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A mechanochemical model of striae distensae

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Abstract

Striae distensae, otherwise known as stretch marks, are common skin lesions found in a variety of clinical settings. They occur frequently during adolescence or pregnancy where there is rapid tissue expansion and in clinical situations associated with corticosteroid excess. heralding their onset is the appearance of parallel inflammatory streaks aligned perpendicular to the direction of skin tension. Despite a considerable amount of investigative research, the pathogenesis of striae remains obscure. The interpretation of histologic samples - the major investigative tool - demonstrates an association between dermal lymphocytic inflammation, elastolysis, and a scarring response. Yet the primary causal factor in their aetiology is mechanical; either skin stretching due to underlying tissue expansion or, less frequently, a compromised dermis affected by normal loads. In this paper, we investigate the pathogenesis of striae by addressing the coupling between mechanical forces and dermal pathology. We develop a mathematical model that incorporates the mechanical properties of cutaneous fibroblasts and dermal extracellular matrix. By using linear stability analysis and numerical simulations of our governing nonlinear equations, we show that this quantitative approach may provide a realistic framework that may account for the initiating events.

Highlights:

- We present a mechanochemical model of the onset of striae distensae
- Uniformity of cutaneous ECM may be rendered unstable by variations in parameters

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- Instability strongly affected by changes in skin stiffness and cell contractile force
- Analysis confirmed using numerical simulations of the full nonlinear model
- Able to predict postulated precursors of striae for biologically realistic parameters

Keywords: stretch marks, mathematical modelling, numerical simulation

1. Introduction

Striae distensae, commonly known as stretch marks, are benign skin lesions associated with considerable cosmetic morbidity. heralding their onset is the appearance of parallel inflammatory streaks aligned perpendicular to the direction of skin tension. The evolution of striae is characterised by at least two phases: an initial inflammatory phase known as striae rubra and a later, chronic phase known as striae alba [1]. Striae may occur in a wide variety of clinical settings but most commonly develop initially in either adolescence [2] or pregnancy [3]. Striae may also occur in conditions where the dermis is abnormal: Cushing's syndrome [4], prolonged application of topical steroids [5], or Marfan's syndrome [4] are examples. Finally, striae may develop in association with changes to body habitus such as weight loss [6], cachexia [7], obesity [8], or body-building.

Despite their ubiquity and considerable investigation into their origins, the pathogenesis of striae distensae remains unknown. Genetic factors are likely to be important since striae have been observed in monozygotic twins [9]. Much emphasis has been placed on the effects of skin stretching in the pathogenesis of striae [10] since the lesions are found to be aligned perpendicular to the direction of skin tension. Some investigators have suggested that mechanical rupture of dermal components is an important initiating event. However, there is some debate about the relative importance of skin stretching in the aetiology of striae: one group could not find any relationship between striae and the increase in abdominal girth among pregnant females [11] and it has been noted that striae are rare over the extensor surfaces of joints (regions of skin over a joint that is stretched when the joint is flexed) where the skin is subject to physiologic stretching [12]. Less controversy exists regarding the possible role of glucocorticoids in the pathogenesis of striae. This is largely due to the known associations between alterations to hormonal status observed in pregnancy, weight changes, and adolescence on one hand and the more obvious effects of hormonal changes observed in Cushing's syndrome and topical steroid application on the other. In addition, the catabolic effect of both **adrenocorticotrophic hormone (ACTH)** and cortisol are well known. These hormones may modulate fibroblast activity directly leading to reduced mucopolysaccharide secretion, possible changes to elastic fibres, and reduced collagen via either reduced production or increased collagenase secretion or both. Finally, increased levels of steroid hormones and their metabolites have been found in patients exhibiting striae [13].

From the pathologic perspective, the earliest changes are subclinical and are only detectable by electron microscopy. These changes involve mast cell degranulation (the release histamine along with other molecules from granules in the mast cell's cytoplasm into the extracellular space) and the presence of activated macrophages in association with mid-dermal elastolysis [14]. While mast cell activation has been

32 reported in association with elastolysis in actinically affected skin (skin that has been exposed to chemically
33 active rays of the electromagnetic spectrum such as uv light and x-rays), anetoderma (laxity of the skin due
34 to loss of dermal elasticity) has been reported in skin affected by mastocytosis. These findings support the
35 concept that the release of enzymes, possibly elastases (enzymes that are capable of degrading the elastin
36 molecule within the dermis), from mast cells plays a very early and important role in the pathogenesis of
37 striae [14]. When lesions initially become visible, collagen bundles begin to show structural alterations,
38 fibroblasts become prominent, and mast cells are absent [14].

39 On light microscopy, the earliest changes in striae rubra involve dermal oedema, perivascular lymphocyte
40 cuffing (the appearance of lymphocytes in the surrounding small blood vessels within the dermis), and an
41 associated increase in the glycosaminoglycan content of the dermis [15]. Examination of early lesions shows
42 fine elastic fibres predominating throughout the dermis in association with thick and tortuous fibres toward
43 the periphery [16]. An integral component of elastic fibres, fibrillin microfibrils, are found to be reduced in
44 striae rubra [17]. In contrast to early inflammatory lesions, striae alba are characterised by epidermal atrophy,
45 loss of appendages, and a densely packed region of thin eosinophilic collagen bundles aligned horizontal to
46 the surface. Later stage lesions are thus indistinguishable, from the perspective of light microscopy, from a
47 dermal scar.

48 A number of studies suggest that fibroblasts play a key role in the pathogenesis of striae. Compared
49 with normal fibroblasts, expression of fibronectin and both type I and III procollagen were found to be signif-
50 icantly reduced in fibroblasts from striae, suggesting that there exist fundamental aberrations of fibroblast
51 metabolism in striae distensae [18]. From a bio-mechanical perspective, ex-vivo fibroblasts from patients
52 with early striae distensae were found to exhibit high levels of alpha-smooth muscle actin and were able to
53 generate higher contractile forces in comparison with fibroblasts from later stage striae [19].

54 Taken together, the foregoing discussion suggests at least two major factors play important roles in
55 the aetiology of striae distensae: mechanical stretching of the skin and pre-existing dermal pathology. The
56 relative effects of these factors are unknown. For example, it is unknown to what extent steroid hormones in
57 pregnancy or adolescence may pre-condition the skin such that it may be predisposed to developing striae
58 when subjected to stretching.

59 In this paper we develop a mathematical model that attempts to capture early changes in striae distensae
60 development. We are encouraged by the success of similar models used to describe wound healing [20] and
61 earlier contact guidance models of striae by Murray [21] and Hariharan [22] and we are motivated by the
62 ability of mathematical models to incorporate the postulated relevant elements of early striae development
63 including fibroblast contractility, fibroblast motility and remodelling of the extra-cellular matrix. Our model
64 allows us to quantify the degree of mechanical stretching (given by a single parameter) and dermal stiffness
65 (given by an independent parameter) so that we are able to explore the relevant contributions of skin
66 stretching and glucocorticoid-affected skin [23, 24]. Finally, as a model of pattern formation in the skin, we
67 are able to investigate how microscopic events may lead to macroscopic patterns.

68 The remainder of this paper is organised as follows: Section 2 describes the derivation of our model and
69 a non-dimensionalisation of our governing equations. In Section 3, we perform a linear stability analysis
70 and investigate mode selection. Section 4 describes our numerical results of the full nonlinear model. We

71 conclude this paper in Section 5 with a discussion of our results and the implications for pathogenesis.

72 2. Model description

73 Our model is based on the simple assumption that in the pre-clinical phase of striae development there
74 exists spatial inhomogeneity in the density of one or more constituents of the dermis. We thus focus on its
75 two most important components: fibroblasts and the extracellular matrix (ECM). Fibroblasts are spindle-
76 shaped cells embedded within the ECM; they play an essential role in dermal homeostasis, wound healing,
77 and recently have been shown to express a Hox code that accounts for the regional specificity of the epidermal
78 phenotype [25]. Proteoglycans and mucopolysaccharides constitute the ECM. While collagen gives the skin
79 its tensile strength, the proteoglycans and mucopolysaccharides are gel-like substances that trap water, thus
80 facilitating molecular diffusion and cell transport.

81 Since striae are frequently observed to align in a direction perpendicular to the direction of skin tension,
82 they are found to develop as parallel inflammatory streaks in the skin. Hence, without loss of generality,
83 we can reduce a potential two-dimensional problem to a one-dimensional problem since the patterning is
84 translationally invariant in the direction perpendicular to the direction of skin tension. We thus consider a
85 one-dimensional model, defined on a periodic domain, similar to the model developed by Oster et al. [26]
86 and modified by Vaughan, Jr. et al. [27] where the skin is treated as a visco-elastic medium. The model
87 derived below and the model in Vaughan, Jr. et al. [27] generalise the model discussed in Oster et al. [26]
88 and Murray [21] by keeping the inertial terms and writing the governing equations in the material frame of
89 reference. This derivation keeps the nonlinear terms that arise from transformation from spatial to material
90 frames of reference in the spatial derivatives, which increases the range of parameters where the solution
91 evolves to a bounded steady state [27].

92 The model consists of conservation equations for the fibroblast cell density, \hat{c} , and the ECM density,
93 $\hat{\rho}$, in a deformed frame of reference coupled through a force balance equation governing the mechanical
94 interaction of the fibroblasts with the ECM, which is defined in the undeformed (reference) frame. In this
95 formulation, we will transform the equations governing the cell and ECM densities from the deformed frame
96 to the reference frame, which will simplify the methods used to solve this model numerically. Note that
97 variables with a hat denote a variable in the deformed frame of reference and variables without a hat are
98 defined in the reference frame.

99 The stress tensor, σ , satisfies the force balance equation in the reference frame,

$$\rho_0 \frac{\partial^2 u}{\partial t^2} = \frac{\partial}{\partial x} (\sigma + \tau(\hat{c}, \hat{\rho})) + \rho_0 F, \quad (1)$$

100 where u is the material displacement, $\tau(\hat{c}, \hat{\rho})$ is the traction due to cell-ECM interactions and depends on
101 the cell and ECM densities in the deformed frame of reference, ρ_0 is the ECM density in the reference frame,
102 and F is an external body force. We follow the Oster-Murray-Harris model [26] and treat the ECM as a

103 linear, isotropic, visco-elastic material. Hence, the stress tensor in one dimension is

$$\sigma = A \frac{\partial^2 u}{\partial t \partial x} + B \frac{\partial u}{\partial x}. \quad (2)$$

104 Here, $A = \mu_1 + \mu_2$, where μ_1 and μ_2 are the shear and bulk viscosities of the ECM, respectively, and

$$B = E \frac{(1 - \nu)}{(1 + \nu)(1 - 2\nu)},$$

105 where E is the Young's modulus and ν is the Poisson's ratio of the ECM. It is assumed that the ECM
 106 material is attached to the subcutaneous fascia by fibrous bands that resist the lateral displacement of the
 107 overlying dermis. We model this attachment as a linear spring and the body force in the force balance
 108 equation, (1), is

$$F = -su, \quad (3)$$

109 where s is a positive spring constant.

110 We model the traction exerted by the cell-matrix interactions in the deformed frame of reference as

$$\tau(\hat{c}, \hat{\rho}) = \tau_0 \frac{\hat{c}}{1 + \lambda \hat{c}^2} \left(\hat{\rho} + \beta_0 \frac{\partial^2 \hat{\rho}}{\partial \hat{x}^2} \right), \quad (4)$$

111 where τ_0 is the traction strength, λ is a constant that accounts, in a phenomenological way, for contact
 112 inhibition, β_0 is the strength of the long-range traction that arises from the fibrous nature of the ECM,
 113 which can extend the range of the traction force exerted by the fibroblasts, and $\partial/\partial \hat{x}$ is the spatial derivative
 114 in the deformed frame of reference.

115 We assume that there is no production or degradation of the ECM, so we can relate the ECM density
 116 in the deformed frame to the ECM density in the reference frame using the relation $\rho_0 = J\hat{\rho}$, where $J =$
 117 $1 + \partial u/\partial x$ is the Jacobian of the deformation gradient and ρ_0 is a constant. Likewise, we transform the
 118 cell density from the deformed frame to the reference frame using the same relation, $c = J\hat{c}$. Here, c is not
 119 assumed to be constant since we will allow for the movement of cells by diffusion.

120 We transform the derivatives in the long-range traction force into the reference frame by taking the
 121 derivative of the definition of the displacement, $u(x, t) = \hat{x}(x, t) - x$, with respect to the spatial coordinate
 122 in the reference frame, to obtain

$$\frac{\partial}{\partial \hat{x}} = \frac{1}{1 + \partial u/\partial x} \frac{\partial}{\partial x}. \quad (5)$$

123 Hence, the traction force in the reference frame is

$$\tau(c, \partial u/\partial x) = \tau_0 \frac{c\rho_0}{(1 + \partial u/\partial x)^2 + \lambda c^2} \left(1 + \beta_0 \frac{\partial}{\partial x} \left(\frac{1}{1 + \partial u/\partial x} \frac{\partial}{\partial x} \left(\frac{1}{1 + \partial u/\partial x} \right) \right) \right). \quad (6)$$

124 See Appendix A for the details of the derivation of this term.

125

Inserting (2) and (6) into (1), the resulting force balance equation in one dimension is

$$\rho_0 \frac{\partial^2 u}{\partial t^2} = \frac{\partial}{\partial x} \left[A \frac{\partial^2 u}{\partial x \partial t} + B \frac{\partial u}{\partial x} + \tau \left(c, \frac{\partial u}{\partial x} \right) \right] - s \rho_0 u = 0. \quad (7)$$

126

It is assumed that fibroblasts move randomly and are advected with the medium. Thus the governing equation for the cell density, \hat{c} , in the deformed frame of reference is

127

$$\frac{\partial \hat{c}}{\partial \hat{t}} + \frac{\partial}{\partial \hat{x}} (\hat{c}v) = D \frac{\partial^2 \hat{c}}{\partial \hat{x}^2}, \quad (8)$$

128

where $v = \partial u / \partial t$ is the velocity of the medium, D is the diffusion coefficient and \hat{t} is time in the deformed frame. Relating the cell density in the deformed configuration, \hat{c} , with the cell density in the initial configuration, c , using the relation $c = J^{-1} \hat{c}$ and the change in the spatial derivative, (5), and the change in the temporal derivatives due to the change in frame,

129

130

131

$$\frac{\partial}{\partial t} = \frac{\partial}{\partial \hat{t}} + v \frac{\partial}{\partial \hat{x}}, \quad (9)$$

132 we obtain

$$\frac{\partial c}{\partial t} = D \frac{\partial}{\partial x} \left[\frac{1}{1 + \partial u / \partial x} \frac{\partial}{\partial x} \left(\frac{c}{1 + \partial u / \partial x} \right) \right]. \quad (10)$$

133

Next, we non-dimensionalise equations (10) and (7) using the relations:

$$\begin{aligned} x^* &= \frac{x}{L}, & t^* &= \frac{t}{T}, & c^* &= \frac{c}{c_0}, & u^* &= \frac{u}{L}, \\ \lambda^* &= \lambda c_0^2, & \alpha^* &= \frac{1}{s T^2}, & a^* &= \frac{A}{L^2 T s \rho_0}, & b^* &= \frac{B}{L^2 s \rho_0}, \\ d^* &= \frac{D T}{L^2}, & \tau^* &= \frac{\tau_0 c_0}{L^2 s}, & \beta^* &= \frac{\beta_0}{L^2}, \end{aligned} \quad (11)$$

134

where L is the characteristic length scale and T is the characteristic time scale. The non-dimensionalised equations, after dropping the stars, are:

135

$$\frac{\partial c}{\partial t} = d \frac{\partial}{\partial x} \left[\frac{1}{1 + \partial u / \partial x} \frac{\partial}{\partial x} \left(\frac{c}{1 + \partial u / \partial x} \right) \right] \quad (12)$$

136 and

$$\alpha \frac{\partial^2 u}{\partial t^2} = a \frac{\partial^3 u}{\partial x^2 \partial t} + b \frac{\partial^2 u}{\partial x^2} + \tau \frac{\partial}{\partial x} \left[\frac{c}{(1 + \partial u / \partial x)^2 + \lambda c^2} \left(1 + \beta \frac{\partial}{\partial x} \left[\frac{1}{1 + \partial u / \partial x} \frac{\partial}{\partial x} \left(\frac{1}{1 + \partial u / \partial x} \right) \right] \right) \right] - u. \quad (13)$$

137

Note that we generalise the Oster-Murray-Harris model by keeping the inertial term, $\alpha \partial^2 u / \partial t^2$. Even though the α term is small for physically relevant parameters, the acceleration of the medium, $\partial^2 u / \partial t^2$, can be large in the numerical simulations and it would not be appropriate to neglect this term for all time.

139

140 The above equations for the dimensionless fibroblast density, c , and the material displacement, u , are
 141 coupled with appropriate boundary conditions. Since striae occur as parallel lines and are translationally
 142 invariant in the transverse direction, we assume in this paper that the solution is periodic in space and,
 143 hence, we enforce periodic boundary conditions on the ends of the domain.

144 3. Linear analysis

145 The linearized versions of equations (12) and (13) around the normalized steady state $c = 1$ and $u = 0$
 146 are

$$\frac{\partial c}{\partial t} = d \left(\frac{\partial^2 c}{\partial x^2} - \frac{\partial^3 u}{\partial x^3} \right) \quad (14)$$

147 and

$$\alpha \frac{\partial^2 u}{\partial t^2} = a \frac{\partial^3 u}{\partial x^2 \partial t} + \left(b - 2 \frac{\tau}{(1 + \lambda)^2} \right) \frac{\partial^2 u}{\partial x^2} + \frac{\tau(1 - \lambda)}{(1 + \lambda)^2} \frac{\partial c}{\partial x} - \frac{\tau\beta}{1 + \lambda} \frac{\partial^4 u}{\partial x^4} - u = 0. \quad (15)$$

148 By defining $\tau_\lambda = \tau / (1 + \lambda)^2$ and $\beta_\lambda = \beta(1 + \lambda)$ and looking for solutions of the form

$$\begin{bmatrix} c \\ u \end{bmatrix} = \mathbf{z} e^{\sigma t + i k x}, \quad (16)$$

149 where \mathbf{z} is the eigenvector, σ is the linear growth rate, and k is the spatial wavenumber, we require $|A| = 0$,
 150 for

$$A = \begin{bmatrix} \sigma + dk^2 & -idk^3 \\ -i\tau_\lambda(1 - \lambda)k & \alpha\sigma^2 + ak^2\sigma + B(k^2) \end{bmatrix}, \quad (17)$$

151 where

$$B(k^2) = \tau_\lambda \beta_\lambda k^4 + (b - 2\tau_\lambda)k^2 + 1. \quad (18)$$

152 The characteristic equation is

$$\alpha\sigma^3 + (a + \alpha d)k^2\sigma^2 + (B(k^2) + adk^4)\sigma + dk^2(B(k^2) + \tau(1 - \lambda)k^2) = 0. \quad (19)$$

153 If we ignore inertial forces ($\alpha = 0$) and cell diffusion ($d = 0$), we recover the characteristic equation for the
 154 basic Oster-Murray-Harris model.

155 If we take the ratio τ_λ/b as the bifurcation parameter, the real part of σ can become positive in three
 156 ways as τ_λ/b . Applying the Routh-Horwitz conditions to (19),

$$B(k^2) + adk^4 < 0,$$

$$d \neq 0 \text{ and } B(k^2) + \tau_\lambda(1 - \lambda)k^2 < 0,$$

(20)

or

$$\alpha dk^2(B(k^2) + \tau_\lambda(1 - \lambda)k^2) \geq (a + \alpha d)k^2(B(k^2) + adk^4)$$

157 are the conditions necessary for linear instability.

158 The first case, $B(k^2) + adk^4 < 0$, will occur when

$$\frac{\tau_\lambda}{b} > \frac{1}{2}, \quad \text{and} \quad [b - 2\tau_\lambda]^2 = 4(\tau_\lambda\beta_\lambda + ad). \quad (21)$$

159 The critical wavemode in this case is

$$k_c^2 = \sqrt{\frac{1}{\tau_\lambda\beta_\lambda + ad}}. \quad (22)$$

160 The presence of diffusion has a stabilizing effect in this case by increasing the critical value of τ_λ required
161 for a bifurcation from

$$\tau_\lambda = \frac{b}{2} + \frac{1}{2} \left(\beta + \sqrt{\beta(2b + \beta)} \right) \quad (23)$$

162 in the original Oster-Murray-Harris model to

$$\tau_\lambda = \frac{b}{2} + \frac{1}{2} \left(\beta + \sqrt{\beta(2b + \beta) + 4ad} \right). \quad (24)$$

163 The second case, $d \neq 0$ and $B(k^2) + \tau_\lambda(1 - \lambda)k^2 < 0$, will occur when

$$\frac{\tau_\lambda}{b} > \frac{1}{1 + \lambda} \quad \text{and} \quad [b - (1 + \lambda)\tau_\lambda]^2 = 4\tau_\lambda\beta_\lambda. \quad (25)$$

164 The critical wavemode is

$$k_c^2 = \sqrt{\frac{1}{\tau_\lambda\beta_\lambda}}. \quad (26)$$

165 Note that this case only arises if there is cell diffusion, $d \neq 0$.

166 The third case will occur when

$$\frac{\tau_\lambda}{b} > \frac{1}{2 + \frac{\alpha d}{a}(1 - \lambda)} \quad \text{and} \quad \left[b - \left(2 + \frac{\alpha d}{a}(1 - \lambda) \right) \tau_\lambda \right]^2 = 4(\tau_\lambda\beta_\lambda + d(a + \alpha d)). \quad (27)$$

167 The critical wavemode is

$$k_c^2 = \sqrt{\tau_\lambda\beta_\lambda + d(a + \alpha d)}. \quad (28)$$

168 Note that when $\lambda \geq 1 + 2a/\alpha d$, this case never produces a bifurcation as τ_λ is increased.

169 If $\lambda \neq 1$, diffusion causes the system to be able to reach a bifurcation for different values of critical τ_λ ,
170 which depend on the parameters b , β , α , a , and d . When $\lambda = 1$, the bifurcation conditions in the second
171 case, (25), are reached first for all values of b , β , α , a , and d and are identical to the bifurcation conditions
172 for the original Oster-Murray-Harris model.

The fixed, dimensional parameter values we use for this system are [28, 29, 30, 21, 31, 32]

$$\begin{aligned} A = 10^5, \text{ poise}, \quad D_0 = 10^{-9} \frac{\text{cm}^2}{\text{sec}}, \quad \beta = 10^{-2} \text{ cm}^2, \\ s = 10^2 \frac{1}{\text{sec}^2}, \quad c_0 = 10^4 \frac{\text{cell}}{\text{cm}^3}, \quad \rho_0 = 10^{-1} \frac{\text{g}}{\text{cm}^3}. \end{aligned} \quad (29)$$

Using the length scale $L = 1$ cm and the time scale $T = 10$ sec, these correspond to the non-dimensional parameters

$$\alpha = 10^{-4}, \quad a = 10^3, \quad d = 10^{-8}, \quad \beta = 10^{-2}. \quad (30)$$

We set $\lambda = 1$ and the two parameters, b and τ , are chosen so that the uniform steady state is linearly unstable to small random perturbations. Figure 1 shows the dispersion relation for the root of (19) with a positive real part for various values of τ with $b = 1.012$ ($B = 10.12$ dynes/cm²). A bifurcation occurs for $\tau > 2.469$ ($\tau_0 > 2.469 \times 10^{-5}$ dynes cm⁴/cell mg) where only one root has a positive real part for $k^2 \neq 0$ and the other two have negative real parts for $k^2 \neq 0$.

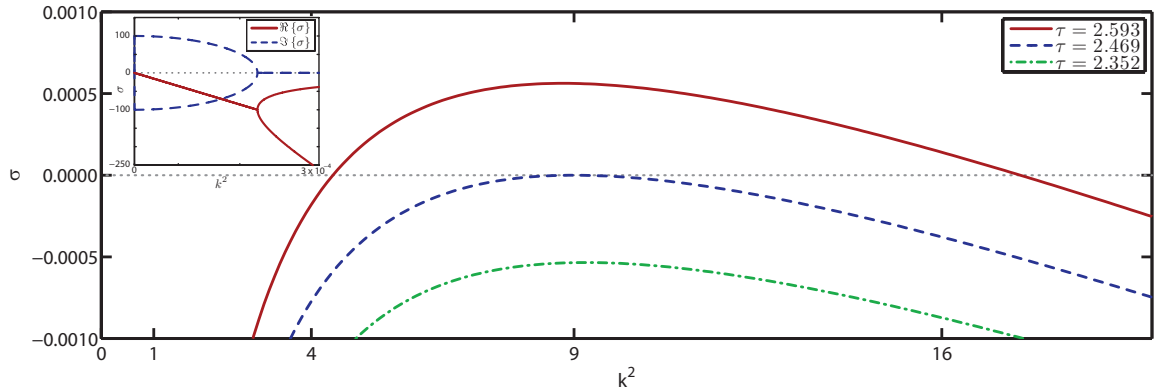


Figure 1: Dispersion relation as the bifurcation parameter τ is increased. The other two roots of the characteristic equation (19), σ_2 and σ_3 , are not shown and $\Re\{\sigma_2, \sigma_3\} \leq 0$. The roots are real except for a narrow region (shown in the inset), $0 \leq k^2 \leq 2 \times 10^{-4}$ for $\tau = 2.5933$, where the roots are complex with a negative real part. For $k^2 = 0$, the root is purely complex.

For $k^2 = 0$, the three roots are $\sigma = 0, \pm i/\sqrt{\alpha}$. The $\sigma = 0$ root corresponds to a uniform increase/decrease in the cell density. Since we can scale any perturbations in the total cell density out, we can neglect this root. The $\sigma = \pm i/\sqrt{\alpha}$ roots correspond to oscillatory translation of the medium. These oscillations do not affect the cell/ECM densities and do not contribute to the nonlinear dynamics of the system.

For $k^2 \neq 0$, the root with a positive real part has a zero imaginary part except for a thin region, $0 < k^2 < 2 \times 10^{-4}$. This narrow range of wavemodes are not admissible unless the domain is of sufficient length and are not admissible for biologically relevant domain sizes in this paper.

188 **4. Numerical Results**

189 We numerically solve equations (12) and (13) using finite differences in space and the backward Euler
 190 method in time. We take the domain to be periodic with length 2π cm discretised using a mesh with 300
 191 grid points and a time step of $\Delta t = 10^{-1}$. The uniform steady state solution for u and c is perturbed using
 192 a uniform random function with mean zero and the amplitude of the perturbations are 10^{-2} . The numerical
 193 solutions are independent of grid size for grids that are sufficiently refined to resolve the significant spatial
 194 frequencies in the problem and are independent of the exact random initial condition subject to a phase
 195 shift.

196 From a biological point of view, we suggest the precursors of clinically evident striae may appear as
 197 regions of ECM density that are below the uniform steady state value ($u = 0, \rho = c = 1$). Figure 2 shows
 198 the dilation and the ECM density after four days for $b = 1.012$ ($B = 10.12$ dynes/cm²) and $\tau = 2.593$
 ($\tau_0 = 2.593 \times 10^{-5}$ dynes cm⁴/cell mg). In the ECM density, we can see the development after four

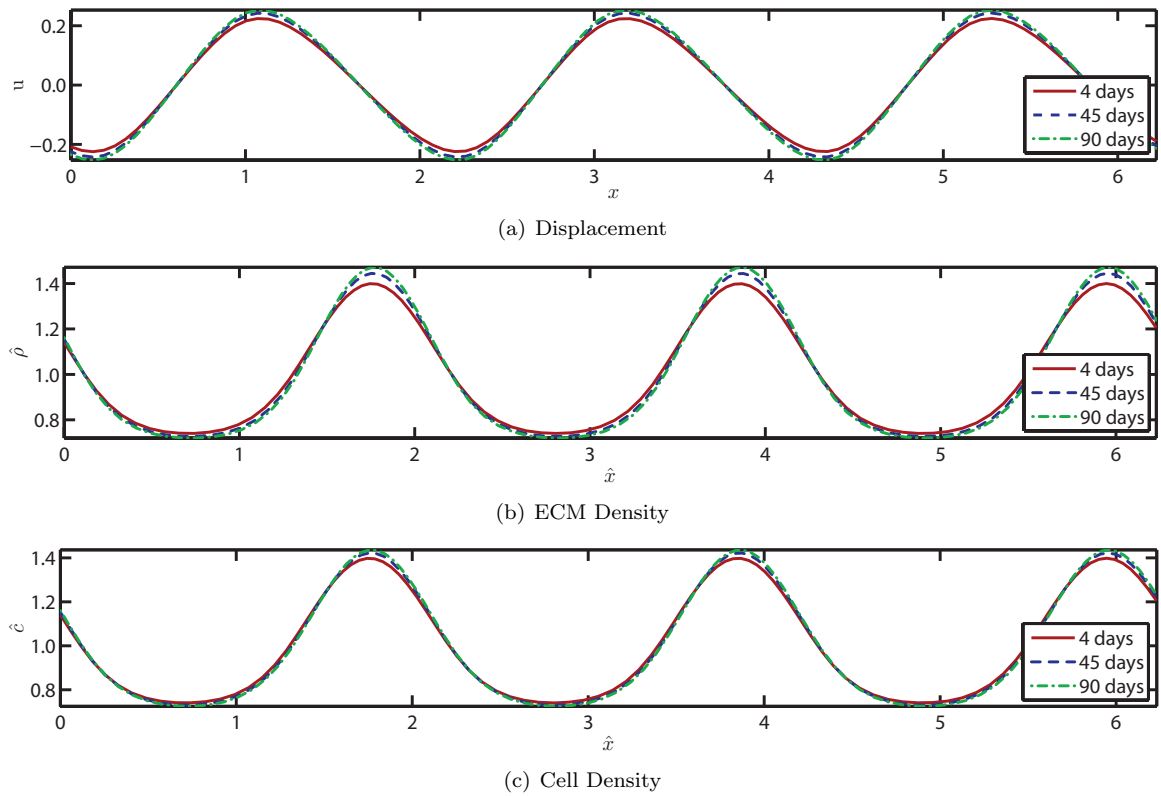


Figure 2: Numerical solution of equations (12) and (13) after 4, 45, and 90 days for $\hat{b} = 1.012$ and $\hat{\tau} = 2.593$. The ECM and cell densities are in the deformed variables in the deformed frame of reference. The minima after four days represent a 26% decrease in ECM density and the interlesional distance is approximately 2.09 cm.

199

200 days of regions where there is a significant decrease in density ($\min \rho \approx 74\% \rho_0$). This corresponds to
201 a $2.6 \times 10^{-2} \text{ g/cm}^3$ decrease in ECM density at the minima. The critical wavemode is $k_c = 3$ and this
202 corresponds to an interlesional distance of approximately 2.09 cm. The above numerical solution is not a
203 steady state solution. There is a second phase of growth that is due to the diffusion of fibroblasts. We can
204 see slow growth in the displacement and cell/ECM densities at 45 and 90 days. Here, the maximum cell
205 density has increased by a total 2% after 90 days. Any appreciable effect due to this slow growth will occur
206 over a long time frame and since we are interested in the onset of patterns that can become precursors of
207 striae distensae, we will focus on the initial development of patterning.

208 5. Discussion

209 We have described in detail a mathematical model proposed to capture the earliest events in the patho-
210 genesis of striae distensae. Motivation for the model is two-fold: first, we are interested in exploring the
211 relative contributions of both skin stretching and corticosteroid effects in the pathogenesis of striae; and
212 second, we are interested in the process of pattern formation in striae. Since stretch marks are frequently
213 observed to align perpendicular to the direction of skin tension, it is likely that the effects of tissue forces play
214 an important role in the genesis of striae, and in determining the patterns that form. These considerations
215 naturally lead to a mechanico-chemical type model.

216 We have shown that an intuitive model incorporating the density of fibroblasts, the density of extracel-
217 lular matrix, and a force balance equation that accounts for the contractile forces generated by fibroblasts
218 is able to predict, for biologically realistic parameters, periodic solutions for ECM density. In this periodic
219 spatial pattern, the precursors of striae are postulated to appear in close proximity to regions of ECM density
220 at their maximal densities.

221 As discussed in Section 1, an important unresolved issue with regard to the pathogenesis of striae
222 distensae is the relative contributions of skin stretching on one hand, and endocrine factors on the other.
223 While some investigators have suggested that striae may simply result from tissue rupture due to mechanical
224 loads, endocrine changes are present in association with many of the clinical situations in which striae are
225 encountered [12]. In our model, we are able to quantify the degree of stretch imposed on the skin by
226 our parameter τ_0 (which is a measure of the contractile force exerted by fibroblasts on the surrounding
227 extracellular matrix) since the fibroblasts respond to external mechanical force by opposing that force.
228 Furthermore, we are able to characterize the skin stiffness by adjusting B , a parameter proportional to the
229 Young's modulus. Since it is recognised that corticosteroid exposure may increase skin extensibility [23, 24],
230 we can model the effects of corticosteroid excess by reducing B . Our results suggest that the uniformity
231 of cutaneous extracellular matrix may be rendered unstable by either increasing τ_0 or decreasing B . We
232 obtain solutions that are qualitatively identical via two distinct mechanisms, and these distinct mechanisms
233 correspond to either stretching of the skin or increasing the extensibility of skin. From the clinical perspective,
234 these results suggest that either factor alone can induce striae. For example, in adolescence striae may
235 develop simply as a result of skin stretching (increasing τ_0); abnormal endocrine factors and changes to skin
236 extensibility are not needed. Conversely, the application of potent topical steroids to the skin results in a
237 reduction in the mucopolysaccharide content [33]—and this may account for the increases in skin extensibility

238 reported [24]—with the subsequent development of striae under normal skin tension. Indeed, the skin is not
239 subject to increased stretch in a number of clinical situations where striae distensae arise: cachexia (**wasting**
240 **syndrome**) and other causes of severe weight loss are two examples where decreased dermal substrate and the
241 resultant increased extensibility of skin may be the sole aetiological factor. Interestingly, of seven patients
242 treated with high dose intravenous corticosteroid for alopecia areata (**hair loss in one or more circular spots**
243 **on the scalp**) [23] one patient developed striae distensae on the thighs within four days of the infusion. In
244 this study skin extensibility was measured and was shown to increase within hours of the infusion, reaching
245 a maximal extensibility at around four days. In ageing skin it has been reported that there is a decrease
246 in extensibility [34] (corresponding to an increase in b in our model) and perhaps explaining why striae are
247 only rarely observed to develop in mature adults. Finally, our model is not inconsistent with both increased
248 stretch and dermal changes acting together in the genesis of striae. For example, in pregnancy lesions
249 may arise due to the synergistic effect of both increased skin extensibility on one hand, and the increased
250 mechanical forces acting on the skin secondary to a rapidly distending abdomen on the other.

251 Striae distensae in the skin often exhibit typical patterns. Although the individual lesions are linear and
252 usually five to ten centimetres long, multiple lesions are the norm and are always aligned perpendicular to
253 the axis of skin tension. Striae develop on the back of adolescent males as parallel streaks perpendicular
254 to the direction of vertical growth. Conversely, striae on the breasts in females often have a radial pattern
255 indicating that the lines of tension in the skin in an enlarging breast are circumferential. Although our
256 model is one-dimensional, we are able to predict the existence of periodic solutions that arise parallel to the
257 direction of tension, consistent with the clinical observations noted above. We have shown the dimensional
258 value for the wavelength of our solutions can be adjusted to approximate one centimetre. This result is in
259 good agreement with the average distance found between striae that are aligned in a parallel arrangement.

260 Although we have demonstrated that increases in the contractile forces exhibited by fibroblasts or in-
261 creases to skin extensibility may lead to dermal inhomogeneity of ECM density, we have been unable to
262 provide a definite link between these changes and the mast cell degranulation that is known to be an early
263 event in the pathogenesis of striae. However, it is known that increases in GAG (glycosaminoglycan) den-
264 sity is a very early finding in striae rubra [15], and it is unclear at present whether these changes predate,
265 coincide with, or follow the mast cell associated elastolysis. Our model, in predicting periodicity in ECM
266 density along the axis of skin tension, adds weight to the hypothesis that local increases in GAG density
267 (since the GAG is part of the ECM) may precede mast cell degranulation. Since fibroblasts in early striae are
268 known to exhibit aberrant gene expression profiles, one possible pathogenic mechanism is apparent: secreted
269 fibroblast products may exist locally in higher concentrations where the ECM density is higher and thus
270 lead to spatially dependent mast cell recruitment and subsequent elastolysis.

271 In summary, we present a conceptually simple but mathematically complex model that attempts to
272 account for the earliest events in the pathogenesis of striae distensae. We suggest that the results are suffi-
273 ciently robust enough to provide evidence for the existence of an important symmetry breaking mechanism
274 that is able to distinguish between two fundamentally different and clinically relevant causes.

275 **Appendix A. Transformation of the Traction Force into the Material Frame of Reference**

276 We begin with the traction force exerted by the fibroblasts on the ECM in the spatial frame of reference:

$$\tau(\hat{c}, \hat{\rho}) = \tau_0 \frac{\hat{c}}{1 + \lambda \hat{c}^2} \left(\hat{\rho} + \beta_0 \frac{\partial^2 \hat{\rho}}{\partial \hat{x}^2} \right) \quad (\text{A.1})$$

277 where the hats designate the spatial frame of reference. We transform \hat{c} and $\hat{\rho}$ from the spatial frame to the
278 reference frame using the Jacobian of the deformation gradient, $J = 1 + \partial u / \partial x$, to get

$$\hat{c} = \frac{c}{1 + u_x} \quad \text{and} \quad \hat{\rho} = \frac{\rho_0}{1 + u_x}, \quad (\text{A.2})$$

279 where $u_x = \partial u / \partial x$. Substituting (A.2) into (A.1) and simplifying yields the equation

$$\tau_0 \frac{c \rho_0}{(1 + u_x)^2 + \lambda c^2} \left(1 + \beta (1 + u_x) \frac{\partial^2}{\partial \hat{x}^2} \left(\frac{1}{1 + u_x} \right) \right). \quad (\text{A.3})$$

280 Next, we convert the spatial derivatives to the reference frame using

$$\frac{\partial}{\partial \hat{x}} = \frac{1}{1 + u_x} \frac{\partial}{\partial x} \quad (\text{A.4})$$

281 to obtain the traction force in the reference frame:

$$\tau \left(c, \frac{\partial u}{\partial x} \right) = \tau_0 \frac{c \rho_0}{(1 + u_x)^2 + \lambda c^2} \left(1 + \beta \frac{\partial}{\partial x} \left(\frac{1}{1 + u_x} \frac{\partial}{\partial x} \left(\frac{1}{1 + u_x} \right) \right) \right). \quad (\text{A.5})$$

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