreover, the pathogenesis of the glaucomatous disease is explained by a 197 TMC loss greater than the physiological age-related cell loss. This loss has been suggested to 198 affect the ability of the human TM to regulate aqueous humor outflow and to lead to IOP 199 elevation. In addition to providing a source of reproducible and valuable cells for the 200 constitution of an *in vitro* model iPSC-TM cells are promising autologous cell sources that 201 can be used to regenerate the declining TMC population and function of IOP regulation^{45,46}.

202

203 TM progenitor cells

204 A population of progenitor cells has been identified in Schwalbe's Ring, which is located at the junction between the corneal endothelium and the anterior portion of the TM⁴⁷. These 205 206 progenitors can be isolated and expanded, and studies have shown that they have the ability to differentiate into both keratocytes and TMC. Zhang et al. developed an optimized method to 207 208 expand multipotent progenitors from human TMC in a two-dimensional (2D) culture followed 209 by three-dimensional (3D) culture in Matrigel using a modified embryonic stem cell medium⁴⁸. The expanded cells expressed TM markers, embryonic stem cell (ESC) markers, 210 211 and neural crest (NC) markers. Although some markers were lost after passage, the cells 212 regained the markers when seeded on 3D Matrigel via activation of the canonical BMP signaling⁴⁸. These cells can be used in an *in vitro* model system to help better understand how 213 214 TM is affected in glaucoma and whether TM progenitor cells may have potential therapeutic 215 applications for glaucoma.

216

217 Generation of Pathological Models:

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As discussed earlier, obtaining cells from the glaucomatous TM is a challenging task forresearchers, which is why different molecules are used to induce a diseased phenotype.

Applying mechanical stress to the TM can induce changes similar to those observed in
glaucoma. This can be done by stretching or compressing the tissue using a variety of devices.
In a previous study, Schlunck et al. demonstrated that the stiffness of the ECM could alter the
structure of the cytoskeleton of trabecular cells as well as the profiles of certain protein
expressions ⁸.

Various chemicals can be used to induce glaucomatous changes in the TM, such as transforming growth factor $\beta 2$ (TGF- $\beta 2$), and have been shown to contribute to the changes in the ECM of the TM.

TGF- β 2 is a profibrotic cytokine known to be elevated in the aqueous humor of patients with glaucoma^{49,50}. It has been used in many studies as a model of pathological TM ^{41,42,51–53}. Studies have revealed that TGF- β 2 can increase intraocular pressure (IOP) by promoting the synthesis of certain ECM components by trabecular cells (collagens, fibronectin) through epithelial-mesenchymal transition in TMC ^{54,55}. Furthermore, TGF- β 2 can increase cell rigidity by triggering the formation of Cross-linked Actin Networks (CLANs) via the Rho-GTPase pathway^{53,56}.

Hydrogen peroxide (H2O2), another molecule used to induce a glaucomatous phenotype, is a
chemical compound with powerful oxidizing properties and has been shown to promote
cellular senescence, rearrange cytoskeletal structure, and increase proinflammatory mediators
such as IL-6, IL-8, and endothelial–leukocyte adhesion molecule 1 (ELAM-1) in the TM⁵⁷.

Endothelin-1 (ET-1) is another biomarker found in the aqueous humor of patients with POAG^{58,59}. It has been shown that ET-1 can induce TMC contraction in culture and that it can affect the outflow facility^{60,61}. Wang et al. showed that in a cultured HTMC model, treatment with ET-1 increased the expressions of fibronectin and collagen IV, and that in an ex-vivo model, IOP increased after ET-1 administration⁶². Zhou et al. also used ET-1 in a whole eye perfusion system and found a decreased outflow after ET-1 exposition and successfully tested
 several pretreatments to reverse this effect⁶³.

Benzalkonium chloride (BAK) *in vitro* induced apoptosis, oxidative stress, and also an IOP
increase, with reduction of aqueous outflow *in vivo*. BAK enhances all characteristics of TM
degeneration typical of glaucoma and causes degeneration in acute experimental conditions,
potentially mimicking TM degeneration⁶⁴. In an *in vitro* 3D TM model, Bouchemi et al.
showed that BAK disorganized the TMC and decreased their number resulting in an
enlargement of spaces between cells ⁶⁵.

253

254 Available 3D models:

255

256 *3D scaffolds culture*

The first successful scaffold was a micro-fabricated SU-8 porous structure, where TMC were 257 cultured to study steroid-induced glaucoma. Scaffolds with pores of approximately 20 258 259 micrometers in thickness, which were seeded with primary HTM cells, were able to imitate 260 some of the normal tissue functions in vivo. This included being able to induce or reverse glaucomatous conditions using medication ^{66,52,67}. A follow-up study showed that applying a 261 262 hyaluronic acid-containing hydrogel coating to the SU-8 scaffold improved cell proliferation. Over time, various technologies and materials have been explored, including traditional 263 polymeric filters, SU-8 membranes, electrospun nanofibers, and other methods. These 264 265 methods offer precise control over morphological characteristics such as porosity and beam 266 thickness in both 2D and 3D environments. However, the stiffness of the scaffold cannot be directly controlled using most of these methods. Despite these limitations, 3D cultures have 267 268 the potential to create an in vivo-like microenvironment for HTM cell growth. Tissue engineering aims to produce functional biomimetic replicas of tissues of interest, but only a 269

limited number of studies have been reported on bioengineered 3D HTM in vitro models. 270 271 These models partially mimic normal tissue function and provide a platform for drug testing 272 and evaluating the effectiveness of different treatment options. Wlodarczyk-Biegun et al. 273 studied the biofabrication technique of melt electrowriting (MEW), a marriage between 274 electrospinning and 3D printing, as a means of producing fibrillar and porous scaffolds with 275 thermoplastic polymers that replicate the multilayer and gradient structure of the natural HTM ⁶⁸. HTMC cultured on these scaffolds maintained the phenotype of native HTMC and 276 277 infiltrated the scaffolds. However, some may argue that these models are more comparable to 278 a 2D cell culture system rather than a true 3D model, as they cannot fully replicate the 3D 279 cell-ECM interface apart from the ECM secreted by the HTM cells grown on top of the 280 synthetic polymer scaffold.

281

282 Hydrogel- based TM scaffolds

283 Recent research has focused on using hydrogels as scaffolds to study the behavior of HTMC in response to environmental changes and disease conditions^{69,70}. Hydrogels are networks of 284 285 crosslinked, hydrophilic polymers used to recapitulate the 3D architecture of organ systems in tissue engineered models. These materials are so useful in cell culture because they provide a 286 biocompatible, degradable, hydrated microenvironment that mimics the cell-ECM 287 288 interactions of natural tissues. Hydrogel scaffolds offer higher control over the morphology, 289 stiffness, and 3D environment compared to photolithography and electrospinning, while also 290 maintaining structural integrity.

Ideally, 3D cell culture matrices can reproduce the features of the ECM to closely resemble the *in vivo* environment. The interaction between cells and ECM is essential for a range of biophysical and biochemical functions, such as the transportation of signaling molecules, nutrients, and waste metabolites, as well as mechanical integrity. Thus, these 295 matrices need to reflect the specific ECM characteristics of each tissue for a given application. 296 Moreover, the mechanical properties of 3D matrices are also significant, as they can directly influence cell adhesion, thereby affecting both the shape and response of cells⁷¹. The 297 utilization of degradable scaffolds presents an opportunity for more molecular research, as 298 299 opposed to permanent ones. Hydrogels can be created using various synthetic or natural 300 components. In tissue engineering, natural polymers are the most commonly used approach to developing hydrogels⁷². Collagen, fibrin, and elastin, which are components of the ECM of 301 302 the TM, have been used as attachment factors for HTMC to study specific functions and interactions ⁶⁵. 3D Corning® Matrigel® Matrix (Corning Life Sciences, Tewksbury, MA, 303 304 United States) contents several proteins found in extracellular matrix (ECM) such as laminin, collagen IV, heparin sulfate proteoglycan, and entactin/nidogen^{73,74}. Several studies 305 306 demonstrated that unlike cells cultured on traditional 2D planar surfaces, cells in 3D scaffolds 307 are more physiologically relevant concerning in vivo characteristics exhibited by in-vivo surrogates⁷⁵ (figure 1). Vernazza et al. conducted a study to compare the response of HTMC 308 309 in 2D and 3D in vitro models following chronic stress exposure. Their results revealed that 310 3D TMC cultured on Matrigel exhibited a higher sensitivity to the production of intracellular 311 reactive oxygen species induced by hydrogen peroxide treatment compared to 2D cultures. 312 Furthermore, the 3D models demonstrated a more precise regulation of apoptosis triggers and cell adaptation mechanisms than the 2D models ³³. Another scaffold-based approach by 313 Osmond and colleagues utilized HTMC cultured on various collagen scaffolds containing 314 315 different glycosaminoglycans (GAGs) and different pore architectures to better understand 316 how HTMC respond to changes in their extracellular environment. The cellular response was 317 assessed by quantifying cellular proliferation and the expression of fibronectin, an important 318 extracellular matrix (ECM) protein. Fibronectin plays a crucial role in organizing ECM 319 proteins both among themselves and with trabecular cells, thereby contributing to the

resistance of outflow ^{76–78}. The pore architecture of the scaffolds was altered using the freeze-320 321 casting technique to make both large and small pores that are aligned or with a non-aligned 322 random structure. The composition of the scaffolds was altered with the addition of 323 chondroitin sulfate and/or hyaluronic acid. It was found that HTMC grown on large pore 324 scaffolds proliferate more than those grown on small pores. There was an increase in the fibronectin expression with the incorporation of GAGs, and its morphology was changed by 325 326 the underlying pore architecture. These works offer a better understanding of how human TMC behave in response to alterations in their microenvironment and highlight the 327 importance of the mimicry of the 3D strucutre⁷⁹⁻⁸¹. However, the study did not explore how 328 329 the constructs would react under conditions that induce glaucoma. Furthermore, if the 330 accumulation of extracellular matrix (ECM) proteins is a characteristic feature of the pathogenic process in glaucoma, it is important to highlight that cell proliferation is not^{77,82}. 331 332 Therefore, it is crucial to determine whether cells can survive under normal conditions on 333 these new substrates. However, it should be noted that the ability to proliferate does not 334 necessarily indicate an appropriate glaucoma model.

335 3D bioprinting can produce a variety of architectural patterns on a wide array of biomaterials. 336 Li and colleagues developed a hydrogel using a tissue-engineering approach for HTM. The 337 hydrogel consisted of ECM biopolymers and normal HTMC obtained from a donor. By 338 mixing HTMC with collagen type I, hyaluronic acid (HA), and elastin-like polypeptide (ELP) 339 - each containing photoactive functional groups - researchers were able to create HTM 340 hydrogels in various sizes and shapes. Short UV cross-linking, mediated by photo-initiators, 341 was used to solidify the hydrogels. To induce glaucomatous changes, dexamethasone (DEX) was administered, and the therapeutic effects of the ROCK inhibitor Y27632 were evaluated⁸³. 342 To create an in vitro 3D TM scaffold for potential use as a tissue scaffold in glaucoma 343 patients after trabeculectomy, Waduthanthri et al. developed a hydrogel peptide called 344

MAX8B which partly mimics the motif of cellular integrins and enables interactions with ECM components ⁸⁴. The scaffold material demonstrated the ability to undergo shear-thinning and exhibited biocompatibility, facilitating appropriate growth and proliferation of TMC in tightly packed cell monolayers resembling typical TMC morphology. Moreover, the MAX8B scaffold was utilized in an in vitro perfusion system to investigate the impact of Dexamethasone on the outflow facility of the trabecular meshwork proving the effectiveness of this three-dimensional (3D) model as a platform for drug testing⁸⁴.

352

353 Spheroids

354 Although 3D culture techniques have gained popularity for their ability to better mimic in 355 vivo environments, there are some limitations when it comes to replicating physiological and 356 pathological conditions of human TM. This is because the use of scaffolds in 3D cultures is 357 not reflective of the absence of such structures in human TM. However, 3D spheroid cell 358 cultures have recently emerged as a promising alternative to conventional 2D cell cultures, 359 particularly as in vivo models for various diseases. These spheroids allow for more 360 intercellular interactions in a 3D space, potentially resulting in protein networks that resemble 361 those found in real tissues. This makes it possible for 3D spheroids to replicate biological features associated with real tissues. 362

The spheroid model of TM refers to a 3D culture system that mimics the structural and functional properties of the TM in the eye. HTMC have been cultured as spheroids *in vitro* to study their role in glaucoma. These spheroids have been shown to exhibit features of the TM *in vivo*, such as the presence of ECM components and cytoskeletal proteins. These spheroids have been shown to respond to mechanical stress and exhibit physiological responses similar to those observed *in vivo*. These spheroids have been shown to be structurally and functionally similar to the TM *in vivo* and have been used to study the effects of various drugs 370 on TMC behavior. 3D HTM spheroids became significantly and differently smaller and stiffer 371 in response to TGF- β 2 or dexamethasone stimulation^{41,85}. Watanabe et al. successfully 372 obtained 3D HTM spheroids and found that TGF β 2 significantly induced the down-sizing and 373 stiffness of 3D spheroids from human orbital fibroblasts, and those effects were substantially 374 inhibited by a ROCK-inhibitor.^{42,86}

375

376 *Outflow studies*

377 Perfusion studies of outflow in HTMC began in the late 1980s and have since evolved to 378 include a range of techniques and models. One of the earliest studies involved the use of filters to culture HTMC and measure hydraulic conductivity using a pressure/flow circuit⁸⁷. 379 380 This study led to further investigations into the biomechanics of HTMC. The perfusion 381 system developed by Yubing Xie's group enabled continuous pressure monitoring at different flow rates to investigate the effects of drugs such as Lat-B, ROCK inhibitors, or TGFβ2^{52,66,67}. 382 383 As previously mentioned, 3D culture models of TMC are superior to 2D models due to the 384 ability to enable cell-cell and cell-ECM interactions. However, these 3D models fail to 385 reproduce the dynamic continuous supply of nutrients, oxygen, and removal of metabolic waste products. Recent advances propose models that combine the benefits of 3D culture with 386 387 milli-fluidic techniques to improve the physiological relevance of the culture and address the 388 issues related to cell responses under static culture conditions. Microfluidic systems allow for 389 the creation of a 3D microenvironment that mimics the *in vivo* conditions of the TM, 390 including the presence of shear stress and fluid flow. Recently, the MAX8 peptide-hydrogel 391 scaffold and a 3D Matrigel® model have been tested in perfusion chambers to evaluate their use as artificial TM scaffolds^{39,84}. 392

393 In their closed-circuit *in vitro* model developed by Tirendi et al., 3D-HTMCs cultured 394 in Matrigel were provided with a continuous medium supply. This was achieved by connecting single-flow bioreactor culture chambers to a peristaltic pump. The milli-fluidic technology as well as the 3D culture model mimicked cell responses found *in vivo* as a result of the increase in outflow resistance ⁵⁷. This type of model can be used to investigate the effects of various factors on TM function, such as mechanical stress and changes in ECM composition.

400

401 *Ex vivo models*

402 Ex vivo models, specifically perfusion studies, utilizing animal eyes such as pigs, cows, and primates, have been instrumental in advancing our understanding of glaucoma ^{88–91}. 403 404 These models offer valuable insights into the dynamics of aqueous humor outflow and 405 provide a platform to investigate the effects of various experimental interventions on the 406 disease. By perfusing the enucleated eyes with a controlled flow of fluid, researchers can 407 mimic physiological conditions and measure parameters such as intraocular pressure and 408 outflow facility. These models have helped elucidate the mechanisms underlying glaucoma and evaluate potential treatments^{63,89}. For example, Zhou et al. developed a platform to 409 410 simultaneously evaluate outflow facility and its time- and dose-dependent responses to 411 treatments of 20 eyes. They used whole porcine and bovine eyes to develop a perfusion system and studied the regulation of outflow facility by endothelin-1, nitric oxide donor, and 412 sphigosine1-phosphate ⁶³. 413

However, it is important to acknowledge the limitations of ex vivo models. They do not fully replicate the complex in vivo environment of the eye, including interactions with surrounding tissues and systemic factors. Additionally, the use of animal eyes may introduce speciesspecific differences that may not fully reflect human physiology. Given these limitations and the fact that they do not represent an alternative to the reliance on animal experimentation, we will primarily focus on human models. The human anterior segment perfusion culture model is a valuable tool for studying the TM and aqueous humor outflow in glaucoma^{89,92,93}. *Ex vivo* models possess several significant benefits compared to other models, including their ability to maintain the structure of pathways and their capacity to facilitate analysis in nearly ideal physiological conditions⁹⁴. Outflow facility measurements can be performed *ex vivo* or *in vivo*, with *ex vivo* measurements offering a simpler approach by avoiding confounding factors that are difficult to control. However, *in vivo* measurements are more representative of real-life conditions.

Bahler et al. used perfusion organ culture of human anterior segments to study the effect of prostaglandin on the trabecular outflow. Since this human anterior segment culture model lacks a choroid or functional ciliary body, the uveoscleral pathway is absent. This simplification facilitates the analysis by directly assessing the sclera's impact on outflow facility⁹³.

432 Peng et al. have created an ex vivo model of human corneal rim for perfusion culture 433 experiments as an alternative to the human anterior segment perfusion culture model. This 434 model can be used to study the TM and aqueous humor outflow in glaucoma while improving 435 cost and availability. The corneal rims were affixed to 3D-printed perfusion culture plates and perfused in constant flow mode. Pressure changes were recorded using a computerized system. 436 437 TM stiffness of corneal rims treated with dexamethasone was significantly higher than in the control group⁹⁵. Additionally, the model allows histology or immunohistochemistry of the 438 439 TM to investigate biomechanical changes or treatments.

Baudouin et al. examined TM specimens using immunohistology and reverse transcriptase– polymerase chain reaction. Trabecular specimens of glaucomatous patients showed extremely low densities of trabecular cells and the presence of cells expressing fractalkine and fractalkine receptor and their respective mRNAs⁶⁴. These explants methods have the advantages to retain tissue architecture and cellular interactions closer to in vivo conditions as opposed to traditional cell culture methods. They are suitable for studying tissue responses and drug effects at the cellular level⁹⁶. The low cell count of TMC in TM explants from glaucomatous patients can be circumvented by using TM from healthy donors and exposing them to TGF- β 2. The addition of TGF- β 2 to healthy TM permits reproduction of the changes in TM cell cytoskeletal organization and ECM compaction, while providing sufficient material for a transcriptomic study^{2,82,97}.

451

452 **Discussion**

This article discusses the importance of developing models of TM, a structure within the eye that plays a crucial role in regulating IOP, to study the pathophysiology of glaucoma. The TM is a dynamic filtration system that helps regulate IOP by controlling the outflow of aqueous humor.

457 Developing new 3D in vitro models is crucial to studying TM pathophysiology in 458 glaucoma. They mimic the physiological microenvironment of the TM, providing a more 459 physiologically relevant model than traditional 2D cell culture methods⁹⁸.

One of the key advantages of these 3D models is that they reduce the need for animal studies, which can be costly, time-consuming, and ethically challenging^{15,30}. *In vitro* models can be used as a complementary tool to animal studies, as they can provide useful data on

463 mechanisms and drug efficacy before moving to animal models or clinical trials.

While *in vitro* models offer several advantages, they also have limitations that need to be considered. One of the main challenges is that *in vitro* models may not fully recapitulate the complexity of the TM *in vivo*, such as interactions with other tissues and the influence of systemic factors. To address this limitation, researchers often use a multiplicity of models to collect data for a particular question. For example, to study the modification of ECM, a natural hydrogel medium that closely resembles the components of TM ECM is more interesting than a synthetic one. As it provides a more physiologically relevant environment
that can better mimic the ECM interactions in the TM. Similarly, a microfluidic bioreactor
can be used to study the effect of sheer stress or biomechanical impact on TMC^{66,99}. This type
of model allows researchers to control the flow of fluids and apply mechanical forces to the
cells, providing more accurate simulations of the TM microenvironment. A comparison of
innovative 3D TM models and measured outcomes is presented in supplementary table 1.

Furthermore, biomimetic 3D in vitro models, in addition to enhancing our 476 477 understanding of TM tissue biology and outflow pathology, have the potential to be used therapeutically for restoring compromised TM function¹⁰⁰. Promising research has 478 479 demonstrated the effectiveness of stem cell therapy in repairing TM tissue and preserving vision in glaucoma patients ⁴⁶. Moreover, the presence of TM progenitor cells capable of 480 differentiating into functional TM cells further supports the potential for tissue repair^{101,102}. 481 482 Advanced biofabrication techniques allow for the creation of scaffolds that closely mimic the 483 native ECM and provide cues for stem or progenitor cell differentiation, replicating cellular responses observed in vivo¹⁰³. By incorporating biomaterials alongside TM progenitor cells, 484 the development of a delivery system for effective stem cell therapy can be facilitated. 485

In conclusion, the use of multiple models that can better replicate the different aspects of the TM *in vivo* can provide more robust and accurate data. By using a combination of *in vitro*, *ex vivo*, and *in vivo* models, researchers can gain a more comprehensive understanding of glaucoma pathology and develop better treatments for this disease.

However, it is important to consider the limitations of non-animal. The progress made
in the alternative models presented in this study does not imply that we can completely
eliminate the need for animal experimentation at present. In vivo experiments enable a
substantial prediction of the effect of hypotonic treatment on IOP, even if the organization of
their outflow system is not totally similar to that of humans¹⁶. These alternative models are

still in their early stages and may not fully replicate the complexity of the TM or its 495 496 interactions within the eye. They may not provide the same comprehensive data as animal 497 models, particularly in terms of assessing IOP, estimating natural flow rate, accessing the 498 outflow facility, evaluating cellularity, tissue integrity, and capturing natural expression 499 profiles as it would be in a living in vivo system. Additionally, organ culture has a significant 500 limitation whereby the regulation of IOP relies solely on external manipulative regulations, 501 lacking intrinsic regulation in enucleated eyes. Nonetheless, despite these current limitations, 502 the progress made in developing these alternative models is encouraging. While they may not 503 completely replace the need for animal models, they do hold the potential to significantly 504 reduce their utilization, provided of course that the trabecular cells used are not derived from 505 animals. 506 Overall, this progress in *in vitro* and *ex vivo* models offers a promising tool for studying the

507 TM in glaucoma and reducing the need for animal studies. While it has limitations, it provides

a more physiologically relevant model than traditional 2D cell culture methods, and its

- 509 potential applications in drug discovery and testing make it a valuable addition to glaucoma
- 510 research.
- 511

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- 836 wrote the initial manuscript. A. L. and C.B. provided critical insights and expertise in the field,
- 837 contributing to the interpretation of the findings and revision of the manuscript. C.B.
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Figure 1: Confocal microscopy image of the 3D cultured pHTMCs in Matrigel. The pHTMC is
organized in a mesh conformation with interconnections and the formation of intercellular spaces.
Actin fibers are stained in red by phalloidin, membranes with DiO (green), and nuclei with DAPI
(blue). Magnification 200X.

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