Tear-film breakup: the role of membrane-associated mucin polymers

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Mucin polymers in the tear film protect the corneal surface from pathogens and modulate the tear-film flow characteristics. Recent studies have suggested a relationship between the loss of membrane-associated mucins and premature rupture of the tear film in various eye diseases. This work aims to elucidate the hydrodynamic mechanisms by which loss of membrane-associated mucins causes premature tear-film rupture. We model the bulk of the tear film as a Newtonian fluid in a two-dimensional periodic domain, and the lipid layer at the air-tear interface as insoluble surfactants. Gradual loss of membrane-associated mucins produces growing areas of exposed cornea in direct contact with the tear fluid. We represent the hydrodynamic consequences of this morphological change through two mechanisms: an increased van der Waals attraction due to loss of wettability on the exposed area, and a change of boundary condition from an effective negative slip on the mucin-covered areas to the no-slip condition on exposed cornea. Finite-element computations, with an arbitrary Lagrangian-Eulerian scheme to handle the moving interface, demonstrate a strong effect of the elevated van der Waals attraction on precipitating tear-film breakup. The change in boundary condition on the cornea has a relatively minor role. Using realistic parameters, our heterogeneous mucin model is able to predict quantitatively the shortening of tear-film breakup time observed in diseased eyes.

I. INTRODUCTION

The tear film is essential to the health of the eye as it protects the sensitive epithelium of the cornea and lubricates the eyelid. Only several micrometers thick, it features a complex structure and interesting rheological properties [1]. On the corneal surface is a mucus layer up to half a micrometer in thickness, outside which is an aqueous layer of 2 to 5 micrometers that forms the bulk of the tear film. Farthest out is a thin lipid layer that interfaces with the ambient air (Fig. 1). The mucus layer comprises two groups of mucin polymers, membrane-associated mucins (MAMs) attached to the epithelial surface that form a glyocalyx covering the cornea, and gel-forming mucins [2, 3]. Each group contains several kinds of mucin molecules. Inside the aqueous layer there is also a small amount of soluble mucins. Altogether the tear film has about 10 mucin species of different molecular weight and properties, produced by different cells and glands in the eye [2, 4]. As this work focuses on the MAMs only, for simplicity we will use these three terms interchangeably hereafter: the mucus layer, the MAMs and the glyocalyx.

The MAMs play an important role in preventing microbial and fungal infections at the ocular surface [6]. Clinical evidence has implicated MAM loss in eye infections and the dry eye syndrome [7, 8]. Their highly glycosylated extracellular domains form a tight mesh structure that protects the cornea from pathogenic invasion. In case of a bacterial attack, these extracellular domains get clipped from the epithelial surface and bind with the pathogen to activate an internal signaling pathway for mucosal maintenance and repair [9, 10]. Thus, a severe infection can remove MAMs from the epithelial glyocalyx barrier and expose the cornea to the aqueous layer and to direct attack by the pathogens (Fig. 1).

Aside from their role as a physical barrier, recent studies have suggested a second role for MAMs in stabilizing the tear film against premature rupture [3, 11]. After each blink, a new tear film is coated over the cornea, which persists in healthy eyes for an extended period before breaking up. This tear-film rupture time $t_{rup}$ falls

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FIG. 1: Schematic representation of a tear film with a partially damaged mucus layer, with the membrane-associated mucins (MAMs) clipped from the central portion of the corneal epithelium and shed into the aqueous layer. The protrusions on the epithelium represent the corneal microplicae [2, 5].

In a wide range in healthy eyes [12], but averages to between 10 and 15 seconds [13, 14]. In diseased eyes, however, $t_{rup}$ is shortened to 5 seconds or less, the most severe cases corresponding to instantaneous breakup [15–17]. The reasons have been hypothesized to be the loss of MAMs [3, 11]. A short $t_{rup}$ exposes the cornea to contaminants and pathogens in the ambient air. Thus, the MAMs may protect the cornea through two separate mechanisms, as a physical barrier to pathogens and as a tear-film stabilizer. It is the latter that has inspired the fluid-mechanical study reported here.

How can the loss of MAMs precipitate tear-film breakup? Multiple mechanisms contribute to tear-film breakup, including evaporation [18, 19], osmotic flux between the tear film and the cornea [20], non-uniformity in the lipid layer [21, 22], gravity-driven drainage [23, 24], and van der Waals attraction between the tear-air interface and the corneal surface [12, 25–27]. Among these, the van der Waals attraction is most likely to be affected by MAM loss. The other mechanisms produce a rapid initial thinning of the tear film down to about 0.5 micrometer, after which the van der Waals force dominates and dictates the slower final stage of breakup [12, 19, 22, 28].

Observations suggest that the mucin loss happens in localized regions of the corneal surface, and that the mucin loss amounts to a loss of hydrophilicity. Gipson et al. [2, 5] presented images showing localized MAM alterations in regions of the corneal epithelium due to keratinization. It is widely accepted that the mucus layer serves to keep the corneal surface hydrophilic [11], and that localized loss of wettability can initiate rapid dewetting and breakup [17, 29]. In particular, Gipson et al. [5] concluded that “loss of mucins leads to loss of hydration and, in turn, to the formation of dry spots”. Winter et al. [30] suggested incorporating “varying wettability on the substrate in order to represent unhealthy parts of the cornea”. Based on the above, we hypothesize that the loss of MAMs occurs in a spatially heterogeneous manner on the corneal surface, and locally changes its wettability and increases the van der Waals attraction between the interfaces, thus bringing about premature tear-film breakup. To test this hypothesis, we will represent the loss of MAMs by an elevated van der Waals potential in a heterogeneous mucin model for tear-film breakup.

We also consider a second mechanism via the change of the boundary condition on the corneal surface. Essentially a polymer meshwork, the glyocalyx should hinder the fluid flow next to the cornea. Such an effect has been demonstrated by calculations of simple shear flows over polymer brushes, where the linear velocity profile in the bulk can be extrapolated toward the substrate to determine an effective negative slip length [31, 32]. With the loss of the MAMs, the exposed cornea surface should take on the usual no-slip boundary condition. The loss of negative slip in the mucin-depleted areas should promote tear-film breakup. Such an effect of the boundary condition may be likened to that of substrate textures on the instability of thin films [33–35].

The objective of this study is to investigate how these two factors affect tear-film breakup. Recently, Dey et al. [12] proposed a continuous viscosity model for tear-film breakup by explicitly representing the mucin concentration profile along the depth of the film. An intact glyocalyx was represented by a viscosity profile with a high-viscosity layer atop the cornea, whereas loss of MAMs corresponded to a more or less uniform viscosity profile. This study seeks a more realistic representation of the loss of mucin through an increase in the van der Waals attraction and a change in the boundary condition on the substrate.

II. PROBLEM FORMULATION AND METHODOLOGY

A. Problem description

We pose our problem in a two-dimensional domain that is periodic along the longitudinal $x$ direction, with a homogeneous fluid forming the bulk of the tear film, a lipid layer on top modelled as an insoluble surfactant and a solid substrate with spatially heterogeneous mucin cov-
The setup of the heterogeneous mucin model contains several simplifications. We have neglected evaporation of the tear film [18, 19], osmotic flux between the tear film and the cornea [20], rapid rupture due to lipid globs [22], and gravity-driven drainage [23, 24]. All can be significant factors in tear-film flow as demonstrated by earlier studies. Braun et al. [19] found that most of these factors contribute to a rapid initial thinning the tear-film down to about 0.5 μm, at which point van der Waals interaction becomes the dominant factor. Our model essentially starts from this point onward [12]. Another noteworthy factor is the rheology of the tear film. The tear film shows shear-thinning thanks to the many bio-polymers.
dissolved in it [39, 40], and shear-thinning has been included in a few prior models [27, 41, 42]. Viscoelasticity can also be a factor [43], although an estimation of the mucin relaxation time and concentration in the tear film suggests a negligible elastic stress. Calculations using an Oldroyd-B model indeed show negligible effect of the non-Newtonian rheology. In view of above, and of the focus of the present work on MAM loss, we have excluded the complicating factors from our heterogeneous mucin model.

B. The effective Hamaker constant

There is considerable discrepancy regarding the effective Hamaker constant $A_1$ for tear film over a healthy cornea with intact MAMs. From the pairwise interfacial energies between the various components (cornea, mucus and water), Sharma et al. [25, 44] estimated a value of $A_1 = 10^{-21} \sim 4 \times 10^{-20}$ J, which was used in later studies of tear-film breakup [45]. More recently, Winter et al. [30] found that a much larger value $A_1 = 6.6 \times 10^{-18}$ J was required in order to reproduce experimental data on the enlargement of dry spots in human tear films. Later, Braun et al. [19] used this value to predict tear-film breakup under evaporation and osmotic liquid flux. In their continuous viscosity model, Dey et al. [12] used the same value to predict $t_{rup}$ in good agreement with clinical measurements in healthy human eyes. Based on the above, we have decided to adopt $A_1 = 6.6 \times 10^{-18}$ J in the current model.

We have found no theoretical estimation or experimental measurement of the effective Hamaker constant $A_2$ for a tear film above a MAM-depleted cornea. Tracing the calculations of Sharma [46] to earlier studies of the apolar Lifshitz-van der Waals interactions between two or more substances [47, 48], we can estimate $A_2$ for a binary cornea–water system from their interfacial tension: $A_2 = 4.65 \times 10^{-20}$ J. Comparing this with the $A_1$ values that Sharma [46] estimated using similar arguments, one obtains a ratio $A_2 : A_1$ in the range of 1.16 – 46.5. Taking this as a rough guideline, we will explore $A_r$ between 1 and 4 in this study. At $A_r = 4$, the heterogeneous mucin model already predicts “instantaneous breakup” with a rupture time of a second. Thus, we will not explore $A_r$ values above 4.

C. Negative slip over mucin-covered cornea

The mucus layer contains the membrane-associated and the gel-forming mucins. Some authors suggested that the gel may act as a lubricant to reduce the viscous friction and produce effective slippage on the cornea [3, 49, 50]. Given the low viscosity of water, however, we find it difficult to imagine the gel-forming mucins lubricating the tear-film flow. Therefore, we will focus solely on the role of MAMs in hindering fluid flow. This assumption is informed by calculations of shear flow over polymer brushes and networks, which show that the additional resistance due to the polymers amounts to a negative slip; see, e.g., Fig. 4a of [32] and Fig. 2a of [31]. Thus, in the heterogeneous mucin model we have assumed a negative slip length $\beta_1$ over intact MAMs. Over the portion of the cornea where MAMs have been shed, we impose the no-slip condition: $\beta_2 = 0$.

The value of $\beta_1$ is determined such that our model produces the correct tear-film rupture time $t_{rup}$ for healthy eyes. Dey et al. [12] have compiled experimental measurements of $t_{rup}$, and further developed a continuous viscosity model that successfully predicts the healthy-eye $t_{rup}$. In their model, the mucus layer is represented by a high-viscosity layer atop the cornea. In our model, $\beta_1$ should be such that the negative slip amounts to the same degree of hinderance to tear-film breakup. Figure 3 compares the predictions of the current model and that of [12]. The current model with $\beta_1 = -5.5$ nm predicts $t_{rup} = 119$ s, in good agreement with experimental data and the previous model prediction of $t_{rup} = 117$ s [12]. This will be used as a baseline value for the rest of the paper.

D. Model parameters

The dimensional parameters involved in the model are summarized in Table I. To facilitate illustration of the highly elongated domain, we adopt two characteristic lengths $L$ and $H$, and scale the longitudinal lengths by $L$ and the thicknesses by $H$. Following Braun et al. [19], we define the characteristic length $L$ by balancing the viscous and capillary forces. Based on estimations by [19] using parameters pertinent to human tear films, we take $L = 0.5$ mm. Following Dey et al. [12], we choose the length of the computational domain $\Lambda$ to be the wave-
length of the fastest linear mode. In the current model, this choice is complicated by the fact that the heterogeneous mucin loss on the substrate alters the growth rate of the linear instability, and the fastest mode cannot be easily determined \textit{a priori}. To be definite, however, we have chosen $\Lambda$ corresponding to the homogeneous and intact substrate with full mucin coverage, with $A_2$ and $\beta_\ast$. The consequences of this choice in tear-film thinning and breakup will be discussed in Section III C. Finally, the Hamaker constant $A_2$ is not listed in the table. We will take $A_r = A_2/A_1 = 2$ as the baseline value, and vary that ratio in the range of 1 to 4.

To make the system of equations and boundary conditions dimensionless, we need two more characteristic quantities besides the length $L$. We define the characteristic velocity and pressure using the van der Waals driving force:

$$V = \frac{A_1}{6\pi \mu HL}, \quad P = \frac{A_1}{6\pi H^3}. \quad (8)$$

Consequently, the characteristic time is $T = L/V$. Then non-dimensionalization yields the following dimensionless variables (the asterisks will be omitted for brevity in presenting results):

$$x^* = \frac{x}{L}, \quad y^* = \frac{y}{H}, \quad h^* = \frac{h}{H}, \quad u^* = \frac{u}{V}, \quad v^* = \frac{L}{HV},$$
$$t^* = \frac{t}{T}, \quad p^* = \frac{p}{F}, \quad \Gamma^* = \frac{\Gamma}{\Gamma_m}, \quad \sigma^* = \frac{\sigma}{\sigma_m},$$

as well as the following dimensionless groups:

- Aspect ratio: $\epsilon = H/L,$
- Domain length: $\lambda = \Lambda/L,$
- Slip lengths: $\beta_{1,2} = \beta_{1,2}/H,$
- Fraction of MAM-depleted substrate: $f = l_2/\Lambda,$
- MAM transition width: $d = \delta/\Lambda,$
- Ratio of Hamaker constants: $A_r = A_2/A_1,$
- Reynolds number: $Re = \rho V L/\mu,$
- Peclet number: $Pe_s = VL/D_s,$
- Marangoni number: $M = S/D_s,$
- Capillary number: $Ca = \mu V/\sigma_m.$

Using the parameter values in Table I, we determine the baseline values of the dimensionless groups: $\epsilon = 10^{-3},$ $\lambda = 1.61, \beta_{1,2} = 0.011, \beta_2 = 0, f = 1/3, A_r = 2, Re = 10^{-6}, Pe_s = 0.02, M = 1.67 \times 10^{-6}$ and $Ca = 3.11 \times 10^{-8}.$ For brevity, we will drop the bar over the dimensionless slip lengths $\beta_{1,2}$ in the following. Note that the lengths of the MAM-depleted and MAM-covered regions are constrained by $2l_1 + l_2 = \Lambda,$ and we have chosen $f = 1/3$ as the baseline. On the basis of the above, we will vary the parameters $A_r, \beta_1$ and $f$ to study the consequences of progressive mucin loss in tear-film breakup. In presenting the results, we will mostly use dimensionless variables. The only exception is the rupture time $t_{rup},$ for which we also give the dimensional values in certain cases to inform intuition and facilitate comparison with measured values. Note that with the parameter values of Table I, the characteristic time is $T = 357$ s. Initially, the interface is flat at $h_0 = 1,$ with a uniform surfactant concentration $\Gamma_0 = 0.5.$ The amplitude of the initial sinusoidal perturbation to the interface is $0.01h_0.$

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Description</th>
<th>Value and source</th>
</tr>
</thead>
<tbody>
<tr>
<td>$L$</td>
<td>Characteristic length</td>
<td>$5 \times 10^{-4}$ m [19]</td>
</tr>
<tr>
<td>$H$</td>
<td>Initial film thickness</td>
<td>$5 \times 10^{-7}$ m [19]</td>
</tr>
<tr>
<td>$\lambda$</td>
<td>Domain length</td>
<td>$8.05 \times 10^{-4}$ m [12]</td>
</tr>
<tr>
<td>$\beta_1$</td>
<td>Slip length on glyocalyx</td>
<td>$-5.5 \times 10^{-9}$ m (this paper)</td>
</tr>
<tr>
<td>$\beta_2$</td>
<td>Slip length on exposed cornea</td>
<td>0 (this paper)</td>
</tr>
<tr>
<td>$\delta$</td>
<td>Transition layer thickness in Eq. (1)</td>
<td>$8.05 \times 10^{-6}$ m (this paper)</td>
</tr>
<tr>
<td>$\rho$</td>
<td>Density of tear film</td>
<td>$1.3 \times 10^3$ kg m$^{-3}$ [51]</td>
</tr>
<tr>
<td>$\mu$</td>
<td>Characteristic viscosity of tear film</td>
<td>$1.3 \times 10^{-3}$ Pa s [39]</td>
</tr>
<tr>
<td>$\sigma_m$</td>
<td>Maximum interfacial tension</td>
<td>$4.5 \times 10^{-2}$ N m$^{-1}$ [52]</td>
</tr>
<tr>
<td>$S$</td>
<td>Maximum spreading pressure</td>
<td>$7.5 \times 10^{-8}$ N m$^{-1}$ [27]</td>
</tr>
<tr>
<td>$\Gamma_m$</td>
<td>Maximum surfactant concentration</td>
<td>$4 \times 10^{-7}$ mol m$^{-2}$ [53]</td>
</tr>
<tr>
<td>$A_1$</td>
<td>Hamaker constant on glyocalyx</td>
<td>$6.6 \times 10^{-18}$ J [30]</td>
</tr>
<tr>
<td>$D_s$</td>
<td>Surface diffusivity of lipids</td>
<td>$3 \times 10^{-8}$ m$^2$ s$^{-1}$ [53]</td>
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We perform numerical simulations of the tear-film breakup by solving the Navier-Stokes equations in a two-dimensional domain (Fig. 1) using the finite-element package COMSOL Multiphysics®. The moving and deforming interface is tracked by using the arbitrary Lagrangian-Eulerian (ALE) approach with a moving mesh and adaptive mesh refinement. The algorithm refines the mesh near sharp gradients and the moving boundary at every time step. The initial conditions are prescribed as follows:

\[ h(x, 0) = 1 + 0.01 \cos\left(\frac{2\pi x}{\lambda}\right), \quad (9) \]

\[ \Gamma(x, 0) = \frac{1}{2} + \Delta \Gamma \cos\left(\frac{2\pi x}{\lambda}\right), \quad (10) \]

where the perturbation amplitude in the lipid concentration field, \( \Delta \Gamma \) is chosen as per linear stability analysis [12, 45]. The simulation ends when the tip of the interfacial cusp becomes so pointed that further mesh refinement fails. This typically happens right before the tip reaches the substrate. Thus the uncertainty in the moment of rupture is small and does not affect the \( t_{rup} \) value significantly.

In Dey et al. [12], we have already validated several key components of the model, including surfactant transport, Marangoni flows and the conjoining pressure due to van der Waals attraction. The new element in the current implementation is the ALE treatment of interfacial motion. As a validation of the ALE algorithm, we have simulated the Rayleigh-Plateau instability during breakup of a cylindrical filament of a Newtonian fluid, and compared the results with those of Ashgriz and Mashayek [54]. Our results are in very good agreement with the published data, and Fig. 4 presents one such comparison for two solutions obtained at two levels of mesh resolution. Note that our solution hardly changes with mesh refinement. The subsequent solutions are at spatial resolutions equal to or finer than that of the \( N = 200 \) solution presented here.

III. RESULTS AND DISCUSSION

In our heterogeneous mucin model, the loss of MAM on the cornea surface is reflected by elevated van der Waals attraction with the tear-air interface, as well as a loss of negative slip on the substrate. In subsection II C, we have tuned the negative slip \( \beta_1 \) to match the \( t_{rup} \) of the healthy tear film. From this starting point, the model predicts progressive shortening of \( t_{rup} \) as more of the corneal surface loses mucin coverage. It turns out that the increased van der Waals force is the main cause of the premature tear-film rupture; the loss of negative slip only plays a minor role. We also compare the predictions of the current model with those of the continuous viscosity model [12] and with experimental data for diseased eyes.

A. General characteristics of tear-film rupture

Figure 5 presents the general flow characteristics during tear-film rupture predicted by the heterogeneous mucin model using the baseline parameters of Table I. For this case, the MAM-depleted region occupies a fraction of \( f = 1/3 \) of the substrate and sits at its center, with a MAM-covered region of equal length on either side. In the initial stage of the simulation, the most salient feature is the appearance of two interfacial slopes atop the internal boundaries between the MAM-depleted and MAM-covered regions (Fig. 5a). The stronger van der Waals attraction inside drives a flow downward, which then crosses the internal boundaries further down and turns upward just outside the internal boundaries, creating a strongly rotational flow. Because of the large gradient of the conjoining pressure across the internal boundaries, the interfacial distortion is highly localized initially, with a dip inside and a hump outside. This flow creates two local troughs just inside the boundaries that are separated by a peak at the center of the domain. The nature of the van der Waals attraction is such that it drives the dip further down while raises the hump further up, exacerbating the interfacial distortion in time (Fig. 5b).

In time, viscous dissipation broadens the flowing regions and moves the two troughs toward each other (Fig. 5b). The downward flow eventually eliminates the central peak and merges the two troughs into a single one (Fig. 5c). Thanks to this change in flow pattern, the central trough is now narrower and well within the two internal boundaries that mark the MAM-depleted portion of the substrate.

From this point onward, the rupture proceeds much as
FIG. 5: Tear-film rupture with the central portion of the substrate between the vertical dotted lines being cleared of mucin $(f = 1/3, A_r = 2, \beta_1 = -0.011, \beta_2 = 0, d = 0.01)$. The greyscale contours and scale bar show the magnitude of velocity with streamlines superimposed, while the vectors represent velocities with logarithmic magnitude as its length. The frames are at times (a) $t = 0.36$ s, (b) $t = 2.57$ s, (c) $t = 4.65$ s and (d) $t = 5.21$ s, shortly before the tear film ruptures at $t = 5.35$ s.

in earlier models with a homogeneous substrate [12, 45], with a single trough that is driven further down by the van der Waals force against the restoring effects of surface tension and Marangoni stress. Figure 5(d) shows the film profile and velocity contour just before the film ruptures. The negative slip-length $\beta_1$ used in the model induces fine vortical structures at the bottom substrate. The wavy streamlines near the bottom in Figs. 5(c) and 5(d) are a manifestation of these fine vortices.

As the film rupture is driven by the gradient of the conjoining pressure across the boundaries between the mucin-intact and mucin-depleted regions, one naturally wonders how the process is affected by the sharpness of the transition (see Eq. 1). Figure 6 compares the solutions for several values of the transition width from $d = 0$ (sharp transition) to $d = 0.1$ (more gradual transition). The film profile changes slightly with $d$, and clearly converges to a “sharp-transition” limit as $d$ falls below $10^{-2}$ (Fig. 6a). The rupture time $T_{rup}$ increases moderately with $d$, as may be expected, but exhibits apparent limits at both bounds of the $d$ range tested (Fig. 6b). For lack of experimental data on the transition width, we have chosen $d = 10^{-2}$ (or 8.05 µm), close to the sharp-transition limit, as our baseline value.

B. Effects of the van der Waals force and the no-slip condition

We explore separately the effects of an enhanced Hamaker constant and the no-slip condition on breakup of diseased tear films. In the first set of simulations, therefore, we vary $A_2$ while keeping $\beta_2 = \beta_1 = -0.011$, i.e. maintaining the negative-slip boundary condition even in the MAM-depleted area. Besides, we keep the relative size of the MAM-depleted region $f$ and the Hamaker constant $A_1$ over the intact MAM fixed.

Figure 7 shows the effect of gradually increasing the Hamaker constant $A_2$ over the MAM-depleted region. The film thins progressively faster with increasing
$A_2$, and the rupture time $t_{rup}$ decreases monotonically (Fig. 7(a)). This trend is more explicitly shown by the dashed curve of Fig. 7(b). As $A_r$ increases from 1 to 2, $t_{rup}$ shortens from 119 s to 7.6 s, by a factor of nearly 16. Increasing $A_r$ to 4 reduces $t_{rup}$ further to 1.9 s. The latter can be considered instantaneous rupture as reported in the experimental literature [15, 17]. The decrease of $t_{rup}$ with increasing $A_r$ is initially very rapid, and then becomes gentler for larger $A_r$.

A secondary feature is the shape of the thinning curves in Fig. 7(a). With mucin heterogeneity ($A_r > 1$), the $h_c(t)$ curves change concavity in time. In other words, the thinning speed $dh_c/dt$ is non-monotonic; it first increases, then decreases before increasing again toward rupture. In contrast, the homogeneous mucin case ($A_r = 1$) has a monotonically increasing thinning speed. The temporary slow-down in thinning is probably related to the temporary central hump between the troughs of Fig. 5(b), and is thus a signature of the spatial heterogeneity of the conjoining pressure.

The solid curve in Fig. 7(b) illustrates the effect of changing the boundary condition in the MAM-depleted region from the negative slip ($\beta_2 = \beta_1$) to no slip ($\beta_2 = 0$). This brings about an additional small decrease in the rupture time $t_{rup}$. Evidently, the loss of negative slip allows faster liquid flow near the substrate, which in turn facilitates the rupture. This effect is more pronounced, in relative terms, for larger values of $A_r$. But overall it remains much weaker than that of changing $A_r$. Therefore, between the enhanced van der Waals attraction and the loss of negative slip on the substrate, the former plays the primary role in shortening the breakup time for diseased tear films.

### C. Progressive loss of mucin coverage

Clinical evidence points to mucin loss in eye diseases such as bacterial infection and the dry eye syndrome [7, 9, 10]. In our heterogeneous mucin model, we can represent progressive mucin loss by the expansion of the mucin-depleted area, parametrized by the fraction $f$. In subsection III B, we have investigated the roles of an elevated Hamaker constant and loss of negative slip in the MAM-depleted zone at a fixed $f = 1/3$, and come to the conclusion that a higher $A_2$ markedly shortens the
rupture time $t_{rup}$, while $\beta_2 = 0$ plays only a minor role. Naturally one wonders if this conclusion holds for larger $f$ values, when more of the substrate features the no-slip condition. Fig. 8 shows the effect of increasing $f$. When we only impose the no-slip condition ($\beta_2 = 0$) on the MAM-depleted area without raising the Hamaker constant (dashed curve with $A_r = 1$), we find a relatively mild decline in the rupture time $t_{rup}$ with increasing $f$. Compared with the case of no mucin loss ($f = 0$), loss of the negative slip over a fraction of the substrate ($f = 1/3$) shortens $t_{rup}$ by a mere 5%, and its complete loss over the entire substrate ($f = 1$) causes a reduction of about 11%. This supports the previous conclusion: the loss of negative slip plays a minor role in hastening tear-film rupture even if it prevails over the entirety of the corneal surface.

Now we come to the more realistic case of an expanding MAM-free zone that features both no slip and an elevated $A_2$. Recall that the domain length $\lambda$ is fixed at the wavelength $\lambda_1$ of the fastest linear mode corresponding to a spatially uniform substrate with Hamaker constant $A_1$. This protocol has been followed in the simulations presented so far, and the corresponding result is represented by the curve marked $\lambda = \lambda_1$ in Fig. 8. The curve reveals a steep decrease in the rupture time $t_{rup}$ as $f$ increases from 0 to 1/3, at which point $t_{rup}$ has shortened by a factor of 22, from 119 s to 5.4 s. This is consistent with the earlier observations in Fig. 7. Curiously, as $f$ increases further, $t_{rup}$ begins to rise gradually, reaching $t_{rup} = 17$ s at $f = 1$. This counterintuitive result is mostly due to the fixed domain length $\lambda = \lambda_1$. Without mucin loss ($f = 0$), $\lambda_1$ is the wavelength of the fastest linear mode, and nonlinear simulation of breakup in such a domain gives a proper estimation of $t_{rup}$. With mucin loss and elevated van der Waals attraction, $\lambda_1$ may no longer represent the fastest mode. Then the nonlinear simulation will yield an overestimation of $t_{rup}$.

To test this idea, we have carried out a linear stability analysis by using an analytical result of Zhang et al. [45] (Eq. 20 therein), for a homogeneous substrate with $\beta_1 = -0.011$ and other parameters at baseline values. The dispersion relation is depicted in Fig. 9(a) for three values of the Hamaker constant. The tear film becomes progressively unstable with increasing $A_r$ and the fastest growing wavenumber increases with $A_r$. The corresponding fastest wavelength $\lambda_m$ declines monotonically with increasing $A_r$ (Fig. 9b). This suggests that in our Fig. 8, as $A_r$ is raised to $A_2$ over increasing portions of the substrate, the wavelength of the fastest linear model should become progressively shorter than $\lambda_1$. Consequently, the breakup times predicted by the simulations are longer than the true values.

Unfortunately, it is impossible to know a priori what the proper wavelength or domain length should be for our heterogeneous substrate with distinct $A_r$ values. As a rough estimation, we define an “effective Hamaker constant” $A_e = (1 - f)A_1 + fA_2$, and carry out a series of simulations in which $\lambda$ is varied according to the fastest wavelength $\lambda_e$ for a homogeneous substrate with $A = A_e$. The results are represented by the curve marked by $\lambda = \lambda_e$ in Fig. 8. Note first that the two curves coincide at $f = 0$ as expected. For $f > 0$, the “effectively fastest mode” always outgrows the $\lambda_1$ mode, with a shorter $t_{rup}$. Therefore, it gives a better approximation to the fastest mode that prevails in reality. In particular, the rise of $t_{rup}$ with increasing $f$ beyond $f = 1/3$ has largely disappeared.

The remaining slight increase is attributable to the complexities in the tear-film flow due to heterogeneity of the substrate. These include the rapid rotational flow driven by differing $A$ across the internal boundaries and the resultant interfacial distortions (Fig. 5a), the appearance and subsequent merging of the dual valleys (Fig. 5b,c), and the non-monotonic evolution of the thinning speed in Fig. 7(a). For one, the two internal boundaries are farther apart at larger $f$ values, and the two valleys need to traverse longer distances before merging. This may have contributed to the slight increase in breakup time with increasing $f$.

### D. Comparison with prior studies

It is interesting to compare the predictions of the heterogeneous mucin model with experimental data of shortened $t_{rup}$, and with prediction of the continuous viscosity model [12]. As the mucin loss extends to about 1/3 of the corneal surface, our model predicts a dramatic shortening of the rupture time from 119 s to 3.3 s (Fig. 8, the $\lambda = \lambda_e$ curve). This prediction is consistent with premature tear-film breakup measured by clinical studies,
stability. But the imperfection of the “effectively fastest mode” implies at most a small over-prediction of $t_{rup}$, and does not affect the main model prediction that mucin loss greatly reduces breakup time of tear films. Taken together, the above comparisons show the advantage of the heterogeneous mucin model over the continuous viscosity model, as well as its potential in explaining the shortening of tear-film rupture time in various eye diseases.

Out of curiosity, we have also explored a “homogeneous mucin loss” model, where we impose an elevated $A_2$ over the entire domain along with the no-slip boundary condition $\beta_2 = 0$. With increasing $A_2$, we shorten the domain length $\lambda$ according to the fastest mode of Fig. 9. Results show that the homogeneous model predicts a longer $t_{rup}$ than the heterogeneous mucin model, and under-predicts the shortening of $t_{rup}$ observed in diseased eyes. For example, the homogeneous model predicts $t_{rup} = 5.8$ s at $A_2 = 2$, while the heterogeneous mucin model predicts $t_{rup} = 3.3$ s with $f = 1/3$ and $\lambda = \lambda_c$ (Fig. 8). This difference may have the same origin as that between $f = 1/3$ and 1 on the $\lambda = \lambda_c$ curve in Fig. 8, and can be attributed to the flow features due to the spatially heterogeneous mucin distribution. The closer agreement between the heterogeneous mucin model and experimental data underscores the observations of localized and nonuniform loss of mucin in diseased eyes [2, 5, 30].

IV. CONCLUSION

The heterogeneous mucin model reported in this paper is motivated by experimental observations of the loss of membrane-associated mucin polymers (MAMs) in various eye diseases and the concomitant premature breakup of the tear film. It has been hypothesized that the loss of MAMs is the cause for the premature tear-film rupture. We have tested this hypothesis by examining two mechanisms that may contribute to shortening of the tear-film breakup time: an elevated van der Waals attraction over corneal surfaces that have lost its MAMs, and a change of the boundary condition from an effective negative slip on MAM-covered surfaces to no slip on MAM-depleted surfaces. Using finite-element simulations of film rupture, we probed the hydrodynamic consequences of these two factors by assuming a heterogeneous substrate made of MAM-intact and MAM-depleted areas. Within the parameter ranges tested herein, the model predicts the following results:

(a) Both an elevated van der Waals attraction and a loss of negative slip contribute to a shortening of the $t_{rup}$. The former works by strengthening the conjoining pressure that drives the interfacial instability and the latter by facilitating fluid flow near the corneal surface.

(b) The elevated van der Waals force has a much stronger effect; doubling the Hamaker constant over a fraction of the corneal surface can reduce
The rupture time $t_{rup}$ continues to decrease with progressive expansion of the MAM-free area to about 1/3 of the substrate. The shortened $t_{rup}$ predicted by the model is consistent with clinical data for diseased eyes. With further expansion, the breakup time remains largely unchanged.

The heterogeneous mucin model is superior to the earlier continuous viscosity model as the latter severely under-predicts the shortening in $t_{rup}$ with mucin loss. Thus, a more realistic account of mucin loss on the corneal surface—via changes in wettability and slip condition as opposed to an increase in effective diffusivity—has produced a better model.

The above notwithstanding, we should point out the weaknesses of the current study. The most serious one is perhaps the uncertainty in evaluating the increase in Hamaker constant that accompanies the loss of MAMs. Currently we rely on theoretical estimations of much simplified systems that suggest a rough range of the increase. A direct measurement in the context of the human tear film and cornea will greatly strengthen the basis for our heterogeneous mucin model and allow it to make more quantitative predictions. Second, we have found no experimental data regarding the degree of mucin loss in various eye diseases. In our model, we have represented the progression of mucin loss by the expansion of the mucin-depleted region as a fraction of the entire substrate. This is consistent with the emerging picture in the experimental literature of spatially localized areas of mucin loss or alteration. But no quantification of the loss seems to have been done. Third, the computation is done in a two-dimensional domain that is periodic in the longitudinal direction with a prescribed length $\lambda$. This $\lambda$ corresponds to the wavelength of the most unstable linear mode for a homogeneous corneal surface with a uniform Hamaker constant. As we increase the area of the MAM-depleted region with an elevated Hamaker constant, the fastest linear mode should take on a shorter wavelength. Since this cannot be prescribed \textit{a priori}, our computation cannot faithfully capture the most rapid rupture starting from the appropriate initial perturbation. This is a well-recognized challenge for numerically simulating complex instability problems for which the linear modes are not known. Fourth, the two-dimensionality of the model precludes 3D variations, especially variations in the plane parallel to the corneal surface. Such 3D patterns have been documented by imaging \cite{55} and explored to a limited extent computationally \cite{44, 56}. How they affect the tear-film rupture time is an open question. Finally, our model has neglected a host of potentially important factors in tear-film breakup, including non-Newtonian rheology of the tear, evaporation, osmotic flux between the tear film and the cornea, non-uniformity in the lipid layer, and gravity-driven drainage. Such simplifications are perhaps justifiable as we try to focus on the effect of MAM loss on the corneal surface. But to make more quantitative predictions, a more general model should account for such factors with realistic parameters.

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