Theoretical, computational, and experimental characterization of nematic elastomers

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ABSTRACT

Nematic elastomers are programmable soft materials that display large, reversible, and predictable deformation under an external stimulus such as a change in temperature or light. They are composed of a lightly crosslinked polymer network with stiff, rod-like liquid crystal molecules incorporated within the polymer chains. In thermotropic nematic elastomers, the liquid crystals undergo a continuous and reversible phase transition between the randomly oriented isotropic state and the highly oriented nematic state. Further, there is a direct thermo-mechanical coupling between the underlying temperature-responsive orientational order of the liquid crystal molecules and the macroscopic shape change of the surrounding elastomer chains. Finally, these materials display an unusually soft behavior. These remarkable properties make them promising materials for applications in aerospace as deployable structures and skins, in biomedical engineering as a soft pump, and in communications as the actuation mechanism in a reconfigurable antenna. Motivated by these applications, this thesis discusses the theoretical, computational, and experimental characterization of nematic elastomers.

We begin by investigating an example of actuation that takes advantage of the programmable, soft nature of these materials as well as instabilities associated with large deformation. We outline the multi-stable equilibrium solutions to a cylindrical balloon subjected to internal inflation, the material’s microstructure formation due to this deformation, and its use as a soft pump with large ejection fraction, which involves a snap-through instability. Then we extend the Agostiniani-DeSimone-Dolzmann relaxed energy to a generalized Mooney-Rivlin constitutive relation and study four examples of Ericksen’s universal deformations—the inflation of cylindrical and spherical balloons, the cavitation of a disk, and the bending of a block.

We then move beyond the modeling of ideal materials and present a new constitutive relation for isotropic-genesis polydomain nematic elastomers. It is based on internal variables that describe the fine-scale domain patterns and evolve according to a kinetic process with dissipation. We discuss the model’s implementation in the commercial finite-element software, ABAQUS, and study the problem of torsion of a cylinder. We identify an interesting instability at large torsional strains as a result of the Poynting effect. Finally, we present the design of a thermo-
mechanical tensile setup and the experimental results for strain-rate dependence and temperature-dependence of samples that we synthesize in-house.
PUBLISHED CONTENT AND CONTRIBUTIONS


V. Lee performed all calculations, created all plots, and co-wrote the manuscript.

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Chapter 1

INTRODUCTION

1.1 Nematic elastomers

Liquid crystal elastomers (LCEs) are remarkable stimuli-responsive materials that have recently been explored for their use in actuation [77]. LCEs are lightly crosslinked elastomer networks with liquid crystal molecules incorporated into the underlying polymer chains. P. G. de Gennes first envisioned the coupling between the orientational order of liquid crystals with the macroscopic shape change of a crosslinked elastomer network in 1975 [21]. There are different types of liquid crystal elastomers, including nematic, cholesteric, and smectic, and they can be responsive to different types of stimuli, including a change in temperature, pH, electric field, and light.

![Figure 1.1: Isotropic-nematic phase transition in nematic elastomers with the isotropic with $r = 1$ at high temperatures and nematic phase with $r > 1$ at low temperatures.](image)

Of particular interest are nematic elastomers, which are liquid crystal elastomers with stiff, rod-like liquid crystal molecules. They undergo an isotropic-to-nematic transition accompanied by a significant stretch (by as much as a factor of two or more) along the nematic director (in the direction parallel to a unit vector $n$) and lateral contraction, as shown schematically in Figure 1.1. In this thesis, we focus on thermotropic nematic elastomers, which undergo the phase transition at a characteristic nematic-isotropic temperature $T_{ni}$. At temperatures above $T_{ni}$, the
material is in the isotropic state, where the liquid crystal molecules are randomly oriented, and at temperatures below $T_{nl}$, the material is in the nematic state, where the liquid crystal molecules are aligned along a preferred direction, denoted by $n$. Because of the direct thermo-mechanical coupling between the nematic orientation and the surrounding polymer network, this phase transition induces a deformation of the underlying polymer chains in the nematic elastomer, so a macroscopic shape change is observed, in which the polymer chains stretch parallel to the mesogen alignment and contract perpendicular to it. The phase transition, and consequently the change of shape, is continuous and completely reversible. The degree of order observed in the liquid crystal molecules determines the degree of anisotropy, denoted by $r$ throughout this thesis.

These materials can be synthesized as a monodomain or polydomain. Monodomains can be thought of as the elastomer equivalent of a single crystal, in which the entire sample is aligned along one direction in the nematic state. A polydomain sample is macroscopically randomly oriented in the nematic state, but at a mesoscale, there is nematic alignment within each domain (see Figure 1.2 for a schematic).

![Schematics of polydomain and monodomain nematic elastomer.](image)

Figure 1.2: Schematics of polydomain and monodomain nematic elastomer.

### 1.2 Actuation
The study of materials for actuation applications traditionally focused on systems driven by mechanisms such as pneumatics, hydraulics, or motors. Active materials such as piezoelectrics, dielectric elastomers, a type of electroactive polymer (EAP), and shape memory alloys which undergo the austenite-martensite phase transition, offer much in the way of actuation without complex hardware and moving parts. Using the material as the machine [6] is a powerful idea motivating the study of active materials.

Nematic elastomers are advantageous choices in the field of active materials because they are soft, they can deliver a large energy per unit volume, their shape change is completely reversible in the presence or absence of the stimuli, and their director field
is programmable [2, 48, 49, 57, 76]. Further, it is possible to incorporate photo-active molecules into nematic elastomers, giving rise to photo-active materials that are actuated by light. Some compelling examples include refreshable Braille displays for reading instruments for the visually impaired [14], untethered robotic matter [41], and shape morphing [26]. Additionally, nematic elastomers can be programmed for wrinkle control in thin sheets due to microstructure development, for possible applications in deployable structures for aerospace applications [56].

1.3 Beyond actuation from flat sheets
Nematic elastomers have been exploited for programmable actuation and shape-morphing of thin sheets. Modes, Bhattacharya, and Warner [48, 49] suggested that if sheets of nematic elastomer with prescribed director patterns were fabricated in the nematic state and subsequently heated, they could deform out of plane into three-dimensional shapes. For example, a +1 disclination with an azimuthal director distribution would deform into a cone. This was demonstrated in nematic glasses by de Haan et al. [22]. Ware et al. [76] developed a method of synthesizing nematic elastomers where the director pattern could be written pixel by pixel on flat sheets and demonstrated the formation of these cones. Moreover, they showed that this actuation was extremely robust, as the cone-lifting weights were many hundreds of times larger than the structure itself. Since then, there have been a number of other studies on nematic elastomers [2, 47, 50, 57, 58], including the inverse problem of identifying the director pattern that would lead to a given actuated shape [3, 57]. All of these works address the programming and actuation of initially flat sheets.

Recent advances in 3D printing and other methods of directed synthesis have enabled the synthesis of curved shells [4, 32], and such structures change shape upon heating. In particular, Ambulo et al. demonstrated dramatic snap-through buckling of structures involving regions of positive Gauss curvature and regions of negative Gauss curvature [4]. More recently, magnetic fields have been used to independently control director orientation during 3D printing [68]. These developments in synthesis techniques motivate the work in Chapter 2, where we analyze a balloon made of nematic elastomer, subjected to internal inflation. The study of these materials in different configurations beyond flat sheets opens the doors to applications with more complex boundary conditions and loading configurations.
1.4 Microstructure formation

The interesting material properties of nematic elastomers are due to the interplay amongst features at distinct length scales. The macroscopic shape of the lightly crosslinked polymer network, which can be on the order of centimeters, is determined by the nematic mesogens, which can have a length on the order of nanometers, and domains of nematic alignment can exist in some mesoscale with a characteristic length of micrometers.

![Figure 1.3: Schematic of the experiment in which a monodomain sample is pulled perpendicular to its nematic director, giving rise to stripe domains, which can be seen under polarized light microscopy (in the left-most circle).](image)

The stripe-domain formation in monodomains pulled perpendicular to their nematic alignment exhibit the classical example of fine-scale microstructure formation (see Figure 4 of [42], also shown schematically in Figure 1.3). In this experiment, a monodomain sample with a uniform nematic alignment is clamped on one end and pulled uniaxially in the direction perpendicular to the nematic director. The liquid crystal molecules rotate to align themselves parallel to the direction of stretch, and they do so by forming alternating stripes in which the molecules rotate at opposite angles. When the liquid crystal molecules complete this reorientation process, the sample becomes a monodomain with the nematic director parallel to the direction of stretch. The stripe domains can be seen using polarized light microscopy, an example of which is shown in the left-most circle of Figure 1.3. The stripes are relatively uniform, and the width can vary depending on the specific synthesis methods and possibly also the geometry of the sample. The stripes exhibited by the
samples in our lab are about 70 micrometers in width, whereas the stripes in Figure 4 of [42] have a width closer to 15 micrometers.

At the beginning of the experiment, the clamped monodomain sample appears transparent because all of the liquid crystal molecules are aligned uniformly, but as the experiment progresses, the sample becomes opaque as the microstructure develops, and the liquid crystal molecules are no longer uniformly aligned. Eventually, when the liquid crystal molecules have finished reorienting and are again uniformly aligned, the sample has become a monodomain and therefore appears transparent.

The microstructure formation gives rise to the phenomenon of soft elasticity—the reorientation of the mesogens happens at zero stress, which is depicted by the stress plateau in stress-strain curves of the uniaxial extension of monodomains pulled perpendicular to their nematic alignment (see Figure 9 of [43]). Stripe domains are discussed in the context of the cylindrical balloon explored in Chapter 2.

A related experiment is the uniaxial stretch of a polydomain sample, which undergoes a transition known as the polydomain-to-monodomain transition. The associated stress-strain curve also exhibits soft elasticity, as mesogens within each mesoscale domain reorient towards the direction of most stretch. The soft behavior has largely been studied in uniaxial deformation, and more recently in biaxial deformation [71]. This motivates Chapter 3, where we study the consequences of softness on complex deformations.

1.5 Viscoelasticity and damping in nematic elastomers

It is important to understand and characterize the effect of viscoelasticity, or time dependence, in nematic elastomers if they are to be used in actuation applications, which are typically cyclic in nature. Existing work in this field includes stress-relaxation experiments on these materials, as well as uniaxial experiments at varying strain rates, e.g. through the polydomain-monodomain (P-M) transition [5, 16, 36, 54, 67, 69]. Some of these authors have also used dynamic mechanical analysis (DMA) of monodomain samples under simple shear. Hotta and Terentjev also applied the principle of time-temperature superposition to build a master curve of the material response over a range of frequencies using data from tests at different temperatures, e.g. [5, 35].

The large hysteresis between the loading and unloading stress-strain curves in uniaxial experiments on polydomain samples, e.g. in Figure 5 of [5], is indicative of the material’s ability to dissipate a large amount of energy. Merkel et al. [46]
investigated the effect of temperature, and therefore degree of anisotropy, in the dynamic loading of polydomain nematic elastomers. A combination of the reorientation of the liquid crystal molecules and the viscosity of the polymer chains provides a mechanism to dissipate and absorb applied mechanical energy. Clarke et al. discussed the potential for nematic elastomers in mechanical damping applications due to the internal relaxation of the liquid crystal molecules, independent of the polymer backbone [17]. Examples of applications where efficient dissipation of mechanical energy are desirable include the automotive, aerospace, and white goods (e.g. washing machines, refrigerators) industries. One can imagine tuning the mesogen response for variable vibration and noise suppression.

We address the formulation of a finite-deformation constitutive relation that could model such viscoelastic effects in Chapter 4, which incorporates viscosity associated with the polymer network as well as evolution of some internal variables according to a dissipative kinetic process.

1.6 Nematic elastomers as an engineering material

Studies of monodomain nematic elastomers are extremely useful to characterize a structure or be able to predict material response, especially for actuation applications. However, polydomains, with their domains of fine-scale microstructure, are easier and less expensive to manufacture because they require no director alignment, and can be viewed as an engineering material.

Previous theories largely dealt with simple deformations or idealized materials. The goal of Chapter 4 is to formulate a model for nematic elastomers as an engineering material—with microstructure formation through the relaxed energy and viscoelasticity through the internal variables.

1.7 Thesis outline

This thesis is organized in the following way: First, in Chapter 2, we study the actuation of a cylindrical nematic elastomer balloon. This work is a step away from the established field of actuation from flat sheets, as we focus on actuation from an already three-dimensional reference state. We analyze the deformation of a cylindrical shell of a patterned nematic elastomer under pressure, show that it can undergo an enormous change of volume with changing temperature, and suggest its application as a pump with extremely high ejection fraction.
In Chapter 3, we build upon the framework of DeSimone and Dolzmann [23] and Agostiniani and DeSimone [1] for the relaxed energy of nematic elastomers. We again extend this energy into a generalized Mooney-Rivlin type energy density to capture correctly the elasticity of these materials at very high stretch, and we use this model to solve four examples of Ericksen’s so-called universal deformations: expansion of a spherical balloon, expansion of a cylindrical balloon, cavitation of a disk, and bending of a block. The solutions are presented for varying anisotropy parameters, including the case where the material is isotropic, which corresponds to a rubber without liquid crystals.

Chapter 4 presents an engineering model involving internal variable evolution that captures the macroscopic behavior of isotropic-genesis polydomain nematic elastomers. We present the formulation for the model, its implementation in the commercial finite-element software ABAQUS, and some results under various deformations, including planar extension, biaxial stretch, and torsion. Our numerical results identify an unusual torsional instability driven by the Poynting effect.

In Chapter 5, we present the synthesis technique of thermotropic nematic elastomer samples, the design of our custom-built thermo-mechanical test setup, and the test results from uniaxial extension of these samples in the experimental setup at various crosslinked configurations and temperatures.

Finally, we summarize the findings of this thesis and conclude with comments on the future outlook of this exciting and expansive field of research in Chapter 6. Nematic elastomers play only a small role in the field of stimulus-responsive actuators. There are many promising avenues for further research and development of these materials, experimentally, theoretically, and computationally.
Chapter 2

ACTUATION OF CYLINDRICAL NEMATIC ELASTOMER BALLOONS


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2.1 Introduction

The goal of this work is two-fold. The first is to explore the combination of programmed synthesis of nematic shells and the geometric instabilities associated with the large deformation of slender structures. Similar instabilities have been exploited in other stimuli-responsive materials including electroactive materials [33, 63, 84]. In this work, we focus on the so-called aneurysm instability of pressurized cylinders [28]. As observed in long toy balloons, one observes a discontinuous change of radius (or volume) with an increase of pressure: typically the balloon inflates till it reaches a particular radius, beyond which point a bump (aneurysm) with a significantly larger radius appears in this region, and it propagates through the entire balloon before the radius further increases. We explore the response of a cylindrical shell made of a nematic elastomer and study how the isotropic-nematic phase transition affects this instability. Our work is closely related to that of Giudici and Biggins [31] who recently studied the ballooning instability in both nematic and isotropic LCEs using a Gent-style energy. He et al. [34] have studied the anomalous behavior of (isotropic-genesis polydomain) nematic balloons under tension. We then show how this instability can be used as a high ejection-fraction pump. The second goal is to study actuation and shape-morphing in the presence of mechanical loads. The prior literature has largely focused on free recovery.

We introduce the model of the nematic elastomers at large deformation in Section 2.2 and analyze the deformation of a nematic elastomer cylinder under internal pressure in Section 2.3. We then use the results to motivate a pump with extremely large ejection fraction in Section 2.4.
2.2 Large deformation model of nematic elastomers

We begin with the neo-classical theory of nematic elastomers following Bladon, Terentjev and Warner [10, 77]. The state of a liquid crystal elastomer is described by an anisotropy parameter $r$, a director $n$ and the deformation gradient $F$ relative to a stress-free reference configuration with anisotropy parameter $r_0$ and director $n_0$. The anisotropy parameter is a function of temperature with $r = 1$ in the isotropic state above the transformation, and gradually increases with decreasing temperature so that $r > 1$ in the nematic state. We consider the material to be incompressible so that $\det F = 1$. The neo-classical theory considers the entropy of the polymer chains in the Gaussian approximation, and the free energy density is given as

$$W_{WT}(F, n, r) = \frac{\mu}{2} \left( \text{tr} \left( \ell_n F^T \ell_n^{-1} F \right) - 3 \right),$$

(2.1)

where $\mu$ is the shear modulus of the material, and

$$\ell_n = r^{-1/3} \left( I + (r - 1) n \otimes n \right)$$

(2.2)

$$\ell_{n_0} = r_0^{-1/3} \left( I + (r_0 - 1) n_0 \otimes n_0 \right)$$

(2.3)

are the step-length tensors in the current and reference configurations that collect the anisotropy parameter and the director. It is easy to show that

$$W_{WT}(F, n, r) = W_{NH} \left( \ell_n^{-1/2} F \ell_{n_0}^{1/2} \right),$$

(2.4)

where $W_{NH}(F) = \frac{\mu}{2} (\text{tr} \ C(F)) - 3)$, with $C(F) = F^T F$, is the neo-Hookean energy density which describes the entropy of polymer chains in ordinary rubber in the Gaussian approximation [72].

The neo-classical theory is known to describe complex features of nematic elastomers at finite, but moderate, deformation. However, at extremely large stretches, the Gaussian approximation does not hold, and this theory does not adequately describe the stiffening much like its neo-Hookean counterpart. Various constitutive relations are used to describe rubber in this high-stretch regime. A common feature of many of these models is that the energy density depends only on principle stretches $\lambda_i$ of $F$ (equivalently the eigenvalues $\lambda_i^2$ of $C(F)$):

$$W_E(F) = f(\lambda_1, \lambda_2, \lambda_3).$$

(2.5)

For example, in the Ogden model [53] the energy density is

$$W_O(F) = \sum_{p=1}^{N} \frac{\mu_p}{\beta_p} \left( \lambda_1^{\beta_p} + \lambda_2^{\beta_p} + \lambda_3^{\beta_p} - 3 \right),$$

(2.6)
where $N$, $\mu_p$, and $\beta_p$ are material constants. The shear modulus is $\mu = \frac{1}{2} \sum_{p=1}^{N} \mu_p \beta_p$.

When $N = 1$ and $\beta_1 = 2$, the Ogden energy is the neo-Hookean energy, and when $N = 2$, $\beta_1 = 2$, and $\beta_2 = -2$, the Ogden energy is the Mooney-Rivlin energy. We use the Ogden energy to demonstrate our results following [63], though we can adapt them to any constitutive relation that describes the high stretch behavior. We adopt the elastic energy density (2.5) to nematic elastomers analogously to (2.4). See [1] for similar energies and their relaxation in the ideal case. Other approaches have been proposed to capture the high-stretch regime including the logarithmic correction by Gent [27] which was used by Giudici and Biggins [31] in their work.

Further, the cross-link density and the polymer network may carry an imprint of the initial director, and this leads to a breaking of symmetry (isotropy) leading to the preference of the director to remain in the original orientation. Such an interaction can be described using an additional non-ideal energy density [9]:

$$W_{NI}(F, n) = \alpha \frac{\mu}{2} \text{tr} \left( F (I - n_0 \otimes n_0) F^T n \otimes n \right). \tag{2.7}$$

Note that this energy is minimized when $n = n_0$. Putting these together, we take the energy density of the nematic elastomer to be

$$W(F, n, r) = W_E \left( \ell_n^{-1/2} F \ell_n^{1/2} \right) + W_{NI}(F, n). \tag{2.8}$$

For future use, we note a particular invariance of this energy density. Let $Q$ be a rotation tensor that leaves the reference director invariant: $Q n_0 = \pm n_0$. Then, we claim that

$$W(QFQ^T, Qn, r) = W(F, n, r). \tag{2.9}$$

Note that

$$C((Q \ell_n Q^T)^{-1/2}(QFQ^T)\ell_n^{1/2}) = \ell_n^{1/2} (QFQ^T)^T (Q \ell_n Q^T)^{-1} (QFQ^T) \ell_n^{1/2}$$
$$= \ell_n^{1/2} Q F^{T} \ell_n^{-1} F Q^{T} \ell_n^{1/2}$$
$$= Q \ell_n^{1/2} F^{T} \ell_n^{-1} F \ell_n^{1/2} Q^{T}$$
$$= QC(\ell_n^{-1/2} F \ell_n^{1/2}) Q^{T}, \tag{2.10}$$

where we have used the invariance of $n_0$ under $Q$ in the third equality. It follows that both tensors have the same eigenvalues and thus the same Ogden energy density. A similar calculation holds for the non-ideal energy density as well, thereby establishing (2.9).
2.3 Inflation of a nematic cylinder

Consider a cylindrical shell of initial (reference) length $H$, inner radius $R_i$, and undeformed outer radius $R_o$ subjected to an internal pressure $p$. Following Rivlin [61] and Ericksen [25], the deformation of the cylinder is described by the universal volume-preserving deformation involving radial expansion, axial extension and torsion (see Figure 2.1). The mapping is

\[
\begin{cases}
\rho = \rho(R) \\
\theta = \Theta + DZ \\
z = \xi Z
\end{cases}, \tag{2.11}
\]

where \( \{R, \Theta, Z\} \) and \( \{\rho, \theta, z\} \) denote the cylindrical coordinate system in the reference and deformed coordinate systems respectively. \( \rho(R) \) describes the radial expansion, \( D \) the twist, and \( \xi \) the axial stretch. The deformation gradient in the cylindrical coordinate system is

\[
F = \begin{pmatrix}
\rho' & 0 & 0 \\
0 & \frac{\rho}{R} & \lambda RD \\
0 & 0 & \xi
\end{pmatrix} = \begin{pmatrix}
\frac{1}{\xi} & 0 & 0 \\
0 & \lambda & \lambda RD \\
0 & 0 & \xi
\end{pmatrix}, \tag{2.12}
\]

where we have introduced the hoop stretch \( \lambda(R) = \rho(R)/R \) and used the incompressibility to obtain the second equality. There is an off-diagonal term in the deformation gradient because we would like to allow shear or twist that may accompany director reorientation. We will see later that this indeed plays a role.
We assume that the director both in the reference and deformed configuration are tangential to the cylinder and make an angle $\phi_0$ and $\phi$, respectively, with the azimuthal coordinate. Thus, in cylindrical coordinates,

$$
\mathbf{n}_0 = \begin{pmatrix} 0 \\ \cos \phi_0 \\ \sin \phi_0 \end{pmatrix} \text{ and } \mathbf{n} = \begin{pmatrix} 0 \\ \cos \phi \\ \sin \phi \end{pmatrix}.
$$

(2.13)

The total potential energy of the system is

$$
\Phi = \int_{\Omega} W(\mathbf{F}, \mathbf{n}, r) dV - p \Delta V,
$$

(2.14)

where $\Delta V$ is the difference in the deformed and undeformed volumes. Applied to a balloon with height $H$, we obtain

$$
\Phi = \int_0^H \int_0^{2\pi} \int_{R_i}^{R_o} W(\mathbf{F}, \mathbf{n}, r) RdRd\Theta dZ - p \left( \pi \rho^2 \xi H - \pi R^2 H \right) \bigg|_{R=R_i} \approx 2\pi R_i H W(\mathbf{F}, \mathbf{n}, r) - p \pi R^2 H \left( \xi \lambda^2 - 1 \right).
$$

(2.15)

Above we have assumed that the shell is thin, $T := (R_o - R_i) \ll R_i$, to evaluate the integral.

For a given pressure $p$ and anisotropy parameter $r$, we can now find the equilibrium as

$$
\frac{\partial \Phi}{\partial \lambda} = \frac{\partial \Phi}{\partial \xi} = \frac{\partial \Phi}{\partial D} = \frac{\partial \Phi}{\partial \phi} = 0.
$$

(2.16)

Physically, these equations describe the balance between the hoop stress in the cylinder and the internal pressure, the balance between the axial stress and the internal pressure, the balance of torque, and the balance of internal (material) torque on the director respectively.

To demonstrate the results, we consider a cylinder where the initial director is axial ($\phi_0 = 90^\circ$) and which is mildly nematic with initial anisotropy parameter $r_0 = 2$. The rest of the parameters are shown in Table 2.1, and were chosen to be broadly consistent with an experiment conducted in our laboratory. We fix the current anisotropy parameter $r$ and the hoop stretch $\lambda$ and solve (2.16) for the pressure $p$, axial stretch $\xi$, the twist $D$, and the current director angle $\phi$. We find that the system has two solutions, shown in Figure 2.2 for four different current anisotropy parameters $r^1$ and the classical neo-Hookean case, $r_0 = r = 1$. Note that the pressure has been normalized as $\tilde{p} = \frac{p R_o}{\mu (R_o - R_i)}$, where $\mu = \frac{1}{2} \sum_{p=1}^{3} \mu_p \beta_p = 6.80 \cdot 10^4$ Pa represents the reference shear modulus.

$^1$There is a third unstable solution where the director does not rotate, which we ignore.
Figure 2.2: Inflation of a nematic cylinder: (a) Pressure vs. hoop stretch (solutions 1 and 2), (b) Axial stretch vs. hoop stretch (solutions 1 and 2), (c) Director angle vs. hoop stretch (solution 1), (d) Twist vs. hoop stretch (solution 1), (e) Director angle vs. hoop stretch (solution 2), (f) Twist vs. hoop stretch (solution 2).
Table 2.1: Table of parameters

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Inner radius</td>
<td>$R_i$ 1 cm</td>
</tr>
<tr>
<td>Outer radius</td>
<td>$R_o$ 1.05 cm</td>
</tr>
<tr>
<td>Height of cylinder</td>
<td>$H$ 5 cm</td>
</tr>
<tr>
<td>Initial director angle</td>
<td>$\phi_0$ 90°</td>
</tr>
<tr>
<td>Initial anisotropy parameter</td>
<td>$r_0$ 2</td>
</tr>
<tr>
<td>Non-ideality parameter</td>
<td>$\alpha$ 0.3</td>
</tr>
<tr>
<td>Ogden model shear modulus</td>
<td>$\mu_1$ $1.0 \cdot 10^5$ Pa</td>
</tr>
<tr>
<td>Ogden model shear modulus</td>
<td>$\mu_2$ $1.904762 \cdot 10^2$ Pa</td>
</tr>
<tr>
<td>Ogden model shear modulus</td>
<td>$\mu_3$ $-1.5873 \cdot 10^3$ Pa</td>
</tr>
<tr>
<td>Ogden model constant</td>
<td>$\beta_1$ 1.3</td>
</tr>
<tr>
<td>Ogden model constant</td>
<td>$\beta_2$ 6</td>
</tr>
<tr>
<td>Ogden model constant</td>
<td>$\beta_3$ -3</td>
</tr>
</tbody>
</table>

Consider the first solution, Figure 2.2(a)-(d). We observe that for $r_0 \neq r$, hoop stretch vs. pressure does not pass through $(1, 0)$ but through $((r/r_0)^{-1/6}, 0)$ since the change of the anisotropy parameter gives rise to a spontaneous deformation of the cylinder. The hoop stretch vs. pressure is non-monotone (Figure 2.2(a)): the pressure initially increases but then drops before increasing again with increasing hoop stretch. This reflects the well-known balloon instability: with increasing pressure, the radius increases till it reaches a critical pressure at which it jumps to a large radius. The onset and the extent of this instability is amplified in nematic elastomers due to a rotation of the director. Figure 2.2(c) shows that the director begins to rotate with inflation, reaching the hoop direction asymptotically. To understand this, an increase in radius increases the volume more than an increase in axial stretch, since the former leads to an increase of included area rather than length. Therefore, the pressure seeks to increase the circumference by reorienting the director. This reorientation also leads to a decrease of the axial stretch (Figure 2.2(b)). Consequently the axial stretch is also non-monotone: it decreases during reorientation but increases again as the director stabilizes. Finally, the reorientation leads to a twist in the cylinder (Figure 2.2(d)). The magnitude of all of these trends increases with increasing anisotropy parameter $r$. In particular, the critical pressure decreases and the change of hoop stretch increases with increasing anisotropy parameter $r$.

The reorientation, however, is resisted by the non-ideality as shown by varying the non-ideality parameter $\alpha$ in Figure 2.3. Note that the director rotation from the vicinity of the initial orientation $\phi_0 = \frac{\pi}{2}$ is increasingly delayed as the non-ideality
Figure 2.3: Effect of non-ideality parameter $\alpha$ upon pressure, axial stretch, director angle, and angle of twist for $r_0 = r = 2$.

The parameter increases. The balance between the pressure-assisted reorientation and the non-ideality-mediated resistance leads to the observed behavior.

The second solution, Figure 2.2(a)-(b),(e)-(f), is very similar to the first, except that the reorientation and twist change sign. The pressure vs. hoop-stretch and the axial stretch vs. hoop-stretch curves remain unchanged. Consequently, both solutions have the same pressure vs. volume curves which are shown in Figure 2.4(a). The volume strain is plotted on a logarithmic scale due to the dramatic change of volume during the instability.

It is useful to understand the origin of the two solutions. The material is not chiral, and neither is the initial configuration. Therefore a breaking of the chiral symmetry by rotation of the director has to be accompanied by a symmetry-related counterpart. To elaborate on this, recall the invariance (2.9). Let $Q$ be a 180° rotation about the azimuthal direction,

$$Q = \begin{pmatrix} -1 & 0 & 0 \\ 0 & 1 & 0 \\ 0 & 0 & -1 \end{pmatrix}. \quad (2.17)$$
Note that $Qn_0 = -n_0$ so that it satisfies the requirement for (2.9). It is easy to check that for $F$ and $n$ in (2.12) and (2.13),

$$ QFQ^T = \begin{pmatrix} \frac{1}{\lambda} & 0 & 0 \\ 0 & \lambda & -\lambda D \\ 0 & 0 & \xi \end{pmatrix}, \quad Qn = \begin{pmatrix} 0 \\ \cos \phi \\ -\sin \phi \end{pmatrix}. \tag{2.18} $$

Thus, the invariance (2.9) implies that any solution with chirality has a symmetric counterpart with the same radial and azimuthal stretches.

The presence of the two symmetric solutions enables the formation of stripe domains that avoid overall torsion as shown in Figure 2.4(b). We divide the cylinder into short cylindrical rings and alternate between the two solutions. This leads to a continuous deformation, where one ring twists one way and the other the other way in an alternating pattern, but they meet continuously across the boundaries as indicated by the initially straight fiducial dashed line shown in the figure. The overall torsion is zero while the overall hoop and axial stretch are as before, leading to the pressure-volume curve shown in 2.4(a), where volume strain is defined as the current volume divided by the reference volume.

Stripe domains are widely observed in nematic elastomers, especially in uniaxial tension, where rigid grips prevent any shear [77]. In uniaxial tension of a nematic sheet along an axis that is perpendicular to the initial director orientation, director rotation accommodates stretch but causes shear. However, shear breaks the symmetry and therefore there are two solutions (rotation to the right or left), which alternate to form the stripe domains. The domains are fine, typically with the width of microns, and the interfaces are very sharp, with a width of nanometers. The stripe domains in Figure 2.4(b) are the exact analogs of those in uniaxial tension.

## 2.4 Pump

The pressure-volume curves in Figure 2.4(a) motivate the application of this cylindrical nematic elastomer balloon as a pump. Recall that the anisotropy parameter $r$ depends on temperature, and therefore the four pressure-volume curves represent four distinct temperatures. In a typical monodomain nematic elastomer, $r = 2$ at a high temperature of about 85°C, while $r = 8$ at a low temperature of about 25°C [77]. These two pressure-volume curves are re-plotted in Figure 2.5 as the hot and cold nematic elastomers. An important observation is that the lower-critical pressure (point E) of the pressure-volume curve at the high temperature is higher than the upper-critical pressure (point B) of the pressure-volume curve at the low
Figure 2.4: (a) Pressure-volume relation for the inflation of a nematic cylinder. (b) Formation of stripe domains that avoid any torsion—the lines describe the director, while the dashed line is the deformed shape of a fiducial line that is initially straight and axial.

temperature. This enables the operation as a pump between an inlet pressure $p_i$ and outlet pressure $p_o$, where $p_f < p_i \leq p_o \leq p_E$.

Also shown in the figure are the isotherms (pressure-volume relation) of a fixed mass of fluid, in this case air, at the hot and the cold temperatures of $85^\circ C$ and $25^\circ C$, respectively. These isotherms were calculated using the ideal gas law. The cold fluid isotherm is given by $p_{\text{cold}}(V) = n_C R T_{\text{cold}} / V$, where $n_C$ is the number of moles of air at point C, $R = 8.3145$ is the ideal gas law constant, and $T_{\text{cold}}$ is the cold fluid temperature. The hot fluid isotherm pressure is analogously given by the corresponding number of moles at point D and the hot fluid temperature: $p_{\text{hot}}(V) = n_D R T_{\text{hot}} / V$.

The pump operates as follows. Let us begin at the high temperature with the outlet closed and the inlet open so that the nematic pump is at the point A. Now, cool the pump with the inlet open so that the pressure remains at $p_i$. On cooling fully to the cold temperature when $r = 8$, the only equilibrium solution is point C, which has a very large volume. So, the balloon would draw in a large volume of air from A to C. We note that this process does not proceed smoothly. As the pump is cooled from the high temperature, the volume changes gradually till the temperature when the upper critical pressure (the point corresponding to B at the intermediate
temperature) equals $p_i$. At this point, there will be an instability (likely accompanied by an aneurysm), and the volume jumps to something close to C. This instability has been analyzed by Giudici and Biggins [31], and may be of interest in microfluidics. Subsequent cooling takes it to point C.

Now close the inlet and start heating the pump. The mass of fluid in the pump is fixed, and so its behavior shifts from that of the cold isotherm to that of the hot isotherm. In the interim, the pressure-volume curve of the pump also changes to that of the hot material. Therefore, the equilibrium shifts from C to D. Now, open the outlet so that the pressure decreases to $p_o$. The only available state in the hot pump is at F, and so the pump goes from point D, with very large volume, to point F, with small volume, expelling the fluid. This is again accompanied by an instability from E to F. Closing the outlet and opening the inlet takes us from F to A, resetting the pump.

A pump can be characterized by its ejection fraction. In this case, the ejection fraction is

$$\frac{(\text{filled volume}) - (\text{empty volume})}{\text{filled volume}} = \frac{V_C - V_F}{V_C} = 98.6\%, \quad (2.19)$$
which means that 98.6% of the fluid is pumped out of a filled balloon during each cycle. This is extremely high: a normal human heart has a left ventricular ejection fraction between 50% and 70%, according to the American College of Cardiology [39]. A plot of ejection fraction for fixed \( r_0 = 2 \) and varying \( r \) can be seen in Figure 2.6.

### 2.5 Conclusion

We have introduced a modified formulation of the standard Warner-Terentjev energy density incorporated into a higher-order Ogden model to more accurately describe the behavior of nematic elastomers at large deformation. Furthermore, this work has initiated the study of actuation from geometries beyond flat, two-dimensional sheets by exploring a curvilinear three-dimensional geometry. We have outlined the deformation of a nematic elastomer balloon under simple expansion and twist. The material is actuated remotely by changing the temperature to dictate the degree of anisotropy, and the response is tunable. The foundation for our understanding of nematic elastomer actuation from flat geometries has already been well established with respect to the design, optimization, manufacturing, and tuning (e.g. voxelated sheets [76], wrinkling-resistant membranes [56], and moving inchworm [81]). Future applications based on more complex geometries and loading conditions, for instance incorporation of disclination defects and gradients of director or temperature across the thickness, can build upon this framework. Finally, the actuation can be effected by light instead of temperature [79]. For example, the pump described here would function with a light-driven actuation from \( r = 8 \) to \( r = 2 \). A practical
difficulty to be overcome is that many photo-active materials have a low penetration depth and the actuation is in bending, rather than stretching.
3.1 Introduction
The goal of this chapter is to understand how the soft elasticity of nematic elastomers affects complex inhomogeneous deformations. We have already seen in Chapter 2 that nematic elastomers can form fine-scale patterns called stripe domains. Stripe domains were first observed by Kundler and Finkelmann [42] in monodomains, nematic sheets subjected to uniaxial tension in a direction normal to the original director. These domains were associated with soft elasticity. Mathematically, the Bladon-Terentjev-Warner theory [10] leads to an energy that is not convex, and its relaxation leads to fine-scale structure and soft behavior [23]. This has been studied in homogeneous deformations (largely in uniaxial stretch [18, 19, 56], but more recently in biaxial stretch [85]), where we see different regimes depending on the imposed deformation. We expect these regimes to interact when the material is subjected to inhomogeneous deformation, and this motivates the current work.

We begin by describing the Bladon-Terentjev-Warner theory [10] and its relaxation due to Dolzmann and DeSimone [23]. The BTW theory uses a Gaussian approximation to treat the entropy of polymer chains. However, when polymer chains are subjected to large deformation, this approximation is no longer accurate as the polymer chains themselves are stretched, so we propose a generalization based on a Mooney-Rivlin energy. We then find its relaxation, which was independently done by Agosinian and DeSimone [1]. We use this relaxed energy to study a series of problems—the spherical balloon expansion, cylindrical balloon expansion, cavitation, and bending of a beam. The key idea here is to exploit the notion of universal deformations proposed by Ericksen [25]. He showed that there are certain deformations that automatically satisfy the equations of elasticity in all isotropic incompressible elastic bodies, and this is the basis of much work in finite elasticity.

3.2 Energy
Bladon-Terentjev-Warner theory
We begin by recalling the Bladon-Terentjev-Warner (BTW) theory for ideal nematic elastomers, introduced in Chapter 2. We pick a reference configuration to be the stress-free isotropic state and lower the temperature so that the material is in a
nematic state. The BTW theory adapts the Gaussian chain model to this situation, and shows that the free energy density is given by

\[
W^{BTW}(F, n) = \begin{cases} 
\frac{\mu}{2} \left[ \text{tr} \left( F^T \ell_n^{-1} F \right) - 3 \right] & \det F = 1, |n| = 1 \\
\infty & \text{else}
\end{cases}
\]

(3.1)

with step-length tensor \( \ell_n = r^{-1/3} (I + (r - 1)n \otimes n) \). The anisotropy parameter \( r \) describes the mesogen ordering: \( r = 1 \) represents the isotropic state, and \( r > 1 \) represents the anisotropic nematic state. Note that \( W^{BTW} = 0 \) if and only if \( F = Q\ell_0R, \hat{n} = Q\hat{e}_0 \) for \( Q, R \in SO(3), \hat{e}_0 \) a fixed unit vector, and \( \ell_0 = r^{1/3} (I + (r - 1)\hat{e}_0 \otimes \hat{e}_0) \).

**DeSimone-Dolzmann relaxation**

The BTW free energy is not convex, and this leads to fine-scale microstructure. DeSimone and Dolzmann computed the relaxation [23]. To do so, minimize over the nematic director \( n \) to obtain

\[
W(F) = \min_{n \text{ s.t. } |n| = 1} W^{BTW}(F, n) = \begin{cases} 
\frac{\mu}{2} \left[ r^{1/3} \left( \frac{s^2}{r} + \frac{r^2}{s^2} + \frac{1}{r^3} \right) - 3 \right] & \det F = 1, |n| = 1 \\
\infty & \text{else}
\end{cases},
\]

(3.2)

where \( s \) is the largest singular value of \( F \), and \( t \) is the largest singular value of \( \text{cof} \ F \). The relaxed energy is given by:

\[
W^{qc}(F) = \begin{cases} 
0 & F \in L \\
\frac{\mu}{2} \left( r^{1/3} + \frac{2t}{r^{1/6}} - 3 \right) & F \in M \\
w(F) & F \in S \\
\infty & \text{else}
\end{cases}.
\]

(3.3)

where \( L \) represents the liquid-like region, \( M \) represents the microstructure-formation region, and \( S \) represents the solid-like region:

\[
L : \{(s, t) : t \leq s^2, t \geq \sqrt{s}, t \leq r^{1/6}\}
\]

\[
M : \{(s, t) : t \geq r^{1/6}, t \leq s^2, t \geq r^{-1/2}s^2\} \quad (3.4)
\]

\[
S : \{(s, t) : t \geq \sqrt{s}, t \leq r^{-1/2}s^2\}
\]
We can rewrite these constraints in terms of the principal stretches of $F$ (details can be seen in the Appendix in Section A.4):

$$L : \{\lambda_{\text{max}}\lambda_{\text{mid}} \leq r^{1/6}\}$$

$$M : \{\lambda_{\text{max}}\lambda_{\text{mid}} \geq r^{1/6}, \frac{\lambda_{\text{max}}}{\lambda_{\text{mid}}} \leq \sqrt{r}\},$$

$$S : \{\frac{\lambda_{\text{max}}}{\lambda_{\text{mid}}} \geq \sqrt{r}\}$$

This describes the energy after the material has formed fine-scale microstructure. Figure 3.1 shows an illustration of the regions $L$, $M$, and $S$.

![Figure 3.1: Regions of $L$, $M$, and $S$ in the phase diagram of $(s,t)$.](image)

Physically, a liquid-like deformation in region $L$ is accommodated by unstressed microstructure in which the nematic director is in three dimensions (not confined to a plane). In region $M$, the deformation is accommodated by stressed microstructure in which the nematic director is planar (confined to a two-dimensional plane). For example, stripe domains are examples of microstructure that form in region $M$. The solid-like region $S$ corresponds to the stressed response of the polymer network without any liquid crystal molecule reorientation, for instance stretching a monodomain parallel to its nematic director.

One would expect that a nematic elastomer in region $L$ to be opaque due to the scattering of light because the nematic directors are reorienting in 3D. A nematic elastomer undergoing planar microstructure in region $M$ would be macroscopically opaque, but under polarized light microscopy there may be patterned features such as stripe domains detectable at the micrometer length scale. A sample in region $S$
would appear transparent, because the nematic mesogens would be fully aligned as a monodomain.

**Generalized Mooney-Rivlin energy**

This generalization and relaxation was independently proposed by Agostiniani and DeSimone [1]. Note that the trace formula is simply an extension of the neo-Hookean model for rubbers. This can be seen clearly when the energy is rewritten as such:

\[ W^{BTW}(F) = \frac{\mu}{2} \left[ \text{tr} (\tilde{F}^T \tilde{F}) - 3 \right], \]  

(3.6)

where

\[ \tilde{F} = \ell_n^{-1/2} F. \]  

(3.7)

Because the trace formula still relies on Gaussian chain modeling, it cannot model the stress build-up at large stretches. We will work to construct an energy density that is based on a generalized Mooney-Rivlin model.

Because a nematic elastomer is isotropic, our free energy density needs to be a function of the three invariants of the left Cauchy-Green tensor \( b \):

\[ W = f(I_1, I_2, I_3). \]  

(3.8)

The three invariants of the second-order tensor \( b = FF^T \) are

\[ I_1 = \text{tr} \ b = \text{tr} \ (FF^T) \]  

(3.9)

\[ I_2 = \frac{1}{2} \left[ (\text{tr} \ b)^2 - \text{tr} \ (b^2) \right] \]  

(3.10)

\[ I_3 = \det b = \det (FF^T) = (\det F)^2. \]  

(3.11)

Due to incompressibility of the material, the third invariant \( I_3 = 1 \), so

\[ W = g(I_1, I_2). \]  

(3.12)

We notice that the first invariant is a function of \( F \) and the second invariant is a function of \( \text{cof} \ F \):

\[ I_1 = \text{tr} \ b = \text{tr} (FF^T) \]  

(3.13)

\[ I_2 = \text{tr} \ b^{-1} = \text{tr} \ (\text{cof} b) = \text{tr} \ [(\text{cof} F) (\text{cof} F)^T]. \]  

(3.14)

Thus, we begin by looking at an energy based on the first invariant. For a rubber with deformation gradient \( F \),

\[ W_{H1}(F) = c |I_1 - 3|^p \]  

(3.15)

\[ = c |\text{tr}(FF^T) - 3|^p. \]  

(3.16)
For a nematic elastomer with deformation gradient \( F \),

\[
W_{I_1}(\tilde{F}) = W_{I_1}(\ell_n^{-1/2} F) = c \left| \text{tr} \left( \ell_n^{-1} b \right) - 3 \right|^p.
\]  

(3.17)

(3.18)

After minimizing the energy over the nematic director \( n \), we obtain:

\[
\min_{n \text{ s.t. } |n|=1} W_{I_1}(\ell_n^{-1/2} F) = c \left| r^{1/3} \left( \frac{s^2}{r} + \frac{t^2}{s^2} + \frac{1}{t^2} \right) - 3 \right|^p = c \left| W_1(s, t) \right|^p.
\]  

(3.19)

Details can be seen in the Appendix in Section A.2. Similarly, the energy based on the second invariant is as follows for rubbers:

\[
W_{I_2}(F) = d |I_2 - 3|^q
\]  

(3.20)

\[
= d \left| \text{tr} \left[ \left( \text{cof } F \right) \left( \text{cof } F \right)^\top \right] - 3 \right|^q.
\]  

(3.21)

For a nematic elastomer with deformation gradient \( F \),

\[
W_{I_2}(\tilde{F}) = W_{I_2}(\ell_n^{-1/2} F) = d \left| \text{tr} \left( \left( \ell_n^{-1} b \right)^{-1} \right) - 3 \right|^q.
\]  

(3.22)

(3.23)

After minimizing the energy over the nematic director \( n \), we obtain:

\[
\min_{n \text{ s.t. } |n|=1} W_{I_2}(\ell_n^{-1/2} F) = d \left| r^{-1/3} \left( \frac{r}{s^2} + \frac{s^2}{t^2} + \frac{1}{t^2} \right) - 3 \right|^q = d \left| W_2(s, t) \right|^q.
\]  

(3.24)

Details can be seen in the Appendix in Section A.3. Thus, we can create a generalized Mooney-Rivlin energy for an isotropic, incompressible nematic elastomer based on \( W_1(s, t) \) and \( W_2(s, t) \) as follows:

\[
W(s, t) = \begin{cases} 
\sum_{i=1}^{M} c_i \left| r^{1/3} \left( \frac{s^2}{r} + \frac{t^2}{s^2} + \frac{1}{t^2} \right) - 3 \right|^{p_i} & \text{det } F = 1 \\
+ \sum_{j=1}^{N} d_j \left| r^{-1/3} \left( \frac{r}{s^2} + \frac{s^2}{t^2} + \frac{t^2}{r^2} \right) - 3 \right|^{q_j} & \text{else}
\end{cases}
\]  

(3.25)

where \( s \) is the largest singular value of \( F \), \( t \) is the largest singular value of \( \text{cof } F \), and \( c_i \ (i = 1 : M) \) and \( d_j \ (j = 1 : N) \) are constants.
Relaxation of the generalized Mooney-Rivlin energy

Based on previous work done in this field [1, 65], the free energy density in Equation 3.25 is not convex in $s$ or $t$.

The relaxed form of this energy is:

$$W^{qc}(s, t) = \sum_{i=1}^{M} c_i (|W_1|^{p_i})^{qc} + \sum_{j=1}^{N} d_j (|W_2|^{q_j})^{qc}, \quad (3.26)$$

where

$$\left. \begin{array}{ll}
0 & F \in L \\
\frac{r^{1/3}}{r^2} + \frac{2t}{r^{1/6}} - 3 & F \in M \\
|W_1|^{p_i} & F \in S \\
\infty & \text{else}
\end{array} \right\} \quad (|W_1|^{p_i})^{qc}(s, t)$$

and

$$\left. \begin{array}{ll}
0 & F \in L \\
\frac{r^{-1/3}}{r^2} + \frac{2t^{1/6}}{r} - 3 & F \in M \\
|W_2|^{q_j} & F \in S \\
\infty & \text{else}
\end{array} \right\} \quad (|W_2|^{q_j})^{qc}(s, t). \quad (3.28)$$

The regions $L$, $M$, and $S$ are given by Equation 3.5, and the restrictions upon the exponents $p_i \geq 1 \ (i = 1 : M)$ and $q_j \geq 1 \ (j = 1 : N)$ must be satisfied.

### 3.3 Stress

Based on its principal values and directions, the left Cauchy-Green tensor is $b = \sum_{i=1}^{3} \lambda_i^2 \hat{v}_i \otimes \hat{v}_i$. If the strain energy density of a material can be written in the form $W = W(\lambda_i)$, where $\lambda_1 \geq \lambda_2 \geq \lambda_3$ are the principal stretches, then the Cauchy stress of an incompressible, isotropic hyperelastic body is

$$\sigma = -\eta I + \sum_{i=1}^{3} \lambda_i \frac{\partial W}{\partial \lambda_i} \hat{v}_i \otimes \hat{v}_i. \quad (3.29)$$

Here, the vectors $\hat{v}_i$ are normalized eigenvectors. We can derive the stresses for the relaxed generalized Mooney-Rivlin energy of Equation 3.26 in all three regions: $W = W^{qc}(s, t)$. In region $L$, the stresses are

$$\sigma^L = -\eta^L I. \quad (3.30)$$
In region $M$, $W^{qc}(s, t) = \sum_{i=1}^{M} c_i |A_M|^{p_i} + \sum_{j=1}^{N} d_j |B_M|^{q_j}$, where

$$A_M = r^{1/3} A_3^2 + \frac{2 \alpha_1 \alpha_2}{r^{1/6}} - 3$$

(3.31)

$$B_M = r^{-1/3} \alpha_1^2 \alpha_2^2 + 2 \lambda_3 r^{1/6} - 3.$$  

(3.32)

The principal Cauchy stresses are

$$\sigma^{M} = -\eta^{M} I + \sum_{i=1}^{3} \lambda_i \frac{\partial W^{qc}}{\partial \lambda_i} \hat{v}_i \otimes \hat{v}_i,$$

(3.33)

where

$$\lambda_1 \frac{\partial W^{qc}}{\partial \lambda_1} = \sum_{i=1}^{M} c_i p_i |A_M|^{p_i-1} 2 r^{-1/6} \lambda_1 \lambda_2 + \sum_{j=1}^{N} d_j q_j |B_M|^{q_j-1} 2 r^{-1/3} \lambda_1^2 \lambda_2^2$$  

(3.34)

$$\lambda_2 \frac{\partial W^{qc}}{\partial \lambda_2} = \lambda_1 \frac{\partial W^{qc}}{\partial \lambda_1}$$  

(3.35)

$$\lambda_3 \frac{\partial W^{qc}}{\partial \lambda_3} = \sum_{i=1}^{M} c_i p_i |A_M|^{p_i-1} 2 r^{1/3} \lambda_3^3 + \sum_{j=1}^{N} d_j q_j |B_M|^{q_j-1} 2 r^{1/6} \lambda_3.$$  

(3.36)

In region $S$, $W^{qc}(s, t) = \sum_{i=1}^{M} c_i |A_S|^{p_i} + \sum_{j=1}^{N} d_j |B_S|^{q_j}$, where

$$A_S = r^{1/3} A_3^2 + r^{1/3} \alpha_2^2 + r^{-2/3} \lambda_1^2 - 3$$  

(3.37)

$$B_S = r^{-1/3} \lambda_1^2 \alpha_2^2 + r^{-1/3} \lambda_1^2 \alpha_3^2 + r^{2/3} \lambda_2^2 \alpha_3^2 - 3.$$  

(3.38)

The principal Cauchy stresses are

$$\sigma^{S} = -\eta^{S} I + \sum_{i=1}^{3} \lambda_i \frac{\partial W^{qc}}{\partial \lambda_i} \hat{v}_i \otimes \hat{v}_i,$$

(3.39)

where

$$\lambda_1 \frac{\partial W^{qc}}{\partial \lambda_1} = \sum_{i=1}^{M} c_i p_i |A_S|^{p_i-1} 2 r^{-2/3} \lambda_1^2 + \sum_{j=1}^{N} d_j q_j |B_S|^{q_j-1} 2 r^{-1/3} \lambda_1^2 (\lambda_2^2 + \lambda_3^2)$$  

(3.40)

$$\lambda_2 \frac{\partial W^{qc}}{\partial \lambda_2} = \sum_{i=1}^{M} c_i p_i |A_S|^{p_i-1} 2 r^{1/3} \lambda_2^3 + \sum_{j=1}^{N} d_j q_j |B_S|^{q_j-1} 2 \lambda_2^3 (r^{-1/3} \lambda_1^2 + r^{2/3} \lambda_3^2)$$  

(3.41)

$$\lambda_3 \frac{\partial W^{qc}}{\partial \lambda_3} = \sum_{i=1}^{M} c_i p_i |A_S|^{p_i-1} 2 r^{1/3} \lambda_3^3 + \sum_{j=1}^{N} d_j q_j |B_S|^{q_j-1} 2 \lambda_3^2 (r^{-1/3} \lambda_1^2 + r^{2/3} \lambda_2^2).$$  

(3.42)
3.4 Ericksen’s “universal deformations"

J. L. Ericksen established the problem of determining all deformations which can be produced in every isotropic, incompressible, hyperelastic body by the application of surface tractions alone (no body forces) [25]. Because these universal relations are independent of constitutive relation, they are a powerful tool in continuum mechanics.

Below are the solution families known thus far for a point \((x, y, z)\), \((\rho, \theta, z)\), or \((\rho, \theta, \phi)\) with respective material point \((X, Y, Z)\), \((R, \Theta, Z)\), or \((R, \Theta, \Phi)\), and constant \(a, b, c, d, e, f\):

**Family 0**: Homogeneous deformations (for spatial point \(x\) with material point \(X\) and deformation gradient \(F\), constant vector \(c\))

\[
x = FX + c
\]  
(3.43)

**Family 1**: Bending, stretching, and shearing of a rectangular block

\[
\rho = \sqrt{2aX}, \theta = bY, z = \frac{Z}{ab} - bcY
\]  
(3.44)

**Family 2**: Straightening, stretching, and shearing of a sector of a tube

\[
x = \frac{1}{2}ab^2R^2, y = \frac{\Theta}{ab}, z = \frac{Z}{b} - \frac{c\Theta}{ab}
\]  
(3.45)

**Family 3**: Inflation, bending, torsion, extension, and shearing of an annular wedge, with \(a(cf - de) = 1\)

\[
\rho = \sqrt{aR^2 + b}, \theta = c\Theta + dZ, z = e\Theta + fZ
\]  
(3.46)

**Family 4**: Inflation or eversion of a sector of a spherical shell

\[
\rho = [\pm R^3 + a]^{1/3}, \theta = \pm\Theta, \phi = \Phi
\]  
(3.47)

**Family 5**: Inflation, bending, extension, and azimuthal shearing of an annular wedge [38, 66]

\[
\rho = a^{1/2}R, \theta = d\ln (bR) + c\Theta, z = eZ, ace = 1
\]  
(3.48)

In this work, we address three of the families, as applied to nematic elastomers: the bending of a block is a Family 1 deformation, the inflation of a cylindrical balloon is a Family 3 deformation, and the inflation of a spherical balloon and the cavitation of a disk are classified as Family 4 deformations.
3.5 Spherical balloon

Deformation of spherical balloon expansion

We are interested in the deformation of a balloon, which can be modeled as a spherical shell subjected to an internal pressure. In the undeformed configuration, the balloon has inner radius $R_i$ and outer radius $R_o$. The internal pressure is denoted by $p$. The spherical balloon is assumed to remain spherical throughout the deformation. The undeformed sphere has radial coordinate $R \in [R_i, R_o]$, azimuthal angle $\Theta \in [0, 2\pi)$, and polar angle $\Phi \in [0, \pi)$, while the deformed sphere has coordinate system $\rho \in [\rho_i, \rho_o]$, $\theta \in [0, 2\pi)$, and $\phi \in [0, \pi)$. Following Ericksen, we make the ansatz

$$\begin{align*}
\rho &= \rho(R) \\
\theta &= \Theta \\
\phi &= \Phi
\end{align*} \quad (3.49)$$

The deformation gradient in spherical coordinates is

$$F = \begin{pmatrix}
\frac{d\rho}{dR} & \frac{\rho}{R} & \frac{\rho}{R} \\
0 & \frac{\rho}{R} & 0 \\
0 & 0 & \frac{\rho}{R}
\end{pmatrix}. \quad (3.50)$$

With incompressibility, $\det F = 1$, we obtain this first-order differential equation

$$\frac{d\rho}{dR} \frac{\rho^2}{R^2} = 1. \quad (3.51)$$

Solving for the deformed radius $\rho$ as a function of the undeformed radius $R$:

$$\rho = \left( R^3 + c \right)^{1/3}, \quad (3.52)$$
where $c$ is a constant. Let $\lambda$ denote the azimuthal stretch, $\lambda = \frac{\rho}{R}$, and let $\lambda_o$ denote the azimuthal stretch at the outer radius, $\lambda_o = \frac{\rho(R_o)}{R_o}$. Then,

$$
\lambda_o = \left(\frac{R_o^3 + c}{R_o}\right)^{1/3}.
$$

(3.53)

Solving for $c$ and plugging $c$ back into Equation 3.52 yields

$$
\rho = \left(R^3 + R_o^3 \left(\lambda_o^3 - 1\right)\right)^{1/3},
$$

(3.54)

and the azimuthal stretch is

$$
\lambda = \left(1 + \left(\frac{R_o}{R}\right)^3 \left(\lambda_o^3 - 1\right)\right)^{1/3}.
$$

(3.55)

Thus, the deformation gradient is

$$
F = \begin{pmatrix}
\frac{1}{\lambda^2} \\
\lambda \\
\lambda
\end{pmatrix},
$$

(3.56)

and the left Cauchy-Green tensor, $b = FF^\top$ is

$$
b = \begin{pmatrix}
\frac{1}{\lambda^2} \\
\lambda^2 \\
\lambda^2
\end{pmatrix}.
$$

(3.57)

The principal stretches are $\lambda_1 = \lambda_2 = \lambda$ (corresponding to $e_\theta$ and $e_\phi$) and $\lambda_3 = \frac{1}{\lambda^2}$ (corresponding to $e_\rho$). Thus, $s = \lambda$ and $t = \lambda^2$. The regions are:

$$
L : \{ R \geq R^* \} 
$$

(3.58)

$$
M : \{ R \leq R^*, r \geq 1 \} 
$$

(3.59)

$$
S : \{ r \leq 1 \},
$$

(3.60)

where

$$
R^* = R_o \left(\frac{\lambda_o^3 - 1}{r^{1/4} - 1}\right)^{1/3}.
$$

(3.61)

This leads to the possibility of three cases:

- **Case 1**: $r > 1$ and $R^* \leq R_i$: the entire balloon is in region $L$
- **Case 2**: $r > 1$ and $R_i < R^* < R_o$: the inner region of the balloon $R \in [R_i, R^*]$ is in $M$, and the outer region $R \in [R^*, R_o]$ is in $L$
Figure 3.3: Diagram of all possible cases in the inflation of a nematic elastomer spherical balloon.

- **Case 3:** $r > 1$ and $R^* \geq R_\theta$: the entire balloon is in region $M$
- **Case 4:** $r = 1$ and the entire balloon is in region $S$

A diagram illustrating the various cases can be seen in Figure 3.3.

Note that Figure 3.1 can provide us with insight into the regions that this deformation will experience. The spherical balloon expansion is merely equibiaxial stretch (see the deformation gradient of Equation 3.56). Recalling that $s$ is the largest singular value of $F$ and $t$ is the largest singular value of $\text{cof} \ F$, this means that $t = s^2$ for this deformation. Following along the $t = s^2$ curve in Figure 3.1, we see that the deformation will move progressively through region $L$ then $M$, never touching $S$ for $r > 1$.

**Stress**

The expressions for stress are as follows: In region $L$,

$$\sigma^L = -\eta^L I. \quad (3.62)$$

In region $M$, the non-zero components of the stress are:

$$\sigma^M_{\rho \rho} = -\eta^M + \sum_{i=1}^{M} c_i p_i |A_M|^{p_i-1} \frac{2r^{1/3}}{\lambda^4} + \sum_{j=1}^{N} d_j q_j |B_M|^{q_j-1} \frac{2r^{1/6}}{\lambda^2}$$

$$\sigma^M_{\theta \theta} = -\eta^M + \sum_{i=1}^{M} c_i p_i |A_M|^{p_i-1} \frac{2\lambda^2}{r^{1/6}} + \sum_{j=1}^{N} d_j q_j |B_M|^{q_j-1} 2r^{-1/3} \lambda^4$$

$$\sigma^M_{\phi \phi} = \sigma^M_{\theta \theta}, \quad (3.63)$$

where

$$A_M = \frac{r^{1/3}}{\lambda^4} + \frac{2\lambda^2}{r^{1/6}} - 3$$

$$B_M = r^{-1/3} \lambda^4 + \frac{2r^{1/6}}{\lambda^2} - 3. \quad (3.64)$$
In region $S$, $r = 1$ and the non-zero components of the stress are

$$\sigma_{rr}^S = -\eta^S + \sum_{i=1}^{M} c_i A_S |\rho_i|^{p_i-1} \frac{2}{\lambda^4} + \sum_{j=1}^{N} d_j q_j |B_S|^{q_j-1} \frac{4}{\lambda^2}$$

$$\sigma_{\theta\theta}^S = -\eta^S + \sum_{i=1}^{M} c_i A_S |\rho_i|^{p_i-1} (2\lambda^2) + \sum_{j=1}^{N} d_j q_j |B_S|^{q_j-1} (2\lambda^2) \left( \lambda^2 + \frac{1}{\lambda^4} \right)$$

$$\sigma_{\phi\phi}^S = \sigma_{\theta\theta}^S,$$

where

$$A_S = \frac{1}{\lambda^4} + 2\lambda^2 - 3$$

$$B_S = \lambda^4 + \frac{2}{\lambda^2} - 3.$$  \hspace{1cm} (3.65)

### Solving static equilibrium

The static equilibrium equations (in the absence of body forces) in spherical coordinates are

$$\rho \frac{\partial \sigma_{\rho\rho}}{\partial \rho} + 2 \sigma_{\rho\rho} - \frac{2}{\rho} \frac{\partial \sigma_{\rho\phi}}{\partial \phi} + \frac{1}{\rho} \frac{\partial \sigma_{\rho\phi}}{\partial \rho} + \frac{1}{\rho} \frac{\partial \sigma_{\phi\phi}}{\partial \phi} + \frac{1}{\rho} \sin \phi \frac{\partial \sigma_{\theta\theta}}{\partial \theta} - \frac{1}{\rho} \left( \sigma_{\theta\theta} + \sigma_{\phi\phi} \right) = 0$$

$$\theta \frac{\partial \sigma_{\rho\theta}}{\partial \rho} + 2 \sigma_{\rho\theta} - \frac{2}{\rho} \frac{\partial \sigma_{\rho\phi}}{\partial \phi} + \frac{1}{\rho} \frac{\partial \sigma_{\rho\phi}}{\partial \rho} + \frac{1}{\rho} \frac{\partial \sigma_{\theta\theta}}{\partial \theta} + \frac{\sigma_{\theta\theta}}{\rho} \sin \phi \frac{\partial \phi}{\partial \theta} + \frac{\cot \phi}{\rho} \left( \sigma_{\phi\phi} + \sigma_{\theta\phi} \right) = 0$$

$$\phi \frac{\partial \sigma_{\rho\phi}}{\partial \rho} + 2 \sigma_{\rho\phi} - \frac{2}{\rho} \frac{\partial \sigma_{\rho\phi}}{\partial \phi} + \frac{1}{\rho} \frac{\partial \sigma_{\rho\phi}}{\partial \rho} + \frac{1}{\rho} \frac{\partial \sigma_{\phi\phi}}{\partial \theta} + \frac{\sigma_{\phi\phi}}{\rho} \sin \phi \frac{\partial \theta}{\partial \theta} + \frac{\cot \phi}{\rho} \left( \sigma_{\phi\phi} - \sigma_{\theta\phi} \right) = 0,$$

and the boundary conditions for this problem are as follows:

$$\sigma_{\rho\rho}|_{\rho=\rho_i} = -p$$  \hspace{1cm} (3.67)

$$\sigma_{\rho\rho}|_{\rho=\rho_o} = 0.$$  \hspace{1cm} (3.68)

In all cases, solving the $\phi$ equation yields $\eta = \eta (\rho, \theta)$. Similarly, solving the $\theta$ equation, we find that $\eta = \eta (\rho)$. Finally, to solve the $\rho$ equation, we will rewrite the stress expressions as

$$\sigma_{\rho\rho} = -\eta + \hat{\sigma}_{\rho\rho}$$

$$\sigma_{\theta\theta} = \sigma_{\phi\phi} = -\eta + \hat{\sigma}_{\theta\theta},$$

and so the boundary conditions can be rewritten as follows

$$-\eta(\rho = \rho_i) + \hat{\sigma}_{\rho\rho}(\rho = \rho_i) = -p$$

$$-\eta(\rho = \rho_o) + \hat{\sigma}_{\rho\rho}(\rho = \rho_o) = 0.$$
Case 1

In Case 1, the entire balloon is in region $L$. The two boundary conditions for this case are:

$$-\eta^L(\rho = \rho_i) + \hat{\sigma}^L_{\rho\rho}(\rho = \rho_i) = -p$$  \hspace{1cm} (3.76)

$$-\eta^L(\rho = \rho_o) + \hat{\sigma}^L_{\rho\rho}(\rho = \rho_o) = 0.$$  \hspace{1cm} (3.77)

The stress in this region is found in Equation 3.62. Thus, the $\rho$ equilibrium equation yields

$$\frac{d\sigma^L_{\rho\rho}}{dp} + 2\frac{\sigma^L_{\rho\rho}}{\rho} - \frac{1}{\rho} \left( \sigma^L_{\theta\theta} + \sigma^L_{\phi\phi} \right) = 0 \tag{3.78}$$

$$\frac{d(-\eta^L + \hat{\sigma}^L_{\rho\rho})}{dp} + 2\frac{(-\eta^L + \hat{\sigma}^L_{\rho\rho})}{\rho} - \frac{1}{\rho} \left( -\eta^L + \hat{\sigma}^L_{\theta\theta} - \eta^L + \hat{\sigma}^L_{\phi\phi} \right) = 0 \hspace{1cm} (3.79)$$

$$\int_{\rho_i}^{\rho_o} d \left( -\eta^L + \hat{\sigma}^L_{\rho\rho} \right) = \int_{\rho_i}^{\rho_o} 0 dp \hspace{1cm} (3.80)$$

Thus, for Case 1, the inner pressure is:

$$p = 0.$$ \hspace{1cm} (3.81)

Case 2

In Case 2, $R \in [R_i, R^\ast]$ is in region $M$, and the $R \in [R^\ast, R_o]$ is in $L$. The two boundary conditions can be rewritten specific to the region:

$$-\eta^M(\rho = \rho_i) + \hat{\sigma}^M_{\rho\rho}(\rho = \rho_i) = -p \hspace{1cm} (3.83)$$

$$-\eta^L(\rho = \rho_o) + \hat{\sigma}^L_{\rho\rho}(\rho = \rho_o) = 0,$$ \hspace{1cm} (3.84)

and there is an additional boundary condition for continuity between the two regions:

$$-\eta^L(\rho = \rho^\ast) + \hat{\sigma}^L_{\rho\rho}(\rho = \rho^\ast) = -\eta^M(\rho = \rho^\ast) + \hat{\sigma}^M_{\rho\rho}(\rho = \rho^\ast).$$ \hspace{1cm} (3.85)
First, we solve the equilibrium equations in region \( L \). Again, the stress in region \( L \) is found in Equation 3.62. The \( \rho \) equation yields:

\[
\frac{d\sigma^L_{\rho\rho}}{d\rho} + 2\frac{\sigma^L_{\rho\rho}}{\rho} - \frac{1}{\rho}\left(\sigma^L_{\theta\theta} + \sigma^L_{\phi\phi}\right) = 0 \tag{3.86}
\]

\[
\frac{d(-\eta^L + \hat{\sigma}^L_{\rho\rho})}{d\rho} + 2\frac{(-\eta^L + \hat{\sigma}^L_{\rho\rho})}{\rho} - \frac{1}{\rho}\left(-\eta^L + \hat{\sigma}^L_{\theta\theta} - \eta^L + \hat{\sigma}^L_{\phi\phi}\right) = 0 \tag{3.87}
\]

\[
\frac{d(-\eta^L + \hat{\sigma}^L_{\rho\rho})}{d\rho} = 0 \tag{3.88}
\]

\[
\left(-\eta^L(\rho_o) + \hat{\sigma}^L_{\rho\rho}(\rho_o)\right) - \left(-\eta^L(\rho^*) + \hat{\sigma}^L_{\rho\rho}(\rho^*)\right) = 0 \tag{3.89}
\]

In region \( M \), the stress is that of Eqns. 3.63 and 3.64. The \( \rho \) equation yields:

\[
\frac{d\sigma^M_{\rho\rho}}{d\rho} + 2\frac{\sigma^M_{\rho\rho}}{\rho} - \frac{1}{\rho}\left(\sigma^M_{\theta\theta} + \sigma^M_{\phi\phi}\right) = 0 \tag{3.92}
\]

\[
\frac{d(-\eta^M + \hat{\sigma}^M_{\rho\rho})}{d\rho} + \frac{1}{\rho}\left(2\hat{\sigma}^M_{\rho\rho} - \hat{\sigma}^M_{\theta\theta} - \hat{\sigma}^M_{\phi\phi}\right) = 0 \tag{3.93}
\]

\[
\int_{\rho_i}^{\rho^*} d\left(-\eta^M + \hat{\sigma}^M_{\rho\rho}\right) = -\int_{\rho_i}^{\rho^*} \frac{2}{\rho}\left(\hat{\sigma}^M_{\rho\rho} - \hat{\sigma}^M_{\theta\theta}\right) d\rho \tag{3.94}
\]

\[
\left(-\eta^M(\rho^*) + \hat{\sigma}^M_{\rho\rho}(\rho^*)\right) - \left(-\eta^M(\rho_i) + \hat{\sigma}^M_{\rho\rho}(\rho_i)\right) = \int_{\rho_i}^{\rho^*} \frac{2}{\rho}\left(\hat{\sigma}^M_{\rho\rho} - \hat{\sigma}^M_{\theta\theta}\right) d\rho \tag{3.95}
\]

\[
p = \int_{\rho_i}^{\rho^*} \frac{2}{\rho}\left(\hat{\sigma}^M_{\rho\rho} - \hat{\sigma}^M_{\theta\theta}\right) d\rho. \tag{3.96}
\]

The right-hand side requires a change of integration variable from \( \rho \) to \( R \). Noting that \( d\rho = \frac{1}{\lambda(R)} dR \), we obtain an expression for the inner pressure:

\[
p = \int_{R_i}^{R^*} \frac{2}{\lambda(R)^3} \left[\hat{\sigma}^M_{\theta\theta}(\rho = \lambda(R)R) - \hat{\sigma}^M_{\rho\rho}(\rho = \lambda(R)R)\right] dR. \tag{3.97}
\]

**Case 3**

In Case 3, all of the balloon is in region \( M \). The two boundary conditions for this case are:

\[
-\eta^M(\rho = \rho_i) + \hat{\sigma}^M_{\rho\rho}(\rho = \rho_i) = -p \tag{3.98}
\]

\[
-\eta^M(\rho = \rho_o) + \hat{\sigma}^M_{\rho\rho}(\rho = \rho_o) = 0. \tag{3.99}
\]
The stress in region $M$ can be found in Eqns. 3.63 and 3.64. The $\rho$ equilibrium equation yields:

$$\frac{d\sigma_{\rho\rho}^M}{d\rho} + 2\frac{\sigma_{\rho\rho}^M}{\rho} - \frac{1}{\rho} \left( \sigma_{\theta\theta}^M + \sigma_{\phi\phi}^M \right) = 0$$

(3.100)

$$\frac{d(-\eta^M + \hat{\sigma}_{\rho\rho}^M)}{d\rho} + \frac{1}{\rho} \left( 2\hat{\sigma}_{\rho\rho}^M - \hat{\sigma}_{\theta\theta}^M - \hat{\sigma}_{\phi\phi}^M \right) = 0$$

(3.101)

$$\int_{\rho_i}^{\rho_o} d\left(-\eta^M + \hat{\sigma}_{\rho\rho}^M\right) = -\int_{\rho_i}^{\rho_o} \frac{2}{\rho} \left( \hat{\sigma}_{\rho\rho}^M - \hat{\sigma}_{\theta\theta}^M \right) d\rho$$

(3.102)

$$\left(-\eta^M(\rho_o) + \hat{\sigma}_{\rho\rho}(\rho_o)\right) - \left(-\eta^M(\rho_i) + \hat{\sigma}_{\rho\rho}(\rho_i)\right) = \int_{\rho_i}^{\rho_o} \frac{2}{\rho} \left( \hat{\sigma}_{\theta\theta}^M - \hat{\sigma}_{\rho\rho}^M \right) d\rho$$

(3.103)

$$p = \int_{\rho_i}^{\rho_o} \frac{2}{\rho} \left( \hat{\sigma}_{\rho\rho}^M - \hat{\sigma}_{\rho\rho}^M \right) d\rho.$$  

(3.104)

As in Case 2, we employ a change of integration variable from $\rho$ to $R$ and obtain the inner pressure as:

$$p = \int_{R_i}^{R_o} \frac{2}{\lambda(R)^3} \left[ \hat{\sigma}_{\theta\theta}(\rho = \lambda(R)R) - \hat{\sigma}_{\rho\rho}^M(\rho = \lambda(R)R) \right] dR.$$  

(3.105)

**Case 4**

In Case 4, all of the balloon is in region $S$. Note that only the isotropic state ($r = 1$) falls under this case. The two boundary conditions specific to this case are

$$-\eta^S(\rho = \rho_i) + \hat{\sigma}_{\rho\rho}^S(\rho = \rho_i) = -p$$

(3.106)

$$-\eta^S(\rho = \rho_o) + \hat{\sigma}_{\rho\rho}^S(\rho = \rho_o) = 0.$$  

(3.107)
The stress in region \( S \) can be found in Eqns. 3.65 and 3.66. The \( \rho \) equilibrium equation yields:

\[
\frac{d\sigma_{\rho\rho}^M}{d\rho} + 2 \frac{\sigma_{\rho\rho}^M}{\rho} - \frac{1}{\rho} \left( \sigma_{\theta\theta}^M + \sigma_{\phi\phi}^M \right) = 0
\]

(3.108)

\[
\frac{d(-\eta^M + \hat{\sigma}_{\rho\rho}^M)}{d\rho} + \frac{1}{\rho} \left( 2\hat{\sigma}_{\rho\rho}^M - \hat{\sigma}_{\theta\theta}^M - \hat{\sigma}_{\phi\phi}^M \right) = 0
\]

(3.109)

\[
\int_{\rho_i}^{\rho_o} d \left( -\eta^M + \hat{\sigma}_{\rho\rho}^M \right) = - \int_{\rho_i}^{\rho_o} \frac{2}{\rho} \left( \hat{\sigma}_{\theta\theta}^M - \hat{\sigma}_{\rho\rho}^M \right) d\rho
\]

(3.110)

\[
\left( -\eta^M(\rho_o) + \hat{\sigma}_{\rho\rho}^M(\rho_o) \right) - \left( -\eta^M(\rho_i) + \hat{\sigma}_{\rho\rho}^M(\rho_i) \right) = \int_{\rho_i}^{\rho_o} \frac{2}{\rho} \left( \hat{\sigma}_{\theta\theta}^M - \hat{\sigma}_{\rho\rho}^M \right) d\rho
\]

(3.111)

\[
p = \int_{\rho_i}^{\rho_o} \frac{2}{\rho} \left( \hat{\sigma}_{\theta\theta}^M - \hat{\sigma}_{\rho\rho}^M \right) d\rho.
\]

(3.112)

As in Case 2, we employ a change of integration variable from \( \rho \) to \( R \) and obtain the inner pressure as:

\[
p = \int_{R_i}^{R_o} \frac{2}{\lambda(R)^3 R} \left[ \hat{\sigma}_{\theta\theta}^M(\rho = \lambda(R)R) - \hat{\sigma}_{\rho\rho}^M(\rho = \lambda(R)R) \right] dR.
\]

(3.113)

**Results**

The calculations were performed in MATLAB. The solutions are plotted in Figures 3.4–3.5b with the inner radius \( R_i = 1 \) cm and outer radius \( R_o = 1.1 \) cm. The following parameters for the generalized Mooney-Rivlin model were used: \( M = 2, N = 1, c_1 = 1.0 \cdot 10^5 \) Pa, \( c_2 = 1.90 \cdot 10^2 \) Pa, \( d_1 = 1.59 \cdot 10^{-2} \) Pa, \( p_1 = 1.3, p_2 = 5, \) and \( q_1 = 2. \) This yields an effective shear modulus \( \mu = \frac{1}{2} \left( \sum_{i=1}^{N} c_i p_i + \sum_{j=1}^{M} d_j q_j \right) = 6.52 \cdot 10^4 \) Pa.

The pressure \( p \), normalized by the effective shear modulus \( \mu \), is plotted as a function of the azimuthal stretch at the outer radius \( \lambda_o \). Figure 3.4 shows the comparison between the generalized Mooney-Rivlin model of this work and the BTW (neo-Hookean) model of previous work in the field [23] for an anisotropy parameter of \( r = 8 \). The neo-Hookean model is unable to capture the correct effect of elasticity in the material at very high values of stretch (e.g. for \( \lambda_o \) greater than about 6). As seen in experiments of rubber balloons [73], the balloon experiences a subsequent
Figure 3.4: Comparison of this work’s generalized Mooney-Rivlin model with the trace formula model of [23] for inflation pressure of a spherical balloon.

stiffening due to the further stretching of the polymer, or “effects of the limited extensibility of the network”, as indicated by the increase in pressure at high stretch, which is correctly captured by our generalized Mooney-Rivlin model.

Figure 3.5: Spherical balloon results: (a) Pressure-stretch curves at varying anisotropy parameter. (b) Progression of the spherical balloon solution through individual case numbers.

Figure 3.5a shows the results for varying anisotropy parameter $r$. The response of the balloon is stiffest in the isotropic state ($r = 1$), and gets correspondingly softer as $r$ increases, as expected. As the pressure $p$ increases, the balloon undergoes deformation according to the various cases, as seen in Figure 3.5b. For $r = 1$ (rubber), the balloon is in Case 1 (region $L$) at $\lambda_o = 1$ when there is no deformation, but for the rest of the deformation the balloon is entirely in region $S$, corresponding to a solid-like response with no microstructure formation, since the material is a rubber with no liquid crystals. For $r > 1$, the solutions move progressively from Case 1.
to 2 to 3 throughout the deformation. At a stretch of \( \lambda_o = 1 \), the balloon is again in Case 1, corresponding to being entirely in the \( L \) region because no deformation has occurred. Immediately upon inflation of the balloon, the balloon jumps to Case 2, where the inner part of the balloon experiences region \( M \), developing fine-scale microstructure. Then shortly after, the balloon will become entirely in region \( M \), and the rest of the balloon will develop fine-scale microstructure in response to the pressure. This formation of microstructure by the liquid crystal molecules creates a softer response than the rubber without liquid crystals.

### 3.6 Cylindrical balloon

#### Deformation of cylindrical balloon expansion

We are interested in the deformation of a cylindrical shell composed of the nematic elastomer, capped off at both ends, subjected to an internal pressure. The shell has undeformed height \( H \), inner radius \( R_i \), and undeformed outer radius \( R_o \). Note that there is no thin-wall approximation used in this formulation. The internal pressure is denoted by \( p \), as seen in Figure 3.6a. The cylindrical shell is assumed to deform uniformly, i.e. the cylinder remains a cylinder throughout the deformation. The undeformed cylinder has coordinate system \( R, \Theta, \) and \( Z \), while the deformed cylinder has coordinate system \( \rho, \theta, \) and \( z \), as seen in Figure 3.6b. The liquid crystals are in an isotropic reference state and free to move throughout the deformation. The nematic elastomer is assumed to be incompressible.

![Figure 3.6: (a) Cross-section of cylindrical shell. (b) Schematic showing cylindrical coordinates.](image)

We assume that the cylinder remains a cylinder throughout the deformation, so the mapping that describes this progression is \( \varphi = \rho e_\rho + z e_z \). Following Ericksen, we
look for a solution of the form
\[
\begin{cases}
\rho = \rho(R) \\
\theta = \Theta \\
z = \xi \end{cases}
\] (3.114)

Following the convention found in the paper by Rudykh et al. on electroactive balloons [63], let \( \lambda = r/R \) be the hoop stretch ratio. Let \( \xi \) denote the axial stretch ratio. Thus the associated deformation gradient is:
\[
\mathbf{F} = \begin{pmatrix}
\frac{1}{\lambda\xi} & 0 & 0 \\
0 & \lambda & 0 \\
0 & 0 & \xi
\end{pmatrix},
\] (3.115)

where
\[
\lambda = \sqrt{\frac{1}{\xi} + \frac{R_o^2}{R^2} \left( \lambda_o^2 - \frac{1}{\xi} \right)}.
\] (3.116)

For fixed axial ratio \( \xi = 1 \), the principal stretches \( \lambda_1 \geq \lambda_2 \geq \lambda_3 \) are \( \lambda_1 = \lambda \), corresponding to \( \hat{e}_\theta \), \( \lambda_2 = 1 \), corresponding to \( \hat{e}_z \), and \( \lambda_3 = \frac{1}{\lambda} \), corresponding to \( \hat{e}_\rho \).

Simplifying the regions, we find:
\[
L : R \geq R_2
\] (3.117)
\[
M : R_1 \leq R \leq R_2
\] (3.118)
\[
S : R \leq R_1,
\] (3.119)

where \( R_1 = \sqrt{\frac{R_o^2(\lambda_o^2 - 1)}{r - 1}} \) and \( R_2 = \sqrt{\frac{R_o^2(\lambda_o^2 - 1)}{r^{1/3} - 1}} \).

This leads to the possibility of the following cases:

- **Case 0**: \( r > 1 \) and \( R_1 \leq R \leq R_i \): the entire balloon is in region \( L \)
- **Case 1**: \( r > 1 \) and \( R_1 \leq R_i < R_2 < R_o \): the inner portion of the balloon is in region \( M \) and the outer portion is in \( L \)
- **Case 2**: \( r > 1 \) and \( R_1 \leq R_i < R_o \leq R_2 \): the entire balloon is in region \( M \)
- **Case 3**: \( r > 1 \) and \( R_i < R_1 < R_o \leq R_2 \): the inner portion of the balloon is in region \( S \) and the outer portion is in \( M \)
- **Case 4**: \( r > 1 \) and \( R_i < R_1 < R_2 < R_o \): the balloon is in regions \( S \), then \( M \), then \( L \) from inside to outside
- **Case 5**: \( r \geq 1 \) and \( R_o \leq R_1 \leq R_2 \): the entire balloon is in region \( S \)
A diagram illustrating the various cases can be seen in Figure 3.7.

As in the spherical balloon deformation, we can gain insight from Figure 3.1 regarding the regions that this deformation will experience. The cylindrical balloon expansion with fixed axial ration $\xi = 1$ is the same as a planar extension deformation (the deformation gradient of Equation 3.115 is $F = \text{diag}(1/\lambda, \lambda, 1)$). Recalling that $s$ is the largest singular value of $F$ and $t$ is the largest singular value of $\text{cof} F$, this means that $t = s$ for this deformation. If we were to follow along the $t = s$ line in Figure 3.1, we see that the deformation will move progressively through region $L$ then $M$ then $S$.

**Stress**

The stress in region $L$ is

$$\sigma^L = -\eta^L \mathbf{I}. \quad (3.120)$$

The non-zero components of stress in region $M$ are

$$\sigma^M_{\rho\rho} = -\eta^M + \sum_{i=1}^{M} c_i p_i |A_M|^{p_i - 1} \frac{r^{1/3}}{\lambda^2} + \sum_{j=1}^{N} d_j q_j |B_M|^{q_j - 1} \frac{2r^{1/6}}{\lambda}$$

$$\sigma^M_{\theta\theta} = -\eta^M + \sum_{i=1}^{M} c_i p_i |A_M|^{p_i - 1} \frac{2\lambda}{r^{1/6}} + \sum_{j=1}^{N} d_j q_j |B_M|^{q_j - 1} 2r^{-1/3} \lambda^2$$

$$\sigma^M_{zz} = \sigma^M_{\theta\theta}, \quad (3.121)$$

where

$$A_M = \frac{r^{1/3}}{\lambda^2} + \frac{2\lambda}{r^{1/6}} - 3$$

$$B_M = r^{-1/3} \lambda^2 + \frac{2r^{1/3}}{\lambda} - 3. \quad (3.122)$$
The non-zero components of stress in region S are

\[
\sigma_{\rho\rho}^S = -\eta^S + \sum_{i=1}^{M} c_i \rho_i |A_S|^{\rho_i - 1/2} r^{1/3} + \sum_{j=1}^{N} d_j q_j |B_S|^{q_j - 1/2} (r^{-1/3} \lambda^2 + r^{2/3})
\]

\[
\sigma_{\theta\theta}^S = -\eta^S + \sum_{i=1}^{M} c_i \rho_i |A_S|^{\rho_i - 1/2} \lambda^2 r^{-2/3} + \sum_{j=1}^{N} d_j q_j |B_S|^{q_j - 1/2} (2r^{-1/3} \lambda^2 + 1 + \frac{1}{\lambda^2})
\]

\[
\sigma_{zz}^S = -\eta^S + \sum_{i=1}^{M} c_i \rho_i |A_S|^{\rho_i - 1/2} r^{1/3} + \sum_{j=1}^{N} d_j q_j |B_S|^{q_j - 1/2} \left(2r^{-1/3} \lambda^2 + 2 \frac{r^{2/3}}{\lambda^2}\right),
\]

(3.123)

where

\[
A_S = \frac{r^{1/3}}{\lambda^2} + r^{1/3} + r^{-2/3} \lambda^2 - 3
\]

(3.124)

\[
B_S = r^{-1/3} \lambda^2 + r^{-1/3} + \frac{r^{2/3}}{\lambda^2} - 3.
\]

**Solving static equilibrium**

Static equilibrium (in the absence of body forces) is obtained when \( \text{div} \, \sigma = 0 \). In cylindrical coordinates and for a symmetric tensor \( \sigma \),

\[
\frac{\partial \sigma_{\rho\rho}}{\partial \rho} + \frac{1}{\rho} \frac{\partial \sigma_{\rho\theta}}{\partial \theta} + \frac{\sigma_{\rho\rho} - \sigma_{\theta\theta}}{\rho} + \frac{\partial \sigma_{\rho z}}{\partial z} = 0,
\]

\[
\frac{\partial \sigma_{\rho\theta}}{\partial \rho} + \frac{1}{\rho} \frac{\partial \sigma_{\theta\theta}}{\partial \theta} + \frac{2\sigma_{\rho\theta}}{\rho} + \frac{\partial \sigma_{\theta z}}{\partial z} = 0,
\]

\[
\frac{\partial \sigma_{\rho z}}{\partial \rho} + \frac{1}{\rho} \frac{\partial \sigma_{\theta z}}{\partial \theta} + \frac{\sigma_{\rho z}}{\rho} + \frac{\partial \sigma_{zz}}{\partial z} = 0.
\]

(3.125)

The boundary conditions for this problem are as follows:

\[
\sigma_{\rho\rho} |_{\rho = \rho_i} = -p
\]

(3.126)

\[
\sigma_{\rho\rho} |_{\rho = \rho_o} = 0.
\]

(3.127)

For every case, the \( \theta \) and \( z \) equilibrium equations yield that \( \eta = \eta(\rho) \). This leaves only the \( \rho \) equilibrium equation to be solved. We will rewrite the stress expressions as

\[
\sigma_{\rho\rho} = -\eta + \hat{\sigma}_{\rho\rho}
\]

(3.128)

\[
\sigma_{\theta\theta} = -\eta + \hat{\sigma}_{\theta\theta}.
\]

(3.129)
Case 0

In Case 0, the entire balloon is in region $L$. The two boundary conditions for this case are:

$$-\eta^L(\rho = \rho_i) + \hat{\sigma}^L_{\rho \rho}(\rho = \rho_i) = -p$$  \hspace{1cm} (3.130)

$$-\eta^L(\rho = \rho_o) + \hat{\sigma}^L_{\rho \rho}(\rho = \rho_o) = 0.$$  \hspace{1cm} (3.131)

The $\rho$ equilibrium equation yields

$$\frac{d\sigma^L_{\rho \rho}}{d\rho} + \frac{1}{\rho} \left( \sigma^L_{\rho \rho} - \sigma^L_{\theta \theta} \right) = 0$$  \hspace{1cm} (3.132)

$$\frac{d(-\eta^L + \hat{\sigma}^L_{\rho \rho})}{d\rho} + \frac{1}{\rho} \left( \hat{\sigma}^L_{\rho \rho} - \hat{\sigma}^L_{\theta \theta} \right) = 0$$  \hspace{1cm} (3.133)

$$\int_{\rho_i}^{\rho_o} d \left( -\eta^L + \hat{\sigma}^L_{\rho \rho} \right) = \int_{\rho_i}^{\rho_o} 0 d\rho$$  \hspace{1cm} (3.134)

$$p = 0,$$  \hspace{1cm} (3.135)

and thus, for Case 1, the inner pressure is:

$$p = 0.$$  \hspace{1cm} (3.136)

Case 1

In Case 1, $R \in [R_i, R_2]$ is in region $M$, and the $R \in [R_2, R_o]$ is in $L$. The two boundary conditions can be rewritten specific to the region:

$$-\eta^M(\rho = \rho_i) + \hat{\sigma}^M_{\rho \rho}(\rho = \rho_i) = -p$$  \hspace{1cm} (3.137)

$$-\eta^L(\rho = \rho_o) + \hat{\sigma}^L_{\rho \rho}(\rho = \rho_o) = 0,$$  \hspace{1cm} (3.138)

and there is an additional boundary condition for continuity between the two regions:

$$- \eta^L(\rho = \rho_2) + \hat{\sigma}^L_{\rho \rho}(\rho = \rho_2) = -\eta^M(\rho = \rho_2) + \hat{\sigma}^M_{\rho \rho}(\rho = \rho_2).$$  \hspace{1cm} (3.139)

First, we solve the equilibrium equations in region $L$. The $\rho$ equation yields:

$$\frac{d\sigma^L_{\rho \rho}}{d\rho} + \frac{1}{\rho} \left( \sigma^L_{\rho \rho} - \sigma^L_{\theta \theta} \right) = 0$$  \hspace{1cm} (3.140)

$$\frac{d(-\eta^L + \hat{\sigma}^L_{\rho \rho})}{d\rho} = 0$$  \hspace{1cm} (3.141)

$$\int_{\rho_2}^{\rho_o} d \left( -\eta^L + \hat{\sigma}^L_{\rho \rho} \right) = \int_{\rho_2}^{\rho_o} 0 d\rho$$  \hspace{1cm} (3.142)

$$\left( -\eta^L + \hat{\sigma}^L_{\rho \rho} \right)_{\rho_2}^{\rho_o} = 0$$  \hspace{1cm} (3.143)

$$\eta^M(\rho_2) - \hat{\sigma}^M_{\rho \rho}(\rho_2) = 0.$$  \hspace{1cm} (3.144)
In region M, the \( \rho \) equation yields:

\[
\frac{d\sigma_{\rho \rho}^M}{d\rho} + \frac{1}{\rho} \left( \sigma_{\rho \rho}^M - \sigma_{\theta \theta}^M \right) = 0
\]  \( (3.145) \)

\[
\frac{d(-\eta^M + \hat{\sigma}_{\rho \rho}^M)}{d\rho} + \frac{1}{\rho} \left( \hat{\sigma}_{\rho \rho}^M - \hat{\sigma}_{\theta \theta}^M \right) = 0
\]  \( (3.146) \)

\[
\int_{\rho_i}^{\rho_2} d \left( -\eta^M + \hat{\sigma}_{\rho \rho}^M \right) = \int_{\rho_i}^{\rho_2} \frac{1}{\rho} \left( \hat{\sigma}_{\theta \theta}^M - \hat{\sigma}_{\rho \rho}^M \right) d\rho
\]  \( (3.147) \)

\[
p = \int_{\rho_i}^{\rho_2} \frac{1}{\rho} \left( \hat{\sigma}_{\theta \theta}^M - \hat{\sigma}_{\rho \rho}^M \right) d\rho.
\]  \( (3.148) \)

The right-hand side requires a change of integration variable from \( \rho \) to \( R \). Noting that \( d\rho = \frac{1}{\lambda(R)^2} dR \), we obtain an expression for the inner pressure:

\[
p = \int_{\rho_i}^{\rho_2} \frac{1}{\lambda(R)^3} \left[ \hat{\sigma}_{\theta \theta}^M (\rho = \lambda(R)R) - \hat{\sigma}_{\rho \rho}^M (\rho = \lambda(R)R) \right] dR.
\]  \( (3.149) \)

**Case 2**

In Case 2, \( R \in [R_i, R_o] \) is in region M. The boundary conditions are:

\[
-\eta(\rho_i) + \hat{\sigma}_{\rho \rho}^M(\rho_i) = -p
\]  \( (3.150) \)

\[
-\eta(\rho_o) + \hat{\sigma}_{\rho \rho}^M(\rho_o) = 0.
\]  \( (3.151) \)

Similar to the previous cases, the \( \rho \) equilibrium equation yields the pressure:

\[
p = \int_{\rho_i}^{\rho_o} \frac{1}{\lambda(R)^3} \left[ \hat{\sigma}_{\theta \theta}^M (\rho = \lambda(R)R) - \hat{\sigma}_{\rho \rho}^M (\rho = \lambda(R)R) \right] dR.
\]  \( (3.152) \)

**Case 3**

In Case 3, \( R \in [R_i, R_1] \) is in region S and \( R \in [R_1, R_o] \) is in region M. The boundary conditions are as follows:

\[
-\eta^M (\rho = \rho_o) + \hat{\sigma}_{\rho \rho}^M (\rho = \rho_o) = 0
\]  \( (3.153) \)

\[
-\eta^S (\rho = \rho_i) + \hat{\sigma}_{\rho \rho}^S (\rho = \rho_i) = -p,
\]  \( (3.154) \)

and there is an additional boundary condition for continuity between the two regions:

\[
-\eta^M (\rho = \rho_1) + \hat{\sigma}_{\rho \rho}^M (\rho = \rho_1) = -\eta^S (\rho = \rho_1) + \hat{\sigma}_{\rho \rho}^S (\rho = \rho_1).
\]  \( (3.155) \)

In region M, the \( \rho \) equilibrium equation yields

\[
\eta^S (\rho_1) - \hat{\sigma}_{\rho \rho}^S (\rho_1) = \text{int}_1
\]  \( (3.156) \)

\[
= \int_{R_i}^{R_o} \frac{1}{\lambda(R)^3} \left[ \hat{\sigma}_{\theta \theta}^M (\rho = \lambda(R)R) - \hat{\sigma}_{\rho \rho}^M (\rho = \lambda(R)R) \right] dR.
\]  \( (3.157) \)
In region $S$, we have

\[
\int_{R_i}^{R_1} d\left(-\eta^S + \hat{\sigma}^S_{\rho\rho}\right) = \int_{R_i}^{R_1} \frac{1}{\rho} \left(\hat{\sigma}^S_{\rho\theta} - \hat{\sigma}^S_{\rho\rho}\right) d\rho \tag{3.158}
\]

\[-\eta^S(\rho_1) + \hat{\sigma}^S_{\rho\rho}(\rho_1) - \hat{\sigma}^S_{\rho\rho}(\rho_i) + \eta^S(\rho_i) = \int_{R_i}^{R_1} \frac{1}{\rho} \left(\hat{\sigma}^S_{\rho\theta} - \hat{\sigma}^S_{\rho\rho}\right) d\rho \tag{3.159}
\]

\[-\int_1 + p = \int_2, \tag{3.160}\]

where

\[
\int_2 = \hat{\sigma}^S_{\rho\rho}(\rho_1) = \int_{R_i}^{R_1} \frac{1}{\lambda(R)^3} \left[\hat{\sigma}^S_{\rho\theta}(\rho = \lambda(R)R) - \hat{\sigma}^S_{\rho\rho}(\rho = \lambda(R)R)\right] dR.
\]

This yields the result

\[p = \int_1 + \int_2, \tag{3.162}\]

or

\[
p = \int_{R_i}^{R_o} \frac{1}{\lambda(R)^3} \left[\hat{\sigma}^M_{\rho\theta}(\rho = \lambda(R)R) - \hat{\sigma}^M_{\rho\rho}(\rho = \lambda(R)R)\right] dR \tag{3.163}
\]

\[+ \int_{R_i}^{R_1} \frac{1}{\lambda(R)^3} \left[\hat{\sigma}^S_{\rho\theta}(\rho = \lambda(R)R) - \hat{\sigma}^S_{\rho\rho}(\rho = \lambda(R)R)\right] dR.
\]

**Case 4**

In Case 4, $R \in [R_i, R_1]$ is in region $S$, $R \in [R_1, R_2]$ is in region $M$, and $R \in [R_2, R_o]$ is in region $L$. The boundary conditions are

\[-\eta^L(\rho = \rho_o) + \hat{\sigma}^L_{\rho\rho}(\rho = \rho_o) = 0 \tag{3.164}\]

\[-\eta^S(\rho = \rho_i) + \hat{\sigma}^S_{\rho\rho}(\rho = \rho_i) = -p, \tag{3.165}\]

and there are additional boundary conditions for continuity between the regions:

\[-\eta^M(\rho = \rho_1) + \hat{\sigma}^M_{\rho\rho}(\rho = \rho_1) = -\eta^S(\rho = \rho_1) + \hat{\sigma}^S_{\rho\rho}(\rho = \rho_1) \tag{3.166}\]

\[-\eta^M(\rho = \rho_2) + \hat{\sigma}^M_{\rho\rho}(\rho = \rho_2) = -\eta^L(\rho = \rho_2) + \hat{\sigma}^L_{\rho\rho}(\rho = \rho_2). \tag{3.167}\]

Region $L$ yields the result

\[\eta^L(\rho_2) - \hat{\sigma}^L_{\rho\rho}(\rho_2) = 0. \tag{3.168}\]
Region $M$ yields

\[
-\eta^M(\rho_2) + \hat{\sigma}_{\rho \rho}^M(\rho_2) - \hat{\sigma}_{\rho \rho}^M(\rho_1) + \eta^M(\rho_1) = \int_{\rho_1}^{\rho_2} \frac{1}{\rho} \left( \hat{\sigma}_{\theta \theta}^M - \hat{\sigma}_{\rho \rho}^M \right) d\rho
\]  

(3.169)

\[
-\eta^L(\rho_2) + \hat{\sigma}_{\rho \rho}^L(\rho_2) - \hat{\sigma}_{\rho \rho}^L(\rho_1) + \eta^L(\rho_1) = \int_{\rho_1}^{\rho_2} \frac{1}{\rho} \left( \hat{\sigma}_{\theta \theta}^L - \hat{\sigma}_{\rho \rho}^L \right) d\rho
\]  

(3.170)

\[
\eta^S(\rho_1) - \hat{\sigma}_{\rho \rho}^S(\rho_1) = \int_{\rho_1}^{\rho_2} \frac{1}{\rho} \left( \hat{\sigma}_{\theta \theta}^M - \hat{\sigma}_{\rho \rho}^M \right) d\rho,
\]  

(3.171)

so

\[
\eta^S(\rho_1) - \hat{\sigma}_{\rho \rho}^S(\rho_1) = \text{int}_1
\]  

(3.172)

\[
= \int_{R_1}^{R_2} \frac{1}{\lambda(R)^3} \left[ \hat{\sigma}_{\theta \theta}^M(\rho = \lambda(R) R) - \hat{\sigma}_{\rho \rho}^M(\rho = \lambda(R) R) \right] dR.
\]  

(3.173)

Region $S$ yields

\[
-\eta^S(\rho_1) + \hat{\sigma}_{\rho \rho}^S(\rho_1) - \hat{\sigma}_{\rho \rho}^S(\rho_1) + \eta^S(\rho_1) = \int_{\rho_1}^{\rho_2} \frac{1}{\rho} \left( \hat{\sigma}_{\theta \theta}^M - \hat{\sigma}_{\rho \rho}^M \right) d\rho
\]  

(3.174)

\[
\text{int}_1 - p = \int_{\rho_1}^{\rho_2} \frac{1}{\rho} \left( \hat{\sigma}_{\theta \theta}^M - \hat{\sigma}_{\rho \rho}^M \right) d\rho
\]  

(3.175)

\[
\text{int}_1 + p = \text{int}_2,
\]  

(3.176)

where

\[
\text{int}_2 = \int_{R_1}^{R_2} \frac{1}{\lambda(R)^3} \left[ \hat{\sigma}_{\theta \theta}^S(\rho = \lambda(R) R) - \hat{\sigma}_{\rho \rho}^S(\rho = \lambda(R) R) \right] dR.
\]  

(3.177)

Thus,

\[
p = \text{int}_1 + \text{int}_2,
\]  

(3.178)

or

\[
p = \int_{R_1}^{R_2} \frac{1}{\lambda(R)^3} \left[ \hat{\sigma}_{\theta \theta}^M(\rho = \lambda(R) R) - \hat{\sigma}_{\rho \rho}^M(\rho = \lambda(R) R) \right] dR
\]

\[
+ \int_{R_1}^{R_2} \frac{1}{\lambda(R)^3} \left[ \hat{\sigma}_{\theta \theta}^S(\rho = \lambda(R) R) - \hat{\sigma}_{\rho \rho}^S(\rho = \lambda(R) R) \right] dR.
\]  

(3.179)

Case 5

In Case 5, $R \in [R_i, R_o]$ is in region $S$. The boundary conditions are as follows:

\[
-\eta^S(\rho = \rho_o) + \hat{\sigma}_{\rho \rho}^S(\rho = \rho_o) = 0
\]  

(3.180)

\[
-\eta^S(\rho = \rho_i) + \hat{\sigma}_{\rho \rho}^S(\rho = \rho_i) = -p.
\]  

(3.181)

The $\rho$ equilibrium equation yields the result

\[
p = \int_{R_i}^{R_o} \frac{1}{\lambda(R)^3} \left[ \hat{\sigma}_{\theta \theta}^S(\rho = \lambda(R) R) - \hat{\sigma}_{\rho \rho}^S(\rho = \lambda(R) R) \right] dR.
\]  

(3.182)
Results

The calculations were performed in MATLAB with inner radius $R_i = 1 \text{ cm}$ and outer radius $R_o = 1.1 \text{ cm}$. The parameters that were used for the generalized Mooney-Rivlin model are the same as for the spherical balloon except with $p_2 = 5$, which yields an effective shear modulus $\mu = \frac{1}{2} \left( \sum_{i=1}^{N} c_i p_1 + \sum_{j=1}^{M} d_j q_j \right) = 6.55 \cdot 10^4 \text{ Pa}$. The results for the inflation a cylindrical balloon with fixed axial stretch can be seen in Figures 3.8–3.9b.

![Figure 3.8](image-url)

Figure 3.8: Comparison of this work’s generalized Mooney-Rivlin model with the trace formula model of [23] for inflation pressure of a cylindrical balloon.

Figure 3.8 compares the results from the generalized Mooney-Rivlin model with that of the BTW theory (which features a neo-Hookean energy structure). Similarly to the spherical balloon, the BTW model is unable to capture the correct effect of elasticity in the material at very high values of stretch, so the two curves deviate starting around a stretch of $\lambda_o \approx 3$. This work’s Mooney-Rivlin model captures the physics of the rubber extension at high stretch correctly.

Figure 3.9a and 3.9b shows the results for the balloon inflation at varying anisotropy parameter $r$. The response of the balloon is stiffest in the isotropic state ($r = 1$), and gets correspondingly softer as $r$ increases, as expected. We also note that for $r = 1$, the deformation of the balloon passes through $(\lambda_o, p/\mu) = (1, 0)$ in its undeformed state and then the pressure begins immediately rising upon $\lambda_o$ becoming greater than 1. However, for $r > 1$, the fact that the balloon moves through Case 0 for finite values of $\lambda_o$ means that the balloon starts inflating at points that are not at $(\lambda_o, p/\mu) = (1, 0)$, as mentioned in Chapter 2. This is due to the fact that there is a spontaneous deformation associated with the nematic state, because the liquid crystal molecules spontaneously orient along a preferred direction and
the surrounding polymer network deforms accordingly, with a stretch along that preferred direction and contraction in the transverse axes. Macroscopically, this means that the cylindrical balloon experiences finite stretch with zero stress.

Figure 3.9b shows the solution’s progression through Cases 0 through 5 throughout the deformation. For \( r = 1 \) (rubber), the balloon is in Case 0 (corresponding to region \( L \)) at \( \lambda_o = 1 \) because there is no deformation, but after that point the balloon is entirely in Case 5 (region \( S \)), corresponding to a solid-like response without microstructure formation. For \( r > 1 \), the balloons begin in Case 0 (entirely in region \( L \)). Then the balloons will develop an inner portion that lies in region \( M \) (Case 1), corresponding to fine-scale microstructure formation in the inner part of the balloon, due to the boundary condition that there is internal pressure at the inner radius of the balloon. Then the balloon will then become entirely in region \( M \) (Case 2) before the inner portion of the balloon will develop a solid-like response because it will be in region \( S \) (Case 3). Finally, the balloon will become entirely in region \( S \), and the balloon will have a purely rubber response (Case 5). Note that we do not encounter the case in which \( L, M, \) and \( S \) are all present within the balloon (Case 4) in these calculations.

3.7 Cavitation

Deformation

The original experiments of cavitation by Gent and Lindley demonstrated that when rubber cylinders bonded to parallel plates are subjected to a tensile load, spherical
ruptures form and grow in radius [30]. A schematic for the deformation of cavitation is shown in Figure 3.10.

Figure 3.10: Schematic showing the cross-section of a disk of nematic elastomer bonded to parallel plates, which are stretched in uniaxial tension.

This experiment can be modeled as a spherical void inside of an infinite medium subjected to a state of triaxial extension, as seen in Figure 3.11. We do not consider nucleation (we assume that the spherical void already exists), and we assume that the cavity remains spherical throughout the deformation, which is consistent with the experiments. We are interested in the behavior as the spherical void becomes infinitesimally small (the limit as $R_i \to 0$) or, equivalently, when the sphere of nematic elastomer is infinitely large (the limit as $R_o \to \infty$). The deformation mapping for the growth of such a spherical void is as follows:

$$
\begin{align*}
\rho &= \rho(R) \\
\theta &= \Theta \\
\phi &= \Phi
\end{align*}
$$

The deformation gradient for this specific mapping then is

$$
F = \frac{\partial \rho}{\partial R} e(\rho) \otimes E(R) + \frac{\partial \theta}{\partial \Theta} \frac{\rho \sin \phi}{R \sin \Theta} e(\theta) \otimes E(\Theta) + \frac{\partial \phi}{\partial \Phi} \frac{\rho}{R} e(\phi) \otimes E(\Phi) \\
= \frac{d \rho}{d R} e(\rho) \otimes E(R) + \frac{\rho}{R} e(\theta) \otimes E(\Theta) + \frac{\rho}{R} e(\phi) \otimes E(\Phi).
$$

(3.183) (3.184) (3.185)
Thus,

\[ \mathbf{F}(ij) = \begin{pmatrix} \frac{d \rho}{d R} & \frac{\rho}{R} \\ \frac{1}{\lambda} & \lambda \\ \lambda & \lambda \end{pmatrix}. \]  

(3.186)

Since nematic elastomers are incompressible, we constrain \( \det \mathbf{F} = 1 \):

\[
\begin{align*}
\frac{d \rho}{d R} \rho^2 &= 1 \\
\rho^2 d \rho &= R^2 d R \\
\rho^3 &= R^3 + c. 
\end{align*}
\]

(3.187)  
(3.188)  
(3.189)

We solve for the constant \( c \) in terms of \( \rho_i \) (the deformed radius at the undeformed radius \( R = R_i \)) and obtain

\[
c = \rho_i^3 - R_i^3,
\]

(3.190)

so we have a new expression for the deformed radius, \( \rho \):

\[
\rho = \left[ R^3 + \rho_i^3 - R_i^3 \right]^{1/3}.
\]

(3.191)

Let the non-dimensional azimuthal stretch at the inner radius be denoted by

\[
\lambda_i = \frac{\rho_i}{R_i}.
\]

(3.192)

Then in terms of \( \lambda_i \), the deformed radius is

\[
\rho = \left[ R^3 + R_i^3 \left( \lambda_i^3 - 1 \right) \right]^{1/3},
\]

(3.193)

and the ratio of the deformed radius to undeformed radius is

\[
\lambda = \frac{\rho}{R} = \left[ 1 + \frac{R_i^3}{R^3} \left( \lambda_i^3 - 1 \right) \right]^{1/3}.
\]

(3.194)

Thus, the deformation gradient is

\[
\mathbf{F}(ij) = \begin{pmatrix} \frac{1}{\lambda^2} & \lambda \\ \lambda & \lambda \end{pmatrix}.
\]

(3.195)

The principal values of \( \mathbf{F} \) are \( \lambda_1 \geq \lambda_2 \geq \lambda_3 \): \( \lambda_1 = \lambda_2 = \lambda \) (which correspond to \( \hat{e}_\theta \) and \( \hat{e}_\phi \)) and \( \lambda_3 = \frac{1}{\lambda^2} \) (corresponding to \( \hat{e}_r \)).

There are the same possible cases as presented in the spherical balloon, except that for cavitation we have a new definition for \( R^* \):

\[
R^* = R_i \left( \frac{\lambda_i^3 - 1}{r^{1/4} - 1} \right)^{1/3}.
\]

(3.196)
Stress
The stress state is the same as that for the spherical balloon, seen in Section 3.5.

Solving static equilibrium
This section is the same as that of the spherical balloon, but with the following boundary conditions imposed:

\[ \sigma_{pp}(\rho = \rho_o) = p \]  \hspace{1cm} (3.197)
\[ \sigma_{pp}(\rho = \rho_i) = 0. \]  \hspace{1cm} (3.198)

We solve static equilibrium in the absence of body forces.

Case 1
In Case 1, \( R \in [R_i, R_o] \) is in region \( L \). The outer pressure then is

\[ p = 0. \]  \hspace{1cm} (3.199)

Case 2
In Case 2, \( R \in [R_i, R^*] \) is in region \( M \) and \( R \in [R^*, R_o] \) is in region \( L \). The two boundary conditions can be rewritten specific to the region:

\[ -\eta^M(\rho = \rho_i) + \hat{\sigma}^M_{pp}(\rho = \rho_i) = 0 \]  \hspace{1cm} (3.200)
\[ -\eta^L(\rho = \rho_o) + \hat{\sigma}^L_{pp}(\rho = \rho_o) = p, \]  \hspace{1cm} (3.201)

and there is an additional boundary condition for continuity between the two regions:

\[ -\eta^L(\rho = \rho^*) + \hat{\sigma}^L_{pp}(\rho = \rho^*) = -\eta^M(\rho = \rho^*) + \hat{\sigma}^M_{pp}(\rho = \rho^*). \]  \hspace{1cm} (3.202)

The result from region \( L \) is

\[ -\eta^L(\rho = \rho^*) + \hat{\sigma}^L_{pp}(\rho^*) = p. \]  \hspace{1cm} (3.203)

The result from region \( M \) is

\[ -\eta^M(\rho^*) + \hat{\sigma}^M_{pp}(\rho^*) - \hat{\sigma}^M_{pp}(\rho_i) + \eta^M(\rho_i) = \int_{\rho_i}^{\rho^*} \frac{2}{\rho} \left( \hat{\sigma}^M_{\theta\theta} - \hat{\sigma}^M_{pp} \right) d\rho, \]  \hspace{1cm} (3.204)

or

\[ p = \int_{R_i}^{R^*} \frac{2}{\lambda(R)R} \left[ \hat{\sigma}_{\theta\theta}(\rho = \lambda(R)R) - \hat{\sigma}^M_{pp}(\rho = \lambda(R)R) \right] \frac{1}{\lambda(R)^2} dR. \]  \hspace{1cm} (3.205)
Case 3

In Case 3, all of the balloon is in region $M$. Note that the isotropic state ($r = 1$ and $\lambda \geq 1$) falls under this case. The two boundary conditions for this case are:

\begin{align}
-\eta^M(\rho = \rho_i) + \hat{\sigma}^M_{\rho\rho}(\rho = \rho_i) &= 0 \quad (3.206) \\
-\eta^M(\rho = \rho_o) + \hat{\sigma}^M_{\rho\rho}(\rho = \rho_o) &= p. \quad (3.207)
\end{align}

The result of solving static equilibrium is

\begin{equation}
\begin{split}
p &= \int_{R_i}^{R_o} \frac{2}{\lambda(R)R} \left[ \hat{\sigma}^M_{\rho\rho}(\rho = \lambda(R)R) - \hat{\sigma}^M_{\rho\rho}(\rho = \lambda(R)R) \right] \frac{1}{\lambda(R)^2} dR. \quad (3.208)
\end{split}
\end{equation}

Case 4

In Case 4, all of the balloon is in region $S$. Note that the isotropic state ($r = 1$) falls under this case. The two boundary conditions for this case are:

\begin{align}
-\eta^S(\rho = \rho_i) + \hat{\sigma}^S_{\rho\rho}(\rho = \rho_i) &= 0 \quad (3.209) \\
-\eta^S(\rho = \rho_o) + \hat{\sigma}^S_{\rho\rho}(\rho = \rho_o) &= p. \quad (3.210)
\end{align}

The result of solving static equilibrium is

\begin{equation}
\begin{split}
p &= \int_{R_i}^{R_o} \frac{2}{\lambda(R)R} \left[ \hat{\sigma}^S_{\rho\rho}(\rho = \lambda(R)R) - \hat{\sigma}^S_{\rho\rho}(\rho = \lambda(R)R) \right] \frac{1}{\lambda(R)^2} dR. \quad (3.211)
\end{split}
\end{equation}

Results

The calculations were performed in MATLAB and the solutions are plotted below. The outer radius is $R_o = 1$ cm, and we take the inner radius to be much smaller than the outer radius with a value of $R_i = 1 \cdot 10^{-8}$ m. The parameters used for the generalized Mooney-Rivlin model were the same as for the spherical balloon.

The pressure $p$, normalized by the shear modulus $\mu$, is plotted as a function of the azimuthal stretch at the inner radius $\lambda_i$. Figure 3.12 shows the comparison between the generalized Mooney-Rivlin model of this work and the neo-Hookean model of previous work [23] for an anisotropy parameter of $r = 8$. The BTW theory and generalized Mooney-Rivlin theory deviate starting around a stretch of $\lambda_i \approx 15$. The BTW model predicts that the cavitation pressure plateaus, whereas the Extended Mooney-Rivlin model does not.

Figure 3.13a shows the cavitation results for varying anisotropy parameter $r$. As expected, the isotropic case, $r = 1$, is the stiffest response, and the response is softer as $r$ increases and becomes more nematic.
Figure 3.12: Comparison of this work’s generalized Mooney-Rivlin model with the trace formula model of [23] for pressure of a growing spherical cavity inside a bulk disk.

Figure 3.13: Cavitation results: (a) Results for cavitation at varying anisotropy parameter. (b) Progression of the cavitation solution through individual case numbers.

Figure 3.13b shows the solution’s case at various values of $\lambda_i$. The disk is in Case 1 where the entire structure is undeformed and in region $L$. Then, for $r = 1$, the rest of the deformation belongs to Case 4, where the disk is entirely in region $S$, having a purely elastomer response with no liquid crystal effects. For $r > 1$, the disk develops microstructure in the area immediately surrounding the void that forms during cavitation. Because the void is so small compared to the length scale of the disk, the structure never moves into Case 3 (where the entire disk would be in region $M$).
3.8 Bending

Deformation of bending

The deformation analyzed in this section is the bending of a rectangular block into an arc of a circle, following Ericksen. A schematic of the two-dimensional cross-section of the body is shown in Figure 3.14. In the reference configuration, the block

![Schematic diagram of bending deformation](image)

Figure 3.14: Schematic depicting the mid-plane of a rectangular block undergoing bending deformation.

has Cartesian coordinates \((X_1, X_2, X_3)\), where

\[
X_1 \in [-W, W], \quad X_2 \in [-L, L], \quad X_3 \in [-H, H].
\] (3.212)

The planes \(X_1 = \text{constant}\) become sectors of the cylindrical surface \(\rho = \text{constant}\), the planes \(X_2 = \text{constant}\) become planes \(\theta = \text{constant}\), and the planes \(X_3 = \text{constant}\) become planes \(z = \text{constant}\). We provide an ansatz for the deformation mapping:

\[
\begin{align*}
\rho &= f(X_1) \\
\theta &= g(X_2) \\
z &= \lambda X_3
\end{align*}
\] (3.213)

where \(f\) and \(g\) are functions only of \(X_1\) and \(X_2\) respectively. Therefore, the Cartesian coordinates in the current configuration become

\[
\begin{align*}
x_1 &= \rho \cos \theta = f(X_1) \cos (g(X_2)) \\
x_2 &= \rho \sin \theta = f(X_1) \sin (g(X_2)) \\
x_3 &= z = \lambda X_3
\end{align*}
\] (3.214)
The deformation gradient in Cartesian coordinates is then

$$
F = \nabla \mathbf{x}
$$

$$
= \begin{pmatrix}
  f'(X_1) \cos (g(X_2)) & -f(X_1)g'(X_2) \sin (g(X_2)) & 0 \\
  f'(X_1) \sin (g(X_2)) & f(X_1)g'(X_2) \cos (g(X_2)) & 0 \\
  0 & 0 & \lambda
\end{pmatrix}.
$$

Due to incompressibility, we must reinforce that \( \det F = 1 \):

$$
\lambda f''(X_1) f(X_1) g'(X_2) \cos^2 (g(X_2)) + \lambda f(X_1) f'(X_1) g'(X_2) \sin^2 (g(X_2)) = 1
$$

$$
\lambda f(X_1) f'(X_1) g'(X_2) = 1
$$

$$
g'(X_2) = \frac{1}{\lambda f(X_1) f'(X_1)}. 
$$

To solve for the functions \( f \) and \( g \), we employ separation of variables, yielding two equations to be solved:

$$
\frac{1}{\lambda f(X_1) f'(X_1)} = \alpha 
\quad (3.220)
$$

$$
g'(X_2) = \alpha. 
\quad (3.221)
$$

Solving Equation 3.220, we obtain

$$
\frac{1}{\alpha \lambda} = \int \frac{df}{dX_1}
$$

$$
\int \frac{1}{\alpha \lambda} dX_1 = \int f \, df 
\quad (3.223)
$$

$$
\frac{1}{\alpha \lambda} X_1 + \hat{c} = \frac{f^2}{2} 
\quad (3.224)
$$

$$
f^2 = \frac{2}{\alpha \lambda} X_1 + \beta. 
\quad (3.225)
$$

Since \( \rho = f \), we have

$$
\rho^2 = \frac{2}{\alpha \lambda} X_1 + \beta, 
\quad (3.226)
$$

where \( \beta \) is a constant. Solving Equation 3.221 yields

$$
\frac{dg}{dX_2} = \alpha \quad (3.227)
$$

$$
\int dg = \int \alpha \, dX_2 
\quad (3.228)
$$

$$
g = \alpha X_2 + \tilde{c}, 
\quad (3.229)$$
where $\bar{c}$ is a constant. Since $\theta = g$, we have

$$\theta = \alpha X_2 + \bar{c}. \quad (3.230)$$

Assuming that the deformation is symmetric around the $X_1$-axis,

$$\theta(X_2) = -\theta(-X_2), \quad (3.231)$$

so $\bar{c} = 0$. We then have

$$\theta = \alpha X_2. \quad (3.232)$$

Finally, our deformation mapping is

$$\begin{cases}
\rho^2 = \frac{2}{aL} X_1 + \beta, \\
\theta = \alpha X_2, \\
z = \lambda X_3
\end{cases} \quad (3.233)$$

with $\rho \in \left[\sqrt{-\frac{2W}{aL} + \beta}, \sqrt{\frac{2W}{aL} + \beta}\right]$, $\theta \in [-\alpha L, \alpha L]$, and $z \in [-\lambda H, \lambda H]$. The deformation gradient will be written in mixed cylindrical/Cartesian coordinates. Thus, we have the reference configuration general coordinates as follows:

$$X^i = \{X_1, X_2, X_3\}, \quad (3.234)$$

and the current configuration general coordinates are

$$\xi^i = \{\rho, \theta, z\}, \quad (3.235)$$

with definitions from Equation 3.233. The position vector of a point in the current configuration is

$$x = \rho \cos \theta E_1 + \rho \sin \theta E_2 + z E_3. \quad (3.236)$$

The covariant basis vectors are defined as $e_i = \frac{\partial x}{\partial \xi^i}$, so we have

$$e_\rho = \frac{\partial x}{\partial \rho} = \cos \theta E_1 + \sin \theta E_2 \quad (3.237)$$

$$e_\theta = \frac{\partial x}{\partial \theta} = -\rho \sin \theta E_1 + \rho \cos \theta E_2 \quad (3.238)$$

$$e_z = \frac{\partial x}{\partial z} = E_3 \quad (3.239)$$

with magnitudes

$$|e_\rho| = 1 \quad (3.240)$$

$$|e_\theta| = \rho \quad (3.241)$$

$$|e_z| = 1. \quad (3.242)$$
We introduce the physical basis
\[ e(i) = \frac{e_i}{|e_i|} = \frac{e^i}{|e^i|} \]  
so that we can obtain our deformation gradient:
\[ F = \frac{\partial e^i}{\partial x^j} |e_i||E_j|^{-1} e(i) \otimes E(j) \]  
\[ = \frac{1}{\alpha \lambda \rho} e(\rho) \otimes E(1) + \alpha \rho e(\theta) \otimes E(2) + \lambda e(z) \otimes E(3). \]  

Note that \( \frac{\partial \rho}{\partial x_i} = \frac{1}{\alpha \lambda \rho} \). The left Cauchy-Green tensor is
\[ b = \frac{\partial e^i}{\partial x^j} \frac{\partial e^k}{\partial x^l} |e_i||e_k||E_j|^{-2} e(i) \otimes e(k) \]  
\[ = \frac{1}{\alpha^2 \lambda^2 \rho^2} e(\rho) \otimes e(\rho) + \alpha^2 \rho^2 e(\theta) \otimes e(\theta) + \lambda^2 e(z) \otimes e(z), \]
and the principal stretches are
\[ \frac{1}{\alpha \lambda \rho}, \alpha \rho, \lambda. \]

If the strain energy density of a material can be written in the form \( W = W(\lambda_i) \), where \( \lambda_i \) are the principal stretches, then the Cauchy stress of an incompressible, isotropic hyperelastic body is
\[ \sigma = -p I + \sum_{i=1}^{3} \lambda_i \frac{\partial W}{\partial \lambda_i} e(i) \otimes e(i). \]  

Static equilibrium (in the absence of body forces) is obtained when \( \text{div} \ \sigma = 0 \). In cylindrical coordinates and for a symmetric tensor \( \sigma \), the equations we need to solve are:
\[
\begin{align*}
(\text{div} \ \sigma)(\rho) &= \frac{\partial \sigma(\rho \rho)}{\partial \rho} + \frac{1}{\rho} \frac{\partial \sigma(\rho \theta)}{\partial \theta} + \frac{\sigma(\rho \rho) - \sigma(\theta \theta)}{\rho} + \frac{\partial \sigma(\rho z)}{\partial z}, \\
(\text{div} \ \sigma)(\theta) &= \frac{\partial \sigma(\rho \theta)}{\partial \rho} + \frac{1}{\rho} \frac{\partial \sigma(\rho \theta)}{\partial \theta} + \frac{2 \sigma(\rho \theta)}{\rho} + \frac{\partial \sigma(\theta z)}{\partial z}, \\
(\text{div} \ \sigma)(z) &= \frac{\partial \sigma(\rho z)}{\partial \rho} + \frac{1}{\rho} \frac{\partial \sigma(\rho z)}{\partial \theta} + \frac{\sigma(\rho z)}{\rho} + \frac{\partial \sigma(\theta z)}{\partial z}.
\end{align*}
\]  

Let \( \rho = \kappa \), where \( \kappa = \frac{1}{\alpha} \) is the radius of the neutral axis in the beam. This neutral axis is the axis that experiences zero extension because the arc length of the beam at
this radius does not change from the reference length throughout deformation. The
length of the beam in the undeformed configuration is \(2L\), as is the arc length of
the neutral axis in the current configuration \((s = \rho \theta)\), i.e. the product of the radius
\(\kappa = \frac{1}{\alpha}\) and the angle \((2\alpha L)\). See Figure 3.15 for details.

![Figure 3.15: Schematic depicting the neutral axis of a rectangular block with arc
length \(s\) equal to the length of the undeformed beam, \(2L\).](image)

For the case of plane strain (the stretch in the \(z\) direction is fixed at \(\lambda = 1\)), the
principal stretches are \(\frac{\kappa}{\rho}\) (corresponding to \(\varepsilon_\rho\)), \(1\) (corresponding to \(\varepsilon_z\)), and \(\frac{\rho}{\kappa}\)
(corresponding to \(\varepsilon_\rho\)). The relative magnitudes of the principal stretches depends
on the relationship between \(\rho\) and \(\kappa\). This leads to a breakdown of the regions \(L\),
\(M\), and \(S\) as in Figure 3.16.

![Figure 3.16: Breakdown of the regions \(L\), \(M\), and \(S\) along the radius \(\rho\) in the bending
deformation.](image)

The radius of the neutral axis, \(\rho = \kappa\) lies in the region \(L\). Let us denote the part
of the beam where \(\rho < \kappa\) by \(\Omegas\), and for \(\rho > \kappa\) by \(\Omicron\). The ends of the beam are located
at \(\rho_1 = \sqrt{-\frac{2W}{a\lambda}} + \beta\) and \(\rho_2 = \sqrt{\frac{2W}{a\lambda}} + \beta\). These radii are an unknown and will be
solved for using the static equilibrium equation. We will define the following radii
for convenience:

\[
\rho_{MS}^1 = \frac{\kappa}{r^{1/2}} \tag{3.249}
\]
\[
\rho_{LM}^1 = \frac{\kappa}{r^{1/6}} \tag{3.250}
\]
\[
\rho_{LM}^2 = \kappa r^{1/6} \tag{3.251}
\]
\[
\rho_{MS}^2 = \kappa r^{1/2}. \tag{3.252}
\]

The boundary conditions in this problem are:

\[
\sigma_{pp}|_{\rho=\rho_1} = \sigma_{pp}|_{\rho=\rho_2} = 0. \tag{3.253}
\]

In addition, continuity in the radial stresses at the boundaries between regions must be enforced, i.e. at \( \rho = \rho_{MS}^1, \rho = \rho_{LM}^1, \rho = \rho_{LM}^2, \rho = \rho_{MS}^2 \). The possible cases that arise are:

- **Case 1:** \( r > 1 \) and \( \rho_{LM}^1 \leq \rho_1 \leq \kappa \leq \rho_2 \leq \rho_{LM}^2 \): All of the beam lies in region \( L \).
- **Case 2:** \( r > 1 \) and \( \rho_{MS}^1 \leq \rho_1 \leq \rho_{LM}^1 \leq \kappa \leq \rho_{LM}^2 \leq \rho_2 \leq \rho_{MS}^2 \): The inner portion of the beam lies in region \( L \), and the outer regions lie in \( M \).
- **Case 3a:** \( r > 1 \) and \( \rho_{MS}^1 \leq \rho_1 \leq \rho_{LM}^1 \leq \kappa \leq \rho_{LM}^2 \leq \rho_2 \leq \rho_{MS}^2 \): The beam, in order of increasing radius, lies in region \( S \), then \( M \), then \( L \), then \( M \).
- **Case 3b:** \( r > 1 \) and \( \rho_{MS}^1 \leq \rho_1 \leq \rho_{LM}^1 \leq \kappa \leq \rho_{LM}^2 \leq \rho_{MS}^2 \leq \rho_2 \): The beam, in order of increasing radius, lies in region \( M \), then \( L \), then \( M \), then \( S \).
- **Case 4:** \( r > 1 \) and \( \rho_1 \leq \rho_{LM}^1 \leq \kappa \leq \rho_{LM}^2 \leq \rho_{MS}^2 \leq \rho_2 \): The beam, in order of increasing radius, lies in region \( S \), then \( M \), then \( L \), then \( M \), then \( S \).
- **Case 5:** \( r = 1 \) and \( \rho_1 \leq \kappa \leq \rho_2 \): The entire beam is in region \( S \).

A diagram illustrating the various cases can be seen in Figure 3.17. The heavy black line shows the beam with ends \( \rho = \rho_1 \) and \( \rho = \rho_2 \).

As in the previous deformations, we can gain insight from Figure 3.1 regarding the regions that this deformation will experience. The plane-strain (\( \lambda = 1 \)) bending of a block is the same as a planar extension deformation (the deformation gradient of Equation 3.244 is \( F = \text{diag}(1/\alpha \rho_1, \alpha \rho_2, 1) \)). Therefore, like the cylindrical balloon with \( \xi = 1 \), this means that \( t = s \) for this deformation (recalling that \( s \) is the largest singular value of \( F \) and \( t \) is the largest singular value of \( \text{cof} \ F \)). If we were to follow along the \( t = s \) line in Figure 3.1, we see that the deformation will move progressively through region \( L \) then \( M \) then \( S \).
Figure 3.17: Diagram of all possible cases in the bending of a nematic elastomer block.

Stress, moment, and forces

Static equilibrium (in the absence of body forces) is obtained when \( \text{div } \sigma = 0 \). In cylindrical coordinates and for a symmetric tensor \( \sigma \),

\[
\frac{\partial \sigma_{\rho \rho}}{\partial \rho} + \frac{1}{\rho} \frac{\partial \sigma_{\rho \theta}}{\partial \theta} + \frac{\sigma_{\rho \rho} - \sigma_{\theta \theta}}{\rho} + \frac{\partial \sigma_{\rho z}}{\partial z} = 0,
\]

\[
\frac{\partial \sigma_{\rho \theta}}{\partial \rho} + \frac{1}{\rho} \frac{\partial \sigma_{\theta \theta}}{\partial \theta} + \frac{2\sigma_{\rho \theta}}{\rho} + \frac{\partial \sigma_{\theta z}}{\partial z} = 0, \tag{3.254}
\]

\[
\frac{\partial \sigma_{\rho z}}{\partial \rho} + \frac{1}{\rho} \frac{\partial \sigma_{\theta z}}{\partial \theta} + \frac{\sigma_{\rho z}}{\rho} + \frac{\partial \sigma_{zz}}{\partial z} = 0.
\]

The boundary conditions for this problem are as follows:

\[
\sigma_{\rho \rho}|_{\rho = \rho_1} = 0 \tag{3.255}
\]

\[
\sigma_{\rho \rho}|_{\rho = \rho_2} = 0. \tag{3.256}
\]

For every case, the \( \theta \) and \( z \) equilibrium equations yield that \( \eta = \eta(\rho) \). This leaves only the \( \rho \) equilibrium equation to be solved. After obtaining the stresses, we can calculate the following three quantities. First, the magnitude of the force in the \( \theta \)-direction is

\[
|F_\theta| = \int_{-\lambda H}^{\lambda H} \int_{\rho_1}^{\rho_2} \sigma_{\theta \theta} d\rho dz. \tag{3.257}
\]

(Nota: \( \lambda = 1 \) here.) Second, the magnitude of the force in the \( z \)-direction is

\[
|F_z| = \int_{-\frac{L}{k}}^{\frac{L}{k}} \int_{\rho_1}^{\rho_2} \sigma_{zz} d\rho d\theta. \tag{3.258}
\]
Finally, the moment is
\[ M = \int_{-\Lambda H}^{\Lambda H} \int_{\rho_1}^{\rho_2} \sigma_{\theta \theta} \rho d\rho dz. \]  
(3.259)

**Solving static equilibrium**

**Case 1**

In this trivial case, the beam lies entirely in the \( L \) region. That is, the radius \( \rho \in [\rho_1, \rho_2] \) is such that
\[ \rho_{LM}^1 \leq \rho_1 \leq \kappa \leq \rho_2 \leq \rho_{LM}^2. \]  
(3.260)

Thus, the stress condition is simply the Lagrange multiplier:
\[ \sigma^L = -\eta^L I. \]  
(3.261)

We can solve for static equilibrium by solving \( \text{div} \ \sigma = 0 \) in cylindrical coordinates, given by Equation 3.248:
\[ z : \eta^L = \eta^L(\rho, \theta) \]  
(3.262)
\[ \theta : \eta^L = \eta^L(\rho) \]  
(3.263)
\[ \rho : \frac{\partial \sigma^L_{\rho \rho}}{\partial \rho} + \frac{1}{\rho} \left( \sigma^L_{\rho \rho} - \sigma^L_{\theta \theta} \right) = 0 \rightarrow \sigma^L_{\rho \rho} = \text{const} \rightarrow \eta^L = \text{const}. \]  
(3.264)

We can use the traction boundary condition, \( \sigma^L_{\rho \rho}|_{\rho = \rho_1} = \sigma^L_{\rho \rho}|_{\rho = \rho_2} = 0 \). Thus, we find that the Lagrange multiplier \( \eta^L = 0 \). The magnitude of the force in the \( \theta \)-direction is calculated from Equation 3.257:
\[ |F_\theta| = \int_{-\Lambda H}^{\Lambda H} \int_{\rho_1}^{\rho_2} \sigma^L_{\theta \theta} d\rho dz = 0. \]  
(3.265)

The magnitude of the force in the \( z \)-direction is calculated from Equation 3.258:
\[ |F_z| = \int_{-\frac{L}{\kappa}}^{\frac{L}{\kappa}} \int_{\rho_1}^{\rho_2} \sigma^L_{zz} d\rho d\theta = 0. \]  
(3.266)

Finally, the moment is calculated from Equation 3.259:
\[ M = \int_{-\Lambda H}^{\Lambda H} \int_{\rho_1}^{\rho_2} \sigma^L_{\theta \theta} \rho d\rho dz = 0. \]  
(3.267)

This case physically refers to a beam with \( \kappa \to \infty \) (the radius of curvature is infinitely large, so the beam is flat). This refers to the undeformed case, which corresponds to zero stresses.
Case 2

In Case 2, the beam has outer regions $M$ and inner regions $L$, and the radius $\rho \in [\rho_1, \rho_2]$ is such that

$$\rho_{MS}^1 \leq \rho_1 \leq \rho_{LM}^1 \leq \kappa \leq \rho_{LM}^2 \leq \rho_2 \leq \rho_{MS}^2.$$  \hfill (3.268)

The boundary conditions (BCs) specific to this case are:

$$\sigma_{pp}^{M^{(\cdot)}}(\rho_1) = 0$$  \hfill (3.269)

$$\sigma_{pp}^{M^{(\cdot)}}(\rho_{LM}^1) = \sigma_{pp}^{L^{(\cdot)}}(\rho_{LM}^1)$$  \hfill (3.270)

$$\sigma_{pp}^{L^{(1)}}(\kappa) = \sigma_{pp}^{L^{(2)}}(\kappa)$$  \hfill (3.271)

$$\sigma_{pp}^{L^{(\cdot)}}(\rho_{LM}^2) = \sigma_{pp}^{M^{(\cdot)}}(\rho_{LM}^2)$$  \hfill (3.272)

$$\sigma_{pp}^{M^{(\cdot)}}(\rho_2) = 0.$$  \hfill (3.273)

We solve the equations of static equilibrium using Equation 3.248, which are in cylindrical coordinates. For every region, we only need to solve the equation in $\rho$:

$$\frac{\partial \sigma_{pp}}{\partial \rho} + \frac{1}{\rho} \left( \sigma_{pp} - \sigma_{\theta\theta}^{L^{(\cdot)}} \right) = 0.$$  \hfill (3.274)

Moving from left to right within the beam, we start at region $M^{(\cdot)}$:

$$- \frac{d \eta^{M^{(\cdot)}}}{d \rho} + \frac{d \hat{\sigma}_{pp}^{M^{(\cdot)}}}{d \rho} + \frac{1}{\rho} \left( \hat{\sigma}_{pp}^{M^{(\cdot)}} - \hat{\sigma}_{\theta\theta}^{M^{(\cdot)}} \right) = 0.$$  \hfill (3.275)

Solving for $\eta^{M^{(\cdot)}}$ yields

$$\eta^{M^{(\cdot)}} = \int \left[ \frac{d \hat{\sigma}_{pp}^{M^{(\cdot)}}}{d \rho} + \frac{1}{\rho} \left( \hat{\sigma}_{pp}^{M^{(\cdot)}} - \hat{\sigma}_{\theta\theta}^{M^{(\cdot)}} \right) \right] d \rho + C_1.$$  \hfill (3.276)

Using the first BC, we can obtain $C_1$. Next, in region $L^{(\cdot)}$, we repeat the same procedure and obtain:

$$\eta^{L^{(\cdot)}} = \int \left[ \frac{d \hat{\sigma}_{pp}^{L^{(\cdot)}}}{d \rho} + \frac{1}{\rho} \left( \hat{\sigma}_{pp}^{L^{(\cdot)}} - \hat{\sigma}_{\theta\theta}^{L^{(\cdot)}} \right) \right] d \rho + C_2.$$  \hfill (3.277)

Using the second BC, we can obtain $C_2$. Repeating the same procedure in $L^{(\cdot)}$, we have

$$\eta^{L^{(\cdot)}} = \int \left[ \frac{d \hat{\sigma}_{pp}^{L^{(\cdot)}}}{d \rho} + \frac{1}{\rho} \left( \hat{\sigma}_{pp}^{L^{(\cdot)}} - \hat{\sigma}_{\theta\theta}^{L^{(\cdot)}} \right) \right] d \rho + C_3.$$  \hfill (3.278)

Using the third BC gives us $C_3$. Finally, in region $M^{(\cdot)}$, we have

$$\eta^{M^{(\cdot)}} = \int \left[ \frac{d \hat{\sigma}_{pp}^{M^{(\cdot)}}}{d \rho} + \frac{1}{\rho} \left( \hat{\sigma}_{pp}^{M^{(\cdot)}} - \hat{\sigma}_{\theta\theta}^{M^{(\cdot)}} \right) \right] d \rho + C_4.$$  \hfill (3.279)
Using the fourth BC gives us $C_4$, and using the last BC gives us the unknown $\beta$, which gives us the deformed radii of the beam, $\rho_1$ and $\rho_2$. Thus, we now have all the stress values and can calculate $|F_\theta|$, $|F_z|$, and $M$ from Eqns. 3.257, 3.258, and 3.259 respectively. The integral in $\rho$ between $\rho_1$ and $\rho_2$ must be broken into individual regions with corresponding bounds:

$$|F_\theta| = \int_{-\lambda H}^{\lambda H} \left[ \int_{\rho_1}^{\rho_1} \sigma_{\theta\theta}^{\text{M}(1)} d\rho + \int_{\rho_1}^{\rho_2} \sigma_{\theta\theta}^{\text{L}(1)} d\rho \right] dz + \int_{-\lambda H}^{\lambda H} \left[ \int_{\rho_1}^{\rho_1} \sigma_{\theta\theta}^{\text{L}(2)} d\rho + \int_{\rho_2}^{\rho_2} \sigma_{\theta\theta}^{\text{M}(2)} d\rho \right] dz$$  \hspace{1cm} (3.280)

$$|F_z| = \int_{-\frac{\lambda}{2}}^{\frac{\lambda}{2}} \left[ \int_{\rho_1}^{\rho_1} \sigma_{zz}^{\text{M}(1)} d\rho + \int_{\rho_1}^{\rho_2} \sigma_{zz}^{\text{L}(1)} d\rho \right] + \int_{-\frac{\lambda}{2}}^{\frac{\lambda}{2}} \left[ \int_{\rho_1}^{\rho_1} \sigma_{zz}^{\text{L}(2)} d\rho + \int_{\rho_2}^{\rho_2} \sigma_{zz}^{\text{M}(2)} d\rho \right] d\theta$$  \hspace{1cm} (3.281)

$$M = \int_{-\lambda H}^{\lambda H} \left[ \int_{\rho_1}^{\rho_1} \sigma_{\theta\theta}^{\text{M}(1)} \rho d\rho + \int_{\rho_1}^{\rho_2} \sigma_{\theta\theta}^{\text{L}(1)} \rho d\rho \right] + \int_{-\lambda H}^{\lambda H} \left[ \int_{\rho_1}^{\rho_1} \sigma_{\theta\theta}^{\text{L}(2)} \rho d\rho + \int_{\rho_2}^{\rho_2} \sigma_{\theta\theta}^{\text{M}(2)} \rho d\rho \right] dz.$$  \hspace{1cm} (3.282)

**Case 3a**

In Case 3a, the beam, in order of increasing radius, has portions that are in region $M$, $L$, $M$, and $S$. The radius $\rho \in [\rho_1, \rho_2]$ is such that

$$\rho_1 \leq \rho_{\text{MS}}^1 \leq \rho_L^1 \leq \kappa \leq \rho_L^2 \leq \rho_2 \leq \rho_{\text{MS}}^2.$$  \hspace{1cm} (3.283)

and the boundary conditions specific to this case are:

$$\sigma_{\theta\theta}^{\text{S}(1)} (\rho_1) = 0,$$  \hspace{1cm} (3.284)

$$\sigma_{\theta\theta}^{\text{S}(1)} (\rho_{\text{MS}}^1) = \sigma_{\theta\theta}^{\text{M}(1)} (\rho_{\text{MS}}^1),$$  \hspace{1cm} (3.285)

$$\sigma_{\theta\theta}^{\text{S}(1)} (\rho_L^1) = \sigma_{\theta\theta}^{\text{L}(1)} (\rho_L^1),$$  \hspace{1cm} (3.286)

$$\sigma_{\theta\theta}^{\text{L}(1)} (\kappa) = \sigma_{\theta\theta}^{\text{L}(2)} (\kappa),$$  \hspace{1cm} (3.287)

$$\sigma_{\theta\theta}^{\text{S}(1)} (\rho_{\text{MS}}^2) = \sigma_{\theta\theta}^{\text{M}(2)} (\rho_{\text{MS}}^2);$$  \hspace{1cm} (3.288)

$$\sigma_{\theta\theta}^{\text{S}(1)} (\rho_L^2) = \sigma_{\theta\theta}^{\text{M}(2)} (\rho_L^2).$$  \hspace{1cm} (3.289)

Solving from left to right within the beam, we start at region $S(1)$:

$$\eta^{\text{S}(1)} = \int \left[ \sigma_{\theta\theta}^{\text{S}(1)} (\rho_1) + 1 \rho \left( \sigma_{\theta\theta}^{\text{S}(1)} (\rho_1) - \sigma_{\theta\theta}^{\text{S}(1)} (\rho_{\text{MS}}^1) \right) \right] d\rho + C_1.$$  \hspace{1cm} (3.290)
We can use the first BC to solve for $C_1$. In region $M\odot$ we have
\[ \eta^{M\odot} = \int \left[ \frac{d\hat{\sigma}^{M\odot}_{pp}}{d\rho} + \frac{1}{\rho} \left( \hat{\sigma}^{M\odot}_{\rho\rho} - \hat{\sigma}^{M\odot}_{\theta\theta} \right) \right] d\rho + C_2. \tag{3.294} \]

Solving the second BC gives us $C_2$. In region $L\odot$ we have
\[ \eta^{L\odot} = \int \left[ \frac{d\hat{\sigma}^{L\odot}_{pp}}{d\rho} + \frac{1}{\rho} \left( \hat{\sigma}^{L\odot}_{\rho\rho} - \hat{\sigma}^{L\odot}_{\theta\theta} \right) \right] d\rho + C_3. \tag{3.295} \]

Solving the third BC gives us $C_3$. In region $L\otimes$ we have
\[ \eta^{L\otimes} = \int \left[ \frac{d\hat{\sigma}^{L\otimes}_{pp}}{d\rho} + \frac{1}{\rho} \left( \hat{\sigma}^{L\otimes}_{\rho\rho} - \hat{\sigma}^{L\otimes}_{\theta\theta} \right) \right] d\rho + C_4. \tag{3.296} \]

Solving the fourth BC gives us $C_4$. Finally, in region $M\otimes$ we have
\[ \eta^{M\otimes} = \int \left[ \frac{d\hat{\sigma}^{M\otimes}_{pp}}{d\rho} + \frac{1}{\rho} \left( \hat{\sigma}^{M\otimes}_{\rho\rho} - \hat{\sigma}^{M\otimes}_{\theta\theta} \right) \right] d\rho + C_5. \tag{3.297} \]

Solving the fifth BC gives us $C_5$, and solving the last BC gives us $\beta$. We can calculate $|F_\theta|$, $|F_z|$, and $M$ from Eqns. 3.257, 3.258, and 3.259 analogously as we did in Case 2, breaking the integrals down to their individual regions between $\rho_1$ and $\rho_2$.

**Case 3b**

Case 3b is very similar to Case 3a. In this case, the beam has portions that are in region $S$, $M$, $L$, and $M$, in order of increasing radius. The radius $\rho \in [\rho_1, \rho_2]$ is such that
\[ \rho_{MS}^1 \leq \rho_1 \leq \rho_{LM}^1 \leq \kappa \leq \rho_{LM}^2 \leq \rho_{MS}^2 \leq \rho_2, \tag{3.298} \]
and the boundary conditions specific to this case are:
\[ \sigma_{pp}^{M\odot}(\rho_1) = 0 \tag{3.299} \]
\[ \sigma_{pp}^{M\otimes}(\rho_{LM}^1) = \sigma_{pp}^{L\otimes}(\rho_{LM}^1) \tag{3.300} \]
\[ \sigma_{pp}^{L\otimes}(\kappa) = \sigma_{pp}^{L\otimes}(\kappa) \tag{3.301} \]
\[ \sigma_{pp}^{L\otimes}(\rho_{LM}^2) = \sigma_{pp}^{M\otimes}(\rho_{LM}^2) \tag{3.302} \]
\[ \sigma_{pp}^{M\otimes}(\rho_{MS}^2) = \sigma_{pp}^{S\otimes}(\rho_{MS}^2) \tag{3.303} \]
\[ \sigma_{pp}^{S\otimes}(\rho_2) = 0. \tag{3.304} \]
Solving from left to right within the beam, we start at region \( M_1 \):

\[
\eta^{M_1} = \int \left[ \frac{d\hat{\sigma}_{\rho\rho}^{M_1}}{d\rho} + \frac{1}{\rho} \left( \hat{\sigma}_{\rho\phi}^{M_1} - \hat{\sigma}_{\theta\theta}^{M_1} \right) \right] d\rho + C_1. \tag{3.305}
\]

We can use the first BC to solve for \( C_1 \). In region \( L_1 \) we have

\[
\eta^{L_1} = \int \left[ \frac{d\hat{\sigma}_{\rho\rho}^{L_1}}{d\rho} + \frac{1}{\rho} \left( \hat{\sigma}_{\rho\phi}^{L_1} - \hat{\sigma}_{\theta\theta}^{L_1} \right) \right] d\rho + C_2. \tag{3.306}
\]

Solving the second BC gives us \( C_2 \). In region \( L_2 \) we have

\[
\eta^{L_2} = \int \left[ \frac{d\hat{\sigma}_{\rho\rho}^{L_2}}{d\rho} + \frac{1}{\rho} \left( \hat{\sigma}_{\rho\phi}^{L_2} - \hat{\sigma}_{\theta\theta}^{L_2} \right) \right] d\rho + C_3. \tag{3.307}
\]

Solving the third BC gives us \( C_3 \). In region \( M_2 \) we have

\[
\eta^{M_2} = \int \left[ \frac{d\hat{\sigma}_{\rho\rho}^{M_2}}{d\rho} + \frac{1}{\rho} \left( \hat{\sigma}_{\rho\phi}^{M_2} - \hat{\sigma}_{\theta\theta}^{M_2} \right) \right] d\rho + C_4. \tag{3.308}
\]

Solving the fourth BC gives us \( C_4 \). Finally, in region \( S_2 \) we have

\[
\eta^{S_2} = \int \left[ \frac{d\hat{\sigma}_{\rho\rho}^{S_2}}{d\rho} + \frac{1}{\rho} \left( \hat{\sigma}_{\rho\phi}^{S_2} - \hat{\sigma}_{\theta\theta}^{S_2} \right) \right] d\rho + C_5. \tag{3.309}
\]

Solving the fifth BC gives us \( C_5 \), and solving the last BC gives us \( \beta \). We can calculate \(|F_\theta|, |F_z|, \) and \( M \) from Eqns. 3.257, 3.258, and 3.259 as in Case 3a.

**Case 4**

In Case 4, in order of increasing radius, the beam lies in regions \( S, M, L, M, \) and \( S \). The radius \( \rho \in [\rho_1, \rho_2] \) is such that

\[
\rho_1 \leq \rho_{MS}^1 \leq \rho_{LMS}^1 \leq \kappa \leq \rho_{LM}^2 \leq \rho_{MS}^2 \leq \rho_2. \tag{3.310}
\]

The boundary conditions specific to this case are:

\[
\sigma_{\rho\rho}^{S_2}(\rho_1) = 0 \tag{3.311}
\]
\[
\sigma_{\rho\rho}^{M_1}(\rho_{MS}^1) = \sigma_{\rho\rho}^{S_2}(\rho_{MS}^1) \tag{3.312}
\]
\[
\sigma_{\rho\rho}^{M_1}(\rho_{LM}^1) = \sigma_{\rho\rho}^{L_1}(\rho_{LM}^1) \tag{3.313}
\]
\[
\sigma_{\rho\rho}(\kappa) = \sigma_{\rho\rho}(\kappa) \tag{3.314}
\]
\[
\sigma_{\rho\rho}^{L_2}(\rho_{LM}^2) = \sigma_{\rho\rho}^{M_2}(\rho_{LM}^2) \tag{3.315}
\]
\[
\sigma_{\rho\rho}^{M_2}(\rho_{MS}^2) = \sigma_{\rho\rho}^{S_2}(\rho_{MS}^2) \tag{3.316}
\]
\[
\sigma_{\rho\rho}(\rho_2) = 0. \tag{3.317}
\]
Solving from left to right within the beam, we start at region $S_1$:

$$
\eta^{S_1} = \int \left[ \frac{d\hat{\sigma}^{S_1 \rho \rho}}{d\rho} + \frac{1}{\rho} \left( \hat{\sigma}^{S_1 \rho \rho} - \hat{\sigma}^{S_1 \theta \theta} \right) \right] d\rho + C_1. \tag{3.318}
$$

We can use the first BC to solve for $C_1$. In region $M_1$ we have

$$
\eta^{M_1} = \int \left[ \frac{d\hat{\sigma}^{M_1 \rho \rho}}{d\rho} + \frac{1}{\rho} \left( \hat{\sigma}^{M_1 \rho \rho} - \hat{\sigma}^{M_1 \theta \theta} \right) \right] d\rho + C_2. \tag{3.319}
$$

Solving the second BC gives us $C_2$. In region $L_1$ we have

$$
\eta^{L_1} = \int \left[ \frac{d\hat{\sigma}^{L_1 \rho \rho}}{d\rho} + \frac{1}{\rho} \left( \hat{\sigma}^{L_1 \rho \rho} - \hat{\sigma}^{L_1 \theta \theta} \right) \right] d\rho + C_3. \tag{3.320}
$$

Solving the third BC gives us $C_3$. In region $L_2$ we have

$$
\eta^{L_2} = \int \left[ \frac{d\hat{\sigma}^{L_2 \rho \rho}}{d\rho} + \frac{1}{\rho} \left( \hat{\sigma}^{L_2 \rho \rho} - \hat{\sigma}^{L_2 \theta \theta} \right) \right] d\rho + C_4. \tag{3.321}
$$

Solving the fourth BC gives us $C_4$. In region $M_2$ we have

$$
\eta^{M_2} = \int \left[ \frac{d\hat{\sigma}^{M_2 \rho \rho}}{d\rho} + \frac{1}{\rho} \left( \hat{\sigma}^{M_2 \rho \rho} - \hat{\sigma}^{M_2 \theta \theta} \right) \right] d\rho + C_5. \tag{3.322}
$$

Solving the fifth BC gives us $C_5$. Finally, in region $S_2$ we have

$$
\eta^{S_2} = \int \left[ \frac{d\hat{\sigma}^{S_2 \rho \rho}}{d\rho} + \frac{1}{\rho} \left( \hat{\sigma}^{S_2 \rho \rho} - \hat{\sigma}^{S_2 \theta \theta} \right) \right] d\rho + C_6 \tag{3.323}
$$

Solving the sixth BC gives us $C_6$, and solving the last BC gives us $\beta$. We can calculate $|F_\theta|$, $|F_\zeta|$, and $M$ from Eqns. 3.257, 3.258, and 3.259.

**Case 5**

In Case 5, the entire beam is in region $S$ ($\rho_1 \leq \kappa \leq \rho_2$), and the anisotropy parameter $r = 1$, corresponding to a rubber material, and not a nematic elastomer. The radius $\rho \in [\rho_1, \rho_2]$ is such that

$$
\rho_1 \leq \kappa \leq \rho_2. \tag{3.324}
$$

The boundary conditions specific to this case are:

$$
\sigma^{S_1 \rho \rho}_\rho (\rho_1) = 0 \quad \text{(3.325)}
$$

$$
\sigma^{S_1 \rho \rho \rho}_\rho (\kappa) = \sigma^{S_2 \rho \rho \rho}_\rho (\kappa) \quad \text{(3.326)}
$$

$$
\sigma^{S_2 \rho \rho \rho}_\rho (\rho_2) = 0. \quad \text{(3.327)}
$$
Notice that at $\rho = \kappa$, the stress in the r-direction will be zero: $\sigma_{rr}^{S\Theta}(\kappa) = \sigma_{r\theta}^{S\Theta}(\kappa) = 0$, corresponding to the neutral axis at the radius of curvature. We begin by solving in the region $r \leq \kappa$ with $S^{\Theta}$:

$$
\eta^{S\Theta} = \int \left[ \frac{d\hat{\sigma}_{rr}^{S\Theta}}{d\rho} + \frac{1}{\rho} \left( \hat{\sigma}_{r\theta}^{S\Theta} - \hat{\sigma}_{\rho\theta}^{S\Theta} \right) \right] d\rho + C_1. \tag{3.328}
$$

We can use the first BC to get $C_1$, then move to the region $r \geq \kappa$ with $S^{\Theta}$:

$$
\eta^{S\Theta} = \int \left[ \frac{d\hat{\sigma}_{rr}^{S\Theta}}{d\rho} + \frac{1}{\rho} \left( \hat{\sigma}_{r\theta}^{S\Theta} - \hat{\sigma}_{\rho\theta}^{S\Theta} \right) \right] d\rho + C_2. \tag{3.329}
$$

We use the second BC to get $C_2$, and then the third BC to get $\beta$. With all the unknowns known, we again calculate $|F_\theta|$, $|F_z|$, and $M$ from Eqns. 3.257, 3.258, and 3.259.

**Results**

The stresses, forces, and moments as well as the deformed radii were all solved for using Mathematica using the following parameters: $c_1 = 1.03 \cdot 10^5$ Pa, $c_2 = 1.96 \cdot 10^2$ Pa, $d_1 = 1.63 \cdot 10^{-2}$ Pa, $p_1 = 1$, $p_2 = 1$, and $q_1 = 2$. The beam’s width is $W = 1$ cm, and its length is $L = 2W$. Figures 3.19 and 3.20 (with the legend in Figure 3.18) plot the bending moment and the cases as a function of $\alpha = 1/\kappa$ (the radius of curvature). A value of $\kappa \to \infty$ (or $\alpha = \frac{1}{\kappa} = 0$) corresponds to the undeformed beam (with an infinite radius of curvature), and as $\kappa$ decreases ($\alpha$ increases), this corresponds to the beam bending more and more.

\[
\begin{array}{ccccccccccc}
 r & 1 & 2 & 4 & 6 & 8 \\
 Case 1 & \times & \times & \times & \times \\
 Case 2 & \circ & \circ & \circ & \circ \\
 Case 3a & \square & \square & \square & \square \\
 Case 3b & + & + & + & + \\
 Case 4 & \star & \star & \star & \star \\
 Case 5 & \star \\
\end{array}
\]

*Figure 3.18: Legend for Figures 3.19 and 3.20.*

The case $r = 1$ yields the highest bending moment, as expected for the isotropic material. The solution is entirely in Case 5, i.e. the beam is entirely in the $S$ region,
so it has a solid-like rubber response. For $r > 1$, we first note the spontaneous deformation associated with the nematic state, similar to what we observed in the balloon inflation deformations. Due to the change in temperature associated with an anisotropy parameter $r > 1$, the block will experience a spontaneous stretching along some preferred direction and contraction perpendicular in the transverse directions. There is a change in the block’s curvature at zero moment associated with this spontaneous deformation.

Next, we turn our attention to the fact that there are sometimes multiple cases that yield an equilibrium solution for a given value of $\alpha = 1/\kappa$, so all possible solutions are plotted. For instance, for $r = 2$, the beam begins in Case 1 at $\alpha = 0$, where all of the beam lies in region $L$ and there is no deformation. Then for every point $\alpha > 0$, the beam has equilibrium solutions in Case 2, where the inner part of the beam is in region $L$, and the outer parts of the beam lie in region $M$. Physically, the beam is developing fine-scale microstructure in the innermost region (which is experiencing compressive stress) and the outermost region (which is experiencing tensile stress). The region in between is still in region $L$, experiencing zero stress. Then for $\alpha$ approximately between 40 and 100, the beam has three possible equilibrium solutions, in Cases 2, 3a, and 4. Case 3a physically corresponds to some development of region $S$ in the innermost part of the beam, and Case 4 corresponds to development of region $S$ in both the innermost and outermost parts.
of the beam. Then Case 3a stops being a viable equilibrium solution, then shortly after that Case 4 stops being a possible equilibrium solution, and for $\alpha > 160$, Case 2 is the only equilibrium solution, and the bending moment decreases.

![Figure 3.20: Progression of the bending solution through individual case numbers.](image)

3.9 Conclusion

The remarkable softness that is characteristic of nematic elastomers has emphasized the need for an energy density that extends beyond the limitations of the neo-Hookean trace formula. In this work we have developed a generalized Mooney-Rivlin energy density and have used this model to solve some examples of universal deformations in nematic elastomers. Nematic elastomers have been studied extensively in classical geometries, such as the deformation of thin sheets, and traditional loading conditions, such as uniaxial tension, but they have yet to be explored in-depth as deformations applied to the bulk material, such as the bending of a block, cavitation of a disk, and internal pressure of a balloon. With the relatively recent developments in synthesizing these materials in the bulk with click chemistry and 3D printing, more complex geometries and loading configurations can be explored, and the solutions to the universal deformations for nematic elastomers is a significant step in this direction.
Chapter 4

A GENERAL CONSTITUTIVE MODEL FOR A NON-IDEAL ISOTROPIC-GENESIS POLYDOMAIN NEMATIC ELASTOMER

4.1 Introduction

The previous chapter addressed ideal nematic elastomers. In these ideal materials, the director can reorient freely. Consequently, the fine-scale domain patterns can form and change freely, and the overall behavior is described by the relaxed energy. However, in reality, there is a resistance to changing domain patterns. Specifically, as a result of the synthesis process, there may be a local preference to director alignment at each material point. This preference is extremely large in nematic-genesis elastomers (materials crosslinked in the nematic state), but relatively small in the isotropic-genesis elastomers (materials crosslinked in the isotropic state) [8, 74]. Consequently, isotropic-genesis elastomers still show soft behavior, though one needs to apply a small stress to change domain patterns—see Figure 4a from [74]. Further, isotropic-genesis elastomers have a fine polydomain structure as synthesized because the director preference is random—see Figure 2a from [74]. This preference can be modeled as non-ideality [7], a term that is small in isotropic-genesis nematic elastomers (also discussed in Chapter 2 of this thesis). Since the material is now possibly heterogenous, one has to relax and homogenize: this can be accomplished through bounds [9] or numerically [85].

In this chapter, we develop a constitutive model to describe the macroscopic response of nematic elastomers (on a scale large compared to domain patterns, but small compared to the scale of application). The idea is to use the relaxation and identify internal variables that describe the fine-scale domain patterns and then impose a kinetic process with dissipation on these internal variables. We validate the model against the experiments of Tokumoto, Takabe, and Urayama, as reported in Tokumoto et al. [71]. Finally, we implement the model in the finite element program ABAQUS. After verifying it using homogenous deformations, we study the problem of torsion of a cylinder. We identify an interesting instability at large torsional strains as a result of the Poynting effect.

This continuum-level model, and its implementation for finite element analysis, is the first of its kind for modeling non-ideal polydomain nematic elastomers in 3D.
It is a powerful tool that can be used to analyze nematic elastomers in arbitrarily complex deformations, which will contribute towards nematic elastomers becoming an accessible engineering material.

4.2 Formulation of the constitutive relation

Revisiting the relaxed energy of an ideal nematic elastomer

We follow the work of DeSimone and Dolzmann [23] using the same three regions of interest, a liquid-like region $L$, a solid-like region $S$, and a region $M$ in which laminated microstructure occurs. Recall the generalized energy density of a nematic elastomer

$$W_{NE}(F, n) = f(F^T \ell_n^{-1} F),$$

(4.1)

where $\ell_n = r^{-1/3}((r - 1)n \otimes n + I)$ is the “step-length tensor” that describes the metric of the nematic elastomer in the “stress-free state”. The corresponding elastic energy is

$$W(F) = \min_n W_{NE}(F, n) = \min_{Q \in SO(3)} f(F^T \ell_n^{-1} F) = \min_{Q \in SO(3)} f(F^T Q \ell_n^{-1} Q^T F).$$

(4.2)

Now, consider a situation where the nematic elastomer has formed a fine-scale domain pattern so that the resulting “stress-free” state is described by a metric $G$. We know from previous chapters that

$$G = QG_0(\lambda, \delta)Q^T,$$

(4.3)

where

$$Q \in SO(3),$$

(4.4)

$$G_0(\lambda, \delta) = \begin{pmatrix} \lambda^2 & 0 & 0 \\ 0 & \frac{\delta^2}{\lambda^2} & 0 \\ 0 & 0 & \frac{1}{\delta^2} \end{pmatrix},$$

(4.5)

and

$$\lambda, \delta \in T := \{(s, t) : t \leq r^{1/6}, t \leq s^2, t \geq \sqrt{s}\}.$$

(4.6)

The triangular region $T$ is depicted in Figure 4.1.

In analogy to (4.2), we may write the energy of a nematic elastomer with fine-scale domain pattern characterized by $\lambda, \delta$ to be

$$W_{PNE}(F, \lambda, \delta) = \min_{Q \in SO(3)} f(F^T Q G_0^{-1} Q^T F),$$

(4.7)
so that the effective energy over arbitrary domain patterns is given by
\[
W_{\text{eff}}(F) = \min_{\lambda, \delta \in T} \min_{Q \in SO(3)} f(F^T Q G_0^{-1} Q^T F).
\]  
(4.8)

**Theorem 1.** With the definitions above,
\[
W_{\text{eff}}(F) = W^{qc}(F).
\]  
(4.9)

**Proof.** Let \( s \) be the largest singular value of \( F \) and \( t \) the product of the largest singular values of \( F \). Then,
\[
W_{\text{eff}}(F) = \min_{\lambda, \delta \in T} f\left(\frac{(s^2)}{\lambda^2} + \frac{t^2}{\lambda^2} - 3 \right)^{p_i} + \sum_j d_j \left(\frac{\lambda^2}{s^2} + \frac{\delta^2}{t^2} - 3\right)^{q_j}.
\]  
(4.10)

Now, recall that
\[
f(A) = \sum_i c_i (\text{tr} A - 3)^{p_i} + \sum_j d_j (\text{tr} (\text{cof} A) - 3)^{q_j}
\]  
(4.11)

where \( p_i, q_j \geq 1 \) so that
\[
W_{\text{PNE}}(F, \lambda, \delta) = \sum_i c_i \left(\frac{s^2}{\lambda^2} + \frac{t^2}{\lambda^2} - 3\right)^{p_i} + \sum_j d_j \left(\frac{\lambda^2}{s^2} + \frac{\delta^2}{t^2} - 3\right)^{q_j}
\]  
(4.12)

and
\[
W_{\text{eff}}(F) = \min_{\lambda, \delta \in T} \left(\sum_i c_i \left(\frac{s^2}{\lambda^2} + \frac{t^2}{\lambda^2} - 3\right)^{p_i} + \sum_j d_j \left(\frac{\lambda^2}{s^2} + \frac{\delta^2}{t^2} - 3\right)^{q_j}\right).
\]  
(4.13)
In light of the constraint $\lambda, \delta \in T$, we have multiple cases.

**Case 1: Attained minimum.** We solve

$$\frac{\partial W_{\text{eff}}}{\partial \lambda} = \frac{\partial W_{\text{eff}}}{\partial \delta} = 0. \quad (4.14)$$

Since $c_i, d_j > 0$, $p_i, q_j \geq 1$, it is sufficient (and necessary in a generic sense) that

$$\frac{\partial}{\partial \lambda} \left( \frac{s^2}{\lambda^2} + \frac{r^2 A^2}{\lambda^2} + \frac{\delta^2}{r^2} \right) = 0, \quad (4.15)$$

$$\frac{\partial}{\partial \delta} \left( \frac{s^2}{\lambda^2} + \frac{r^2 A^2}{\lambda^2} + \frac{\delta^2}{r^2} \right) = 0, \quad (4.16)$$

$$\frac{\partial}{\partial \lambda} \left( \frac{\lambda^2}{s^2} + \frac{\delta^2 s^2}{r^2 A^2} + \frac{t^2}{\delta^2} \right) = 0, \quad (4.17)$$

$$\frac{\partial}{\partial \delta} \left( \frac{\lambda^2}{s^2} + \frac{\delta^2 s^2}{r^2 A^2} + \frac{t^2}{\delta^2} \right) = 0, \quad (4.18)$$

or

$$\frac{\partial W}{\partial \lambda} = 0 \rightarrow \frac{\lambda^4}{s^4} = \frac{\delta^2}{r^2}, \quad \frac{\partial W}{\partial \delta} = 0 \rightarrow \frac{\lambda^2}{s^2} = \frac{\delta^4}{t^4} \quad \iff \lambda = s, \delta = t \quad \Rightarrow W_{\text{eff}}(F) = 0. \quad (4.19)$$

This is possible if and only if $s, t \in T$. Recalling that $W^{qc} = 0$ when $s, t \in T$, we conclude

$$W_{\text{eff}}(F) = W^{qc}(F) \quad \text{in } L. \quad (4.20)$$

**Case 2: $t > r^{1/6}$.** We set $\delta = r^{1/6}$ and solve

$$\frac{\partial W_{\text{eff}}}{\partial \lambda} = 0. \quad (4.21)$$

Arguing as before, we conclude

$$\frac{\lambda}{s} = \frac{r^{1/12}}{t^{1/2}} \quad (4.22)$$

which implies

$$W_{\text{eff}}(F) = \sum_i c_i \left( \frac{2 t r^{1/6}}{r^{1/6}} + \frac{t^{1/3}}{t^2} - 3 \right)^{p_i} + \sum_j d_j \left( \frac{2^{r^{1/6}}}{t} + \frac{t^2}{r^{1/3}} - 3 \right)^{q_j}. \quad (4.23)$$

Note that this coincides with the expression for $W^{qc}$ in $M$. However, for $\delta = r^{1/6}$ and $\lambda$ according to (4.22), $\lambda, \delta \in T$ if and only if $1 \leq s/t^{1/2} \leq r^{1/4}$ or $t \leq s^2 \leq r^{1/2}t$. By assumption, $t > r^{1/6}$. So this is the region $M$. We conclude,

$$W_{\text{eff}}(F) = W^{qc}(F) \quad \text{in } M. \quad (4.24)$$
**Case 3:** \( s^2 > r^{1/2}, \ t > r^{1/6} \). Note that this is the region \( S \). We set \( \lambda = r^{1/3}, \delta = r^{1/6} \), and it is easy to verify that

\[
W_{\text{eff}}(F) = W^{\text{qe}}(F) \quad \text{in } S.
\]  

(4.25)

\[\square\]

**Constitutive relation**

The effective or relaxed energy (4.8) is obtained by assuming that the microstructure evolves instantaneously to minimize the energy. However, microstructure evolves according to a kinetic process which is dissipative. Further, some domains may be locally pinned, and this introduces a hardening energy. Finally, there is viscosity associated with the polymer network. All of these considerations motivate the following constitutive relation. We describe this for the isothermal situation where the temperature is fixed. However, this is easily generalized to a general temperature-dependent situation.

We assume that the state of a non-ideal isotropic-genesis polydomain nematic elastomer is described by the deformation gradient \( F \), internal variables \( \lambda, \delta \), and temperature \( T \). We postulate that the stored energy density of a non-ideal isotropic-genesis polydomain nematic elastomer is given by

\[
W(F, \lambda, \delta, T) = W_{\text{PNE}}(F, \lambda, \delta, T) + W_h(\lambda, \delta, T),
\]  

(4.26)

where

\[
W_{\text{PNE}}(F, \lambda, \delta, T) = \sum_i c_i \left( \frac{s^2}{\lambda^2} + \frac{t^2 \lambda^2}{\delta^2 s^2} + \frac{\delta^2}{t^2} - 3 \right)^{p_i} + \sum_j d_j \left( \frac{\lambda^2}{s^2} + \frac{\delta^2 s^2}{t^2 \lambda^2} + \frac{t^2}{\delta^2} - 3 \right)^{q_j}
\]  

(4.27)

as before, and

\[
W_h(\lambda, \delta, T) = C \frac{\delta - 1}{(r^{1/6} - \delta)^k}
\]  

(4.28)

is the hardening energy. The form of the hardening is chosen to penalize \( \delta \rightarrow r^{1/6} \), i.e. the completion of the polydomain-to-monodomain transition. The moduli \( c_i, d_i \) as well as the parameter \( r \) may depend on temperature.

The Cauchy stress is given by

\[
\sigma(F, \lambda, \delta) = -p I + \frac{\partial W_{\text{PNE}}}{\partial F}(F, \lambda, \delta) F^\top + \beta d,
\]  

(4.29)
where $p$ is an unknown pressure, $d = \frac{1}{2}(\dot{F}F^{-1} + F^{-T}\dot{F}^T)$ is the rate-of-deformation tensor, and $\beta$ is the viscosity. The microstructure parameters $\lambda, \delta$ evolve according to the equations

$$\begin{align*}
\dot{\lambda} &= -\alpha_\lambda \frac{\partial}{\partial \lambda} (W_{\text{PNE}} + W_h) \\
\dot{\delta} &= -\alpha_\delta \frac{\partial}{\partial \delta} (W_{\text{PNE}} + W_h)
\end{align*}$$
subject to $\lambda, \delta \in \mathcal{T}$. \hspace{1cm} (4.30)

Note that $r$, and hence $\mathcal{T}$, depends on temperature. It is convenient to introduce a rate-of-dissipation potential for the microstructure evolution

$$D(\dot{\lambda}, \dot{\delta}, d) = \frac{1}{2} \left( \alpha_\lambda |\dot{\lambda}|^2 + \alpha_\delta |\dot{\delta}|^2 \right).$$ \hspace{1cm} (4.31)

If we discretize the evolution equation by a backward Euler (implicit) time discretization, we can update the variables as

$$\lambda^{n+1}, \delta^{n+1} = \arg\min_{\lambda^{n+1}, \delta^{n+1} \in \mathcal{T}} \left[ W_{\text{PNE}}(F, \lambda^{n+1}, \delta^{n+1}, T) + W_h(\lambda^{n+1}, \delta^{n+1}, T) \right.$$

$$\left. + \Delta t D \left( \frac{\lambda^{n+1} - \lambda^n}{\Delta t}, \frac{\delta^{n+1} - \delta^n}{\Delta t} \right) \right].$$ \hspace{1cm} (4.32)

**Useful calculation** It is useful to compute the rotation associated with the minimization in (4.8). Let $\lambda_i$ be the principal values of $F$ with $\lambda_1 \geq \lambda_2 \geq \lambda_3$. Let

$$C = F^TF = \sum_{i=1}^{3} \lambda_i^2 N_i \otimes N_i, \quad b = FF^T = \sum_{i=1}^{3} \lambda_i^2 n_i \otimes n_i.$$ \hspace{1cm} (4.33)

It follows that

$$F = \sum_{i=1}^{3} \lambda_i n_i \otimes N_i.$$ \hspace{1cm} (4.34)

Set

$$G_0 = \sum_{i=1}^{3} \xi_i e_i \otimes e_i,$$ \hspace{1cm} (4.35)

where $\xi_1 \geq \xi_2 \geq \xi_3$. Therefore,

$$F^T Q G_0^{-1} Q^T F = \sum_{i,j,k} \lambda_i \lambda_k \xi_j^{-1} (n_i \cdot Q e_j)(n_k \cdot Q e_j) N_i \otimes N_k,$$ \hspace{1cm} (4.36)

$$(F^T Q G_0^{-1} Q^T F)^{-T} = \sum_{i,j,k} \lambda_i^{-1} \lambda_k^{-1} \xi_j (n_i \cdot Q e_j)(n_k \cdot Q e_j) N_i \otimes N_k.$$ \hspace{1cm} (4.37)
Now, in light of (4.11), maximizing $f(A)$ over $A$ is equivalent to maximizing the trace of $A$ and cof($A$). Examining the above, we see that we maximize the trace of $F^T Q G_0^{-1} F$, cof($F^T Q G_0^{-1} F$) = $(F^T Q G_0^{-1} F)^{-T}$ exactly when

$$n_i = Q e_i, \quad i = 1, 2, 3. \quad (4.38)$$

Thus, the maximizing $Q$ is

$$Q = \sum_{i=1}^{3} n_i \otimes e_i \quad (4.39)$$

so that for the maximizing $Q$,

$$G = \sum_{i=1}^{3} \xi_i n_i \otimes n_i, \quad (4.40)$$

i.e. $G$ shares an eigenbasis with $b$ and

$$F^T Q G_0^{-1} Q^T F = \sum_{i=1}^{3} \lambda_i^2 \xi_i^{-1} N_i \otimes N_i, \quad (4.41)$$

$$F^T Q G_0^{-1} Q^T F)^{-T} = \sum_{i=1}^{3} \lambda_i^{-2} \xi_i^2 N_i \otimes N_i. \quad (4.42)$$

### 4.3 Validation of the model

We now validate the model against the experiments of Tokumoto, Takabe, and Urayama, as reported in Tokumoto et al. [71]. They subjected sheets of isotropic-genesis polydomain nematic elastomers to biaxial extension while leaving the faces of the sheet traction-free, i.e. deformations of the form

$$F = \begin{pmatrix} \lambda_x & 0 & 0 \\ 0 & \lambda_y & 0 \\ 0 & 0 & (\lambda_x \lambda_y)^{-1} \end{pmatrix} \quad (4.43)$$

where $\lambda_x, \lambda_y$ are imposed stretches. In all their experiments $\lambda_x \lambda_y > 1$, and we assume that $\lambda_x > \lambda_y$. Therefore,

$$s = \lambda_x, t = \lambda_x \lambda_y. \quad (4.44)$$

We neglect viscosity ($\beta = 0$), and so the Cauchy stress $\sigma$ is

$$\begin{pmatrix} -p + \mu_1 \frac{\partial^2 \lambda_x}{\partial^2 \lambda_x} + \mu_2 \left( \frac{\partial^2 \lambda_x}{\partial^2 \lambda_x} + \frac{\partial^2 \lambda_y}{\partial^2 \lambda_y} \right) & 0 & 0 \\ 0 & -p + \mu_1 \frac{\partial^2 \lambda_x}{s^2 \delta^2} + \mu_2 \left( \frac{\partial^2 \lambda_x}{s^2 \delta^2} + \frac{\partial^2 \lambda_y}{s^2 \delta^2} \right) & 0 \\ 0 & 0 & -p + \mu_1 \frac{\partial^2 \lambda_x}{t^2 \lambda_x^2} + \mu_2 \left( \frac{\partial^2 \lambda_x}{t^2 \lambda_x^2} + \frac{\partial^2 \lambda_y}{t^2 \lambda_x^2} \right) \end{pmatrix}.$$
Since the faces of the sheet are traction-free, \( \sigma_{33} = 0 \). It follows,

\[
p = \mu_1 \frac{\delta^2}{t^2} + \mu_2 \left( \frac{\lambda^2}{s^2} + \frac{s^2 \delta^2}{t^2 \lambda^2} \right),
\]

and the two non-zero components of stress are

\[
\sigma_{11} = \mu_1 \left( \frac{s^2}{\lambda^2} - \frac{\delta^2}{t^2} \right) + \mu_2 \left( \frac{t^2}{\delta^2} - \frac{\lambda^2}{s^2} \right), \quad (4.46)
\]
\[
\sigma_{22} = \mu_1 \left( \frac{t^2 \lambda^2}{s^2 \delta^2} - \frac{\delta^2}{t^2} \right) + \mu_2 \left( \frac{t^2}{\delta^2} - \frac{s^2 \delta^2}{t^2 \lambda^2} \right). \quad (4.47)
\]

It remains to solve for the evolution of the internal variables \( \lambda, \delta \). We do so by implementing (4.32) using MATLAB. We use the Bladon-Warner-Terentjev form (generalization of the neo-Hookean), where \( c_1 = \mu_1 / 2, p_1 = 1 \) and \( c_i, d_j = 0 \) for \( i > 1, j \geq 1 \). Table 4.1 summarizes the material parameters used. In MATLAB, a gradient descent code was implemented to fit the various parameters in the theoretical model to the experimental data.

**Table 4.1: Material properties used in MATLAB simulations**

<table>
<thead>
<tr>
<th>Property</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Shear modulus</td>
<td>( \mu_1 = 4.93752 \cdot 10^4 ) Pa</td>
</tr>
<tr>
<td>LCE anisotropy parameter</td>
<td>( r = 9.1393 )</td>
</tr>
<tr>
<td>Hardening coefficient</td>
<td>( C = 298 ) Pa</td>
</tr>
<tr>
<td>Hardening exponent</td>
<td>( k = 2 )</td>
</tr>
<tr>
<td>Dissipation property</td>
<td>( \alpha_{\delta} = 2.1838 \cdot 10^7 ) Pa</td>
</tr>
<tr>
<td>Dissipation property</td>
<td>( \alpha_{\lambda} = \alpha_{\delta} = 2.1838 \cdot 10^5 ) Pa</td>
</tr>
<tr>
<td>Exponent in dissipation potential ( D )</td>
<td>( p = 2 )</td>
</tr>
</tbody>
</table>

We explore four specific deformations in the following sections: uniaxial (U) extension, planar extension (PE), equibiaxial (EB) extension, and unequal-biaxial (UB) extension. The general deformation gradient, following from Equation 4.43 and 4.44, is \( F = \text{diag}(s, t/s, 1/t) \). Figs. 4.2 – 4.7 are plots for each individual deformation with more detail. In each figure, the top left subplot depicts the stress-stretch curve, the top right plot shows the (red) path that the internal variables take through the (black) triangular region \( T \), the bottom left plot shows the internal variables as a function of stretch, and the bottom right plot shows the energy density as a function of stretch.

**Uniaxial extension (U)** In the uniaxial case, the body is subjected to uniaxial stress in the \( x \)-direction and traction-free in the \( y \)- and \( z \)-directions. Thus, \( \sigma_{22} = \sigma_{33} = 0 \). The relationship between the stretches is \( t = \sqrt{s} \). Thus, the deformation gradient is \( F = \text{diag}(s, 1/\sqrt{s}, 1/\sqrt{s}) \).
Planar extension (PE)  In the planar extension deformation, the stretch in the $y$-direction is fixed at a ratio of 1, yielding $\lambda_y = 1$, and the body is traction-free in the $z$-direction. Thus, we have $t = s$, and the deformation gradient is $F = \text{diag}(s, 1, 1/s)$. 

Figure 4.2: Uniaxial extension.

Figure 4.3: Planar extension.
**Equibiaxial extension (EB)**  In the equibiaxial deformation, the stretch in the $x$-direction and $y$-direction are equal, and the body is traction-free in the $z$-direction. Thus, $s = \frac{t}{s}$ or $t = s^2$, and the deformation gradient is $F = \text{diag}(s, s, 1/s^2)$.

![Figure 4.4: Equibiaxial extension.](image)

**Unequal biaxial extension (UB)**  In the case of the unequal-biaxial extension, the body is traction-free in the $z$-direction. The experiments are performed by fixing the ratio of $\frac{e_{x}}{e_{y}} = \frac{\lambda_{x}-1}{\lambda_{y}-1}$ to be a constant $\beta$, where $\beta$ has values equal to $5/3, 5/2, 5/1$. (Note that $\beta = 1$ recovers the equibiaxial case.) This means that we have $\frac{s-1}{t/s-1} = \beta$, so $t = \frac{s}{\beta} (s - 1 + \beta)$. Thus, the deformation gradient is $F = \text{diag} \left( s, \frac{s-1+\beta}{\beta}, \frac{\beta}{s(s-1+\beta)} \right)$. The figures can be seen in Figures 4.5–4.7.
Figure 4.5: Unequal biaxial extension, $\beta = 5/3$.

Figure 4.6: Unequal biaxial extension, $\beta = 5/2$. 
Summary: liquid-like behavior  Figures 4.8 and 4.9 show a comparison of the stress-stretch data, plotted for the experimental data from Urayama’s research group [71] and the theoretical model for various deformations.
**Hysteresis** Finally, Figures 4.10a and 4.10b show the load/unload curve for the U and PE deformations. The model is able to capture energy dissipation between the load and unload curves, depicted by the hysteresis between the dashed and solid curves.

![Theoretical stress plot](image)

*Figure 4.9: Theoretical stress, plotted as a function of $\lambda_z$."

**4.4 Implementation in ABAQUS**

We now discuss the model’s implementation in a custom user-material, or UMAT, for use in ABAQUS. The code for the UMAT, written in FORTRAN, can be found in Appendix D.1. It is useful for numerical purposes to consider a compressible model. Further, our validation showed good agreement with the neo-Hookean. So we take the strain...
energy density to be:

\[
\bar{W} = \frac{\mu}{2} \left[ J^{-2/3} \text{tr} \left( F^T G^{-1} F \right) - 3 \right] + \frac{K}{2} (\ln J)^2 ,
\]  

(4.48)

where \( F \) is the deformation gradient, \( J = \text{det} \, F \) is the determinant of the deformation gradient, and \( b = FF^T \) is the left Cauchy-Green tensor. \( G \) is the tensor of internal variables that follow an evolution law and shares the same principal basis as \( b \). In the following sections, we show that the energy density can be rewritten in terms of a tensor \( \tilde{G} \) which shares the same principal basis as \( C \), the right Cauchy-Green tensor:

\[
W = \frac{\mu}{2} \left[ J^{-2/3} \text{tr} \left( F \tilde{G}^{-1} F^T \right) - 3 \right] + \frac{K}{2} (\ln J)^2 .
\]  

(4.49)

**Notation**

We have the right stretch tensor,

\[
U = \sum_{i=1}^{3} \lambda_i N_i \otimes N_i = \sqrt{C}
\]  

(4.50)

and the right Cauchy-Green tensor

\[
C = \sum_{i=1}^{3} \lambda_i^2 N_i \otimes N_i.
\]  

(4.51)

From polar decomposition, we have

\[
F = RU = VR,
\]  

(4.52)

where \( U \) is the right stretch tensor as above, \( R \in SO(3) \), and \( U \in GL(3) \) symmetric and positive-definite. Because \( U \) is symmetric and real, we can write its eigendecomposition as

\[
U = Q \Lambda Q^T,
\]  

(4.53)

where the orthogonal matrix \( Q \) is

\[
Q = \begin{pmatrix}
N_1 & N_2 & N_3
\end{pmatrix}
\]  

(4.54)

and

\[
\Lambda = \begin{pmatrix}
\lambda_1 \\
\lambda_2 \\
\lambda_3
\end{pmatrix}
\]  

(4.55)
The right Cauchy-Green tensor is

\[ C = U^2 = Q \Lambda^2 Q^\top \]

\[ = \sum_{i=1}^{3} \lambda_i^2 N_i \otimes N_i. \]  

(4.56)  

(4.57)  

(4.58)

The left Cauchy-Green tensor is

\[ b = V^2 = R Q \Lambda^2 Q^\top R^\top \]

\[ = \sum_{i=1}^{3} \lambda_i^2 n_i \otimes n_i, \]  

(4.59)  

(4.60)  

(4.61)

where the eigenvectors of \( b \) are related to the eigenvectors of \( C \) through \( n = RN \).

The internal variable tensor \( G \) shares the same principal basis as \( b \):

\[ G = R Q \Xi Q^\top R^\top \]

\[ = \sum_{i=1}^{3} \xi_i^2 n_i \otimes n_i, \]  

(4.62)  

(4.63)

where

\[ \Xi = \begin{pmatrix} \xi_1 & \lambda^2 \\ \xi_2 & \delta^2 / \lambda^2 \\ \xi_3 & 1 / \delta^2 \end{pmatrix}. \]  

(4.64)

A related internal variable tensor \( \tilde{G} \) shares the same principal basis as \( C \):

\[ \tilde{G} = Q \Xi Q^\top \]

\[ = \sum_{i=1}^{3} \xi_i^2 N_i \otimes N_i. \]  

(4.65)  

(4.66)

**Rewritten energy density**

The original energy density for our constitutive model was

\[ W = \frac{\mu}{2} \left\{ J^{-2/3} \text{tr} \left( F^\top G^{-1} F \right) - 3 \right\} + \frac{K}{2} (\ln J)^2. \]  

(4.67)
We can rewrite the energy as a function of $\tilde{G}$:

$$\begin{align*}
W &= \frac{\mu}{2} \left[ J^{-2/3} \operatorname{tr} \left( G^{-1} b \right) - 3 \right] + \frac{\kappa}{2} (\ln J)^2 \\
&= \frac{\mu}{2} \left[ J^{-2/3} \operatorname{tr} \left( RQ \tilde{Q}^{-1} Q^\top R^\top RQA^2 Q^\top R^\top \right) - 3 \right] + \frac{\kappa}{2} (\ln J)^2 \\
&= \frac{\mu}{2} \left[ J^{-2/3} \operatorname{tr} \left( Q \tilde{Q}^{-1} Q^\top \Lambda^2 Q^\top \right) - 3 \right] + \frac{\kappa}{2} (\ln J)^2 \\
&= \frac{\mu}{2} \left[ J^{-2/3} \operatorname{tr} \left( F \tilde{G}^{-1} F^\top \right) - 3 \right] + \frac{\kappa}{2} (\ln J)^2.
\end{align*}$$

We can also rewrite the energy as a function of the principal values $Z$:

$$\begin{align*}
W &= \frac{\mu}{2} \left[ J^{-2/3} \operatorname{tr} \left( RQ \tilde{Q}^{-1} Q^\top R^\top RQA^2 Q^\top R^\top \right) - 3 \right] + \frac{\kappa}{2} (\ln J)^2 \\
&= \frac{\mu}{2} \left[ J^{-2/3} \operatorname{tr} \left( \tilde{Q}^{-1} \Lambda^2 \right) - 3 \right] + \frac{\kappa}{2} (\ln J)^2 \\
&= \frac{\mu}{2} \left[ J^{-2/3} \left( \frac{\lambda_1^2}{\xi_1^2} + \frac{\lambda_2^2}{\xi_2^2} + \frac{\lambda_3^2}{\xi_3^2} \right) - 3 \right] + \frac{\kappa}{2} (\ln J)^2.
\end{align*}$$

or

$$\begin{align*}
W &= \frac{\mu}{2} \left[ J^{-2/3} \left( \frac{\lambda_1^2}{\xi_1^2} + \frac{\lambda_2^2}{\xi_2^2} + \frac{\lambda_3^2}{\xi_3^2} \right) - 3 \right] + \frac{\kappa}{2} (\ln J)^2,
\end{align*}$$

where $J = \det F = \lambda_1 \lambda_2 \lambda_3$.

**Adding in viscosity**

Here, we note that we have added a viscous part of the stress. We assume an additive decomposition of the Piola-Kirchhoff stress $P$ into an elastic portion and a viscosity portion:

$$P = P^e + P^v,$$

where

$$P^e = \frac{\partial W}{\partial F},$$

and $P^v = \beta J dF^\top$ is the viscosity portion. Thus, we have the Cauchy stress:

$$\sigma = \sigma^e + \sigma^v$$

$$= \frac{1}{J} \frac{\partial W}{\partial F} F^\top + \beta d.$$

**Stress measures**

**Second Piola-Kirchhoff stress**

Let us calculate the second Piola-Kirchhoff stress. Let a symmetric, 3x3 tensor $C$ have eigenvalues $\mu_i$ and eigenvectors $N_i$ so that

$$CN_i = \mu_i N_i \quad \text{(no sum over $i$)}.$$
Then, from [37],
\[
\frac{\partial \mu_i}{\partial C} = N_i \otimes N_i. \tag{4.81}
\]
Here, the eigenvalues of the right Cauchy-Green tensor are \( \mu_i = \lambda_i^2 \). Thus, we obtain
\[
S = 2 \frac{\partial W}{\partial C} \tag{4.82}
\]
\[
= 2 \left( \frac{\partial W}{\partial \lambda_1^2} \frac{\partial \lambda_1^2}{\partial C} + \frac{\partial W}{\partial \lambda_2^2} \frac{\partial \lambda_2^2}{\partial C} + \frac{\partial W}{\partial \lambda_3^2} \frac{\partial \lambda_3^2}{\partial C} \right) \tag{4.83}
\]
\[
= 2 \left( \frac{\partial W}{\partial \lambda_1^2} N_1 \otimes N_1 + \frac{\partial W}{\partial \lambda_2^2} N_2 \otimes N_2 + \frac{\partial W}{\partial \lambda_3^2} N_3 \otimes N_3 \right) \tag{4.84}
\]
\[
= \mu J^{-2/3} \left( \frac{1}{\xi_1^2} N_1 \otimes N_1 + \frac{1}{\xi_2^2} N_2 \otimes N_2 + \frac{1}{\xi_3^2} N_3 \otimes N_3 \right) \tag{4.85}
\]
\[
- \frac{\mu}{3} J^{-5/3} \left( \frac{\lambda_1^2}{\xi_1^2} N_1 \otimes N_1 + \frac{\lambda_2^2}{\xi_2^2} N_2 \otimes N_2 + \frac{\lambda_3^2}{\xi_3^2} N_3 \otimes N_3 \right) \tag{4.86}
\]
\[
+ \kappa \ln \frac{J}{\lambda_1} \left( \frac{\lambda_2 \lambda_3}{\lambda_1} N_1 \otimes N_1 + \frac{\lambda_1 \lambda_3}{\lambda_2} N_2 \otimes N_2 + \frac{\lambda_1 \lambda_2}{\lambda_3} N_3 \otimes N_3 \right) \tag{4.87}
\]
Recognizing that
\[
\tilde{G}^{-1} = \frac{1}{\xi_1^2} N_1 \otimes N_1 + \frac{1}{\xi_2^2} N_2 \otimes N_2 + \frac{1}{\xi_3^2} N_3 \otimes N_3 \tag{4.88}
\]
\[
\text{tr}(\tilde{G}^{-1}C) = \frac{\lambda_1^2}{\xi_1^2} + \frac{\lambda_2^2}{\xi_2^2} + \frac{\lambda_3^2}{\xi_3^2} \tag{4.89}
\]
\[
JC^{-1} = \frac{\lambda_2 \lambda_3}{\lambda_1} N_1 \otimes N_1 + \frac{\lambda_1 \lambda_3}{\lambda_2} N_2 \otimes N_2 + \frac{\lambda_1 \lambda_2}{\lambda_3} N_3 \otimes N_3, \tag{4.90}
\]
we are left with
\[
S = \mu J^{-2/3} \tilde{G}^{-1} - \frac{\mu}{3} J^{-5/3} \text{tr}(\tilde{G}^{-1}C)JC^{-1} + \kappa \ln \frac{J}{\lambda_1} JC^{-1}, \tag{4.91}
\]
or
\[
S = \mu J^{-2/3} \left[ \tilde{G}^{-1} - \frac{1}{3} \text{tr}(\tilde{G}^{-1}C)C^{-1} \right] + \kappa (\ln J)C^{-1}. \tag{4.92}
\]
Note that we also reached this stress expression by computing the tensor derivative as follows:
\[
S = 2 \frac{\partial W(C)}{\partial C}, \tag{4.93}
\]
where
\[
W(C) = \frac{\mu}{2} \left[ (\det C)^{-1/3} \text{tr}(\tilde{G}^{-1}C) - 3 \right] + \frac{\kappa}{8} [\ln(\det C)]^2 \tag{4.94}
\]
with \( \frac{\partial \tilde{G}}{\partial C} = 0 \).
First Piola-Kirchhoff stress

The first Piola-Kirchhoff stress can be computed as follows:

\[ P = FS \]  

(4.95)

\[ P = \mu J^{-2/3} \left[ F \tilde{G}^{-1} - \frac{1}{3} \text{tr} \left( \tilde{G}^{-1} C \right) F^{-\top} \right] + \kappa (\ln J) F^{-\top}. \]  

(4.96)

Cauchy stress

The Cauchy stress expression is needed for ABAQUS. We find

\[ \sigma = \frac{1}{J} PF^{\top} = \frac{1}{J} FSF^{\top} \]  

(4.97)

\[ \sigma = \mu J^{-5/3} \left[ F \tilde{G}^{-1} F^{\top} - \frac{1}{3} \text{tr}(\tilde{G}^{-1} C) I \right] + \kappa \frac{\ln J}{J} I. \]  

(4.98)

Note that when \( \tilde{G} = I \), the Cauchy stress recovers the neo-Hookean stress, as expected.

Kirchhoff stress

The Kirchhoff stress is simply

\[ \tau = J\sigma = FSF^{\top} \]  

(4.99)

\[ \tau = \mu J^{-2/3} \left[ F \tilde{G}^{-1} F^{\top} - \frac{1}{3} \text{tr}(\tilde{G}^{-1} C) I \right] + \kappa (\ln J) I. \]  

(4.100)

Derivation of the material Jacobian, DDSDDE

Jaumann rate of the Kirchhoff stress

The Jaumann rate of the Kirchhoff stress is

\[ \dot{\tau}^{(JK)} = \dot{\tau} - \omega \tau + \tau w, \]  

(4.101)
where \( w = \frac{1}{2} (\ell - \ell^\top) \) is the spin tensor. In symbolic form, we have

\[
\begin{align*}
\psi^{(JK)} \tau &= -\frac{2\mu}{3} J^{-2/3} (\text{tr} \, \ell) \left[ F \hat{G}^{-1} F^\top - \frac{1}{3} \text{tr} \left( \hat{G}^{-1} C \right) I \right] \\
&\quad - \frac{\mu}{3} J^{-2/3} \left[ \text{tr} \left( \hat{G}^{-1} C \right) \delta_{ij} \right] \\
&\quad + \mu J^{-2/3} \left[ dF \hat{G}^{-1} F^\top + F \hat{G}^{-1} F^\top d \right] \\
&\quad + \kappa \left( \text{tr} \right) \ell I \\
&\quad + \mu J^{-2/3} \left[ F \hat{G}^{-1} F^\top - \frac{1}{3} \text{tr} \left( \hat{G}^{-1} C \right) I \right].
\end{align*}
\]

\[ (4.102) \]

**Consistent Jacobian, DDSDDE**

The consistent Jacobian matrix (called DDSDDE in ABAQUS), is defined as

\[
\begin{align*}
\psi^{(JK)} \tau &= J(\text{DDSDDE}) : d, \\
\psi^{(JK)} \tau_{ij} &= J \text{ DDSDDE}_{ijkl} d_{kl}. 
\end{align*}
\]

or

We can rewrite \( \psi^{(JK)} \tau \) in the following way so that the DDSDDE expression is easily extricated. Note that we will make use of Section B.2.

\[
\begin{align*}
\psi^{(JK)} \tau_{ij} &= -\frac{2\mu}{3} J^{-2/3} \left[ \left( F \hat{G}^{-1} F^\top \right)_{ij} \delta_{kl} - \frac{1}{3} \text{tr} \left( \hat{G}^{-1} C \right) \delta_{ij} \delta_{kl} \right] d_{kl} \\
&\quad - \frac{2\mu}{3} J^{-2/3} \left[ \left( F \hat{G}^{-1} F^\top \right)_{kl} \delta_{ij} \right] d_{kl} \\
&\quad + \frac{\mu}{2} J^{-2/3} \left[ \left( F \hat{G}^{-1} F^\top \right)_{ij} \delta_{ik} + \left( F \hat{G}^{-1} F^\top \right)_{ik} \delta_{jl} + \left( F \hat{G}^{-1} F^\top \right)_{il} \delta_{jk} + \left( F \hat{G}^{-1} F^\top \right)_{jk} \delta_{il} \right] d_{kl} \\
&\quad + \kappa \left[ \delta_{ij} \delta_{kl} \right] d_{kl} \\
&\quad + \mu J^{-2/3} \left[ \left( F \hat{G}^{-1} F^\top \right)_{ij} - \frac{1}{3} \text{tr} \left( \hat{G}^{-1} C \right) \delta_{ij} \right].
\end{align*}
\]

\[ (4.105) \]

For the last line, we have used the auxiliary Section B.3, where

\[
A_{ij} = \left[ \left( F \hat{G}^{-1} F^\top \right)_{ij} - \frac{1}{3} \text{tr} \left( \hat{G}^{-1} C \right) \delta_{ij} \right].
\]

\[ (4.106) \]
Thus, the material Jacobian, from Equation 4.104, is
\[
\text{DDSDDE}_{ijkl} = -\frac{2\mu}{3} J^{-5/3} \left[ \left( F\tilde{G}^{-1} F^\top \right)_{ij} \delta_{kl} + \left( F\tilde{G}^{-1} F^\top \right)_{kl} \delta_{ij} \right] \\
+ \frac{2\mu}{9} J^{-5/3} \text{tr} \left( \tilde{G}^{-1} C \right) \delta_{ij} \delta_{kl} \\
+ \frac{\mu}{2} J^{-5/3} \left[ \left( F\tilde{G}^{-1} F^\top \right)_{ij} \delta_{ik} + \left( F\tilde{G}^{-1} F^\top \right)_{ik} \delta_{ij} \right] \\
+ \frac{\mu}{2} J^{-5/3} \left[ \left( F\tilde{G}^{-1} F^\top \right)_{ij} \delta_{jk} + \left( F\tilde{G}^{-1} F^\top \right)_{jk} \delta_{il} \right] \\
+ \mu J^{-5/3} \left[ \frac{1}{|d|^2} (A_{ij} d_{kl} + d_{ij} A_{kl}) - \frac{1}{|d|^4} A_{mn} d_{mn} d_{ij} d_{kl} \right].
\]

Note that we can check that the material Jacobian satisfies the symmetries
\[
\text{DDSDDE}_{ijkl} = \text{DDSDDE}_{klij} = \text{DDSDDE}_{ijlk}.
\]

We can also confirm that when \( \tilde{G} = I \) (and therefore \( \dot{\tilde{G}} = 0 \)), we recover the neo-Hookean material Jacobian.

Also note that in approximating the spatial velocity gradient \( \ell \) and consequently the rate-of-deformation tensor \( d \) and spin tensor \( w \), we use the approximation from Weber, Anand (1999) [78]:
\[
\ell_n = \frac{1}{\Delta t} \left( F_n F_{n-1}^{-1} - I \right).
\]

We considered other approximations; the first was a finite difference approximation:
\[
\dot{F}_n = \frac{F_n - F_{n-1}}{\Delta t},
\]
and the second used the increment DSTRAN, as suggested in Nguyen and Waas [52]:
\[
d = \frac{\text{DSTRAN}}{\Delta t}.
\]

**Optimization using NLOpt**

Inside the UMAT, we will be using an external optimization algorithm called NLOpt (documentation here) to conduct the constrained optimization of the internal variables. For algorithms that use a gradient-based method, we need to compute the derivatives of the objective function with respect to the internal variables \( \lambda \) and \( \delta \).

Recall the minimization problem from Equation 4.32. Written explicitly, we have
\[(\lambda, \delta) = \arg \inf_{(\lambda, \delta) \in T} \left\{ \frac{\mu}{2} \left[ J^{-2/3} \left( \frac{\lambda_1^2}{\xi_1^2} + \frac{\lambda_2^2}{\xi_2^2} + \frac{\lambda_3^2}{\xi_3^2} \right) - 3 \right] + \frac{\kappa}{2} (\ln J)^2 \right\} - C \frac{\delta - 1}{(r^{1/6} - \delta)^k} + \Delta t \left( \frac{\alpha_t}{2} \dot{\lambda}^p + \frac{\alpha_\delta}{2} \delta^p \right) \}, \quad (4.112)\]

where \(\lambda_i\) \((i = 1 : 3)\) are the principal stretches of \(\mathbf{F}\), \(J = \det \mathbf{F} = \lambda_1 \lambda_2 \lambda_3\), \(\mathbf{G} = \mathbf{Q} \mathbf{\Xi} \mathbf{Q}^T\), and \(\xi_1 = \lambda^2, \xi_2 = \frac{\delta^2}{\lambda^2}, \xi_3 = \frac{1}{\delta^2}\). We wish for the internal variables to evolve at different speeds, so the parameters of dissipation must satisfy the relation

\[\alpha_t \ll \alpha_\delta.\]  

\[\text{Strain rate}\]

This is the loading for a uniaxial block. We begin with the equation relating stretch and strain:

\[\varepsilon = \frac{L - L_0}{L_0} = \frac{\Delta L}{L_0} = \frac{L}{L_0} - 1 = \lambda - 1.\]  

We choose a reference length for the model \(L_0\), e.g. in some simulations we have \(L_0 = 0.01\text{m}, \text{or 1 cm}\). We choose the desired final strain \(\varepsilon_f = 3\), or a desired final stretch of \(\lambda_f = \varepsilon_f + 1 = 4\). Then the distance that the block needs to stretch is \(\Delta L = \varepsilon_f L_0\). The velocity that the block needs to move is determined by \(v = \dot{\varepsilon} L_0\), or in other words the strain rate is determined by \(\dot{\varepsilon} = \frac{v}{L_0}\). The total time for the simulation is \(t^* = \frac{\Delta L}{v}\), and the stretch as a function of time \(\lambda(t)\) is

\[\lambda(t) = \dot{\varepsilon} t + 1 = \frac{v}{L_0} t + 1 = \frac{\varepsilon_f}{t^*} t + 1.\]  

\[\text{Eigenvector/eigenvalue coding}\]

We find the eigendecomposition of \(\mathbf{C}\), the right Cauchy-Green tensor using a subroutine called Jacobi. We find the eigenvalues of \(\mathbf{C}\) and then order them from greatest to least (using a bubble sort): \(\lambda_1^2 \geq \lambda_2^2 \geq \lambda_3^2\), and the square root of the eigenvalues are collected into the principal stretch matrix is

\[\mathbf{\Lambda} = \begin{pmatrix} \lambda_1 \\ \lambda_2 \\ \lambda_3 \end{pmatrix}.\]  

(4.116)
We also order the corresponding normalized eigenvectors \( N_1, N_2, N_3 \). The eigenvectors are collected into the orthogonal eigenvector matrix \( Q \):

\[
Q = \begin{pmatrix} N_1 & N_2 & N_3 \end{pmatrix}.
\]

Then \( C = QAQ^T \). Recall the energy density (where \( J = \det F = \lambda_1 \lambda_2 \lambda_3 \))

\[
W = \frac{\mu}{2} \left[ J^{-2/3} \left( \frac{\lambda_1^2}{\xi_1^2} + \frac{\lambda_2^2}{\xi_2^2} + \frac{\lambda_3^2}{\xi_3^2} \right) - 3 \right] + \frac{\kappa}{2} (\ln J)^2.
\]  

(4.118)

So each \( \lambda_i \) correlates with \( \xi_i, \ i = 1 : 3 \). Thus we have the internal variable matrix \( \dot{\mathbf{G}} = Q \Xi Q^T \), where \( Q \) is the same as Equation 4.117 and the eigenvalues \( \xi_1, \xi_2, \xi_3 \) correlate to \( \lambda_1 \geq \lambda_2 \geq \lambda_3 \):

\[
\Xi = \begin{pmatrix} \xi_1 & \xi_2 & \xi_3 \end{pmatrix} = \begin{pmatrix} \lambda_1^2 & \frac{\lambda_2^2}{\xi_2^2} & \frac{\lambda_3^2}{\xi_3^2} \end{pmatrix}.
\]

(4.119)

This allows us to compute the quantity \( F \dot{\mathbf{G}}^{-1} F^T \) and therefore the energy density.

### 4.5 Results from the ABAQUS implementation

#### Verification of the model

We compute the results for a single element undergoing the following deformations: uniaxial (U), planar extension (PE), equibiaxial (EB), and unequal biaxial (UB) with stretch ratios of 5/3, 5/2, and 5/1. In all of these cases, we have a match between the theoretical model, solved in MATLAB, and the ABAQUS simulation results. The parameters used in these results are the same as in Table 4.1, with additional parameters shown in Table 4.2. In each deformation, we see that the results match.

<p>|</p>
<table>
<thead>
<tr>
<th>Table 4.2: Additional material properties used in ABAQUS simulations</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bulk modulus</td>
</tr>
<tr>
<td>Viscosity</td>
</tr>
<tr>
<td>Proportionality constant for</td>
</tr>
<tr>
<td>( \dot{\mathbf{G}}^{-1} = \alpha C^{-1} )</td>
</tr>
<tr>
<td>( \kappa = 4.93752 \cdot 10^8 \text{ Pa} )</td>
</tr>
<tr>
<td>( \beta = 1 \cdot 10^6 \text{ Pa} \cdot \text{s} )</td>
</tr>
<tr>
<td>( \alpha = 5 \cdot 10^{-6} )</td>
</tr>
<tr>
<td>( \dot{\mathbf{G}}^{-1} = \alpha C^{-1} )</td>
</tr>
</tbody>
</table>

We repeated the calculations with multiple elements (e.g. cubes meshed with \( 5 \times 5 \times 5 \) and \( 10 \times 10 \times 10 \) elements) with the same boundary conditions and loading conditions as above and confirmed that the results agreed perfectly with the \( 1 \times 1 \times 1 \) simulations.
Figure 4.11: Uniaxial single-element simulation results plotted against theoretical results.

Figure 4.12: Planar extension single-element simulation results plotted against theoretical results.
Figure 4.13: Equibiaxial single-element simulation results plotted against theoretical results.

Figure 4.14: Unequal biaxial (with stretch ratio 5/3) single-element simulation results plotted against theoretical results.
Figure 4.15: Unequal biaxial (with stretch ratio 5/2) single-element simulation results plotted against theoretical results.

Figure 4.16: Unequal biaxial (with stretch ratio 5/1) single-element simulation results plotted against theoretical results.
Dependence upon strain rate and dissipation coefficients in uniaxial stretch

This section summarizes the results from single-element uniaxial stretch for three different strain rates—fast ($1 \times 10^{-2}$/s), medium ($1 \times 10^{-3}$/s), and slow ($1 \times 10^{-4}$/s)—and three different values of $\alpha_d$—big ($2.1838 \times 10^8$ Pa), medium ($2.1838 \times 10^7$ Pa), and small ($2.1838 \times 10^6$ Pa).

Figure 4.17: Uniaxial load and unload curves for varying $\alpha_d$ and fixed strain rates: (a) slow ($1 \times 10^{-4}$/s), (b) medium ($1 \times 10^{-3}$/s), and (c) fast ($1 \times 10^{-2}$/s).

As seen in Figure 4.18, the material response has correspondingly less hysteresis for slower strain rates, as is expected. We can also see that the smaller values of $\alpha_d$ correspond to less hysteresis as well.
Torsion of a cylinder

We move beyond the deformation of thin sheets or ribbons and study the bulk response of a solid cylinder of nematic elastomer under torsion, which have not been modeled before using FEA. The cylinder has height $H$, diameter $D$, and varying height-to-diameter ratios $H : D$, as shown in Table 4.3. We use a C3D8H element in ABAQUS with hybrid formulation, and the mesh is depicted in Figure 4.19.

Table 4.3: Dimensions of the cylinder under torsion

<table>
<thead>
<tr>
<th>$H : D$ ratio</th>
<th>Height, $H$ [m]</th>
<th>Diameter, $D$ [m]</th>
<th>Number of elements</th>
</tr>
</thead>
<tbody>
<tr>
<td>1:1</td>
<td>0.01</td>
<td>0.01</td>
<td>1652</td>
</tr>
<tr>
<td>2:1</td>
<td>0.02</td>
<td>0.01</td>
<td>3509</td>
</tr>
<tr>
<td>3:1</td>
<td>0.03</td>
<td>0.01</td>
<td>4859</td>
</tr>
</tbody>
</table>

The boundary conditions are such that the right face of the cylinder is entirely fixed in displacement and rotation in all three directions ($U_1 = U_2 = U_3 = 0$ and $VR_x = VR_y = VR_z = 0$ in ABAQUS) and the left face is zero entirely except for a prescribed angular velocity $\omega = 0.0026$ rad/s ($VR_z = 0.0026$ in ABAQUS). There is a tie constraint between the cylinder’s face and a rigid plate at each end of the cylinder.

The angle $DH$ describes the angle that one end of the cylinder twists with respect to the other end, as seen in Figure 4.20. We can define the strain rate at the outer rim as $\dot{\gamma}_{R_o}$, where $R_o$ is the half the diameter. For example, for $\omega = \dot{DH} = 0.0026$ rad/s and $H : D = 1 : 1$, the strain rate at the outer rim is 0.0013 rad/s.
Figure 4.19: Meshed cylindrical bodies for varying $H : D$ ratio.

Figure 4.20: Schematic of the twist $DH$ in the cylinder torsion deformation.

The moment and normal force obtained from the ABAQUS results are plotted in Figs. 4.21 and 4.22, separated by anisotropy parameter and $H : D$ ratio. The fact that the normal force $F$ is nonzero and compressive is due to the Poynting effect [59]. Without $F$, the cylinder would elongate in the axial direction.

Figure 4.21: Moment $M$ and normal force $F$ of the cylinder under torsion for varying anisotropy parameters $r$. The $H : D$ ratio is fixed at 1 : 1.

Based on the moment $M$ and normal force $F$, we can also define two helpful quantities: torsional stress at the outer rim $\tau = \frac{2M}{\pi R_o^3}$ and normal stress $N = \frac{2F}{\pi R_o^2}$. Rivlin and Saunders [62] found that for rubbers, $M$ was proportional to $D$ and $|N|$
Figure 4.22: Moment $M$ and normal force $F$ of the cylinder under torsion for varying height-to-diameter ($H : D$) ratios. The anisotropy parameter is fixed at $r = 3$.

was proportional to $D^2$. So in Figure 4.23, we plot $\left| \frac{N}{\tau} \right|$ and indeed see that it is linear with $DR_0$ for rubber (anisotropy parameter $r = 1$). For the nematic elastomers ($r > 1$), the behavior of $\left| \frac{N}{\tau} \right|$ is nonlinear. There are two distinct regimes of linear behavior with different slopes. The first, higher-slope regime corresponds to the reorientation of the liquid crystals, and the second, lower-slope regime corresponds to the response of the polymer chains.

For both the nematic elastomers ($r > 1$) and ordinary rubber ($r = 1$), the cylinders develop a kinking instability at certain critical values of twist, which corresponds to the abrupt change from positive to negative slope in the $|N/\tau|$ plots. The onset of kinking is a function of the $H : D$ ratios: the larger $H : D$ is, the easier it is for the structure to form a kink. Figs. 4.24a and 4.24b show views of the kink for anisotropy parameter $r = 3$ and height-to-diameter ratio $H : D = 3 : 1$. Note that the cross-section develops a pinch and no longer remains circular.

We also plot the evolution of the internal variables, $\lambda$ and $\delta$, throughout the deformation. Figures 4.26 and 4.27 show $\lambda$ and $\delta$ for varying nodes of radii $r_1$ through $r_5$ in the cross-section halfway through the height of the cylinder (see Figure 4.25).

Due to the Poynting effect, twisting the incompressible cylinder will produce a lengthening in the axial direction, but since the cylinder is constrained in the length, the cylinder instead continuously develops an instability. Similar instabilities were observed experimentally in rubber cylinders that were subjected to twist, with and without axial stretch [29, 70]. Coyne [20] and Thompson and Champneys [70]
Figure 4.23: Plot of the absolute value of the ratio between torsional stress at the outer rim $\tau$ and normal stress $N$ for varying anisotropy parameters $r$ and height-to-diameter ratios $H : D$.

Figure 4.24: Kinking instability for $r = 3$, $H : D = 3 : 1$: (a) Cross-sectional view, halfway through the height of the cylinder. (b) Side view.

Figure 4.25: Nodes with increasing radii $r_1$ through $r_5$ in the cross-section halfway through the height of the cylinder.
Figure 4.26: Evolution of the internal variables, depicted in the triangular region $\mathcal{T}$ for nodes of varying radii.

Figure 4.27: Evolution of the internal variables as a function of twist for nodes of varying radii.
describe the development of a quasi-static localization or helix similar to what we observe in Figure 4.24b. The kink has directionality based on the direction the cylinder is twisted; the helical shape develops as a continuation of the twist direction.

It is not surprising that the nematic elastomer cylinders kink, as the ordinary rubbers do so as well. If the height-to-diameter ratio were longer, we would expect that the cylinders would form loops and upon further loading, the loops would twist, similar to the experiments of [70]. However, the behavior of the normal stress and torsional stress from Figure 4.23 is a new finding.

To understand the instability further, we conducted an eigenvalue analysis in ABAQUS of the cylinder under torsion at various height-to-diameter ratios and anisotropy parameters. The cylinder is first preloaded under the torsion deformation described above, then a buckling step is conducted to gain the eigenvalues and eigenmodes. The viscosity is fixed at $\beta = 0$ throughout the simulation. The first five modes for $r = 3$ and $H : D = 2 : 1$ are shown in Figure 4.28. We observe shear banding as the deformation mode, which is very similar across the five modes. The corresponding eigenvalues, from Mode 1 to 5, are (4.5688, 4.5689, 4.5711, 4.5715, 4.5746). The eigenvalues are very close together, which speaks to an imperfection-sensitive structure [24].

Figure 4.28: First five eigenmodes for $r = 3$ and $H : D = 2 : 1$. The colors represent the magnitude of displacement, from zero displacement (blue) to high displacement (red).
The eigenvalues from the first mode are plotted as a function of anisotropy parameter in Figure 4.29, where the eigenvalues for $r > 1$ start to move away from the isotropic ($r = 1$) eigenvalue.

![Graph](image)

Figure 4.29: Eigenvalues from the first eigenmode plotted against anisotropy parameter for fixed $H : D = 1 : 1$.

### 4.6 Conclusion

In this chapter, we have formulated a constitutive relation to describe non-ideal isotropic-genesis polydomain nematic elastomers, which builds upon the work of DeSimone and Dolzmann [23]. We introduced internal variables that evolve according to a dissipative kinetic process that represent the material behavior throughout the classic regions of interest: a liquid-like region, a region in which fine-scale microstructure develops, and a solid-like region. We verify the model in MATLAB, performing the constrained optimization within the triangular region $\mathcal{T}$ using the \texttt{fmincon} function, then validate it with comparison against experimental results for various biaxial tension tests performed by Kenji Urayama’s group. Finally, we adapt the constitutive relation for input as a user-defined material code in ABAQUS and study the deformation of torsion in a solid cylindrical body. We now have a tool to study these non-ideal polydomain nematic elastomers under arbitrarily complex loading configurations and boundary conditions in the future.
Chapter 5

EXPERIMENTAL CHARACTERIZATION OF NEMATIC ELASTOMERS

5.1 Introduction

The previous chapters of this thesis have addressed the theoretical and computational characterization of nematic elastomers, discussing various deformations of both monodomains and polydomains. The goal of this chapter is to characterize the rate dependence and temperature dependence of these nematic elastomers experimentally.

The synthesis techniques of temperature-responsive nematic elastomers began with thin films. There has been much work in developing ways to align the nematic director field within the plane of the thin film of nematic elastomer, e.g. through applied magnetic field, mechanical strain, and textured surface patterning [12, 13, 45, 74–76, 81–83]. The advent of “click chemistry” allowed bulk nematic elastomers to be made partially crosslinked in molds, after which another crosslinking process could induce nematic alignment, without the limitations of thin films [55, 64, 80]. Further, the capability to 3D print of nematic elastomers with shear-alignment of the liquid crystal mesogens have been developed [4, 40].

In this thesis, we make samples of nematic elastomers in-house using the click-chemistry method of synthesizing nematic elastomers following [64], discussed in Section 5.2. This technique involves an easy-to-follow, one-pot chemistry recipe in which the polymer mixture is poured into open-faced molds, and after curing, polydomain nematic elastomers are formed. If a UV crosslinker is added to the mixture, then there exists the option to undergo a second crosslinking step to create a monodomain, using applied mechanical strain as the alignment technique.

In Chapter 5.3, we discuss the design and build of a thermo-mechanical tensile test setup capable of testing nematic elastomers at various temperatures while capturing strain, stress, temperature, and imaging data. Figure 5.1a shows the SolidWorks design of the experimental setup, and Figure 5.1b shows a picture of the actual setup in the lab. We have implemented an imaging setup using polarized light microscopy. Due to the underlying optical properties of the liquid crystals, useful information can be gained from the optical properties of nematic elastomers. Viewing a sample
of nematic elastomer between cross-polarizers can help determine the amount of ordering existing in the system due to the material’s birefringence. Macroscopically, a monodomain sample in its nematic state appears transparent, and a polydomain sample appears cloudy and opaque. Using a microscope can help study a sample more closely and reveal information about anisotropy and domains.

![SolidWorks model](a)

![Picture](b)

Figure 5.1: (a) SolidWorks model of a portion of the experimental setup. (b) Picture of the experimental setup.

Finally, Section 5.4 shows the stress-strain curves resulting from tests at varying temperatures, and the code used to run the experiments can be found in the Appendix D.2.

5.2 Sample preparation

Materials

For sample preparation, we synthesize main-chain polydomain nematic elastomers, following [64]. We include the photoinitiator HHMP to allow for a second crosslinking, used for director alignment into a polydomain. We use the following chemicals for the synthesis procedure:
Table 5.1: Table of chemicals

<table>
<thead>
<tr>
<th>Chemical name and purpose</th>
<th>Full chemical formula</th>
<th>Manufacturer</th>
</tr>
</thead>
<tbody>
<tr>
<td>RM257, di-acrylate mesogen</td>
<td>1,4-Bis-[4-(3-acyloyloxypropyloxy) benzoyloxy]-2-methylbenzene</td>
<td>Wilshire Technologies</td>
</tr>
<tr>
<td>EDDET, di-thiol spacer</td>
<td>2,2’-(ethylenedioxy) diethanethiol</td>
<td>Sigma Aldrich</td>
</tr>
<tr>
<td>PETMP, tetra-thiol crosslinker</td>
<td>Pentaerythritol tetrakis (3-mercaptopropionate)</td>
<td>Sigma Aldrich</td>
</tr>
<tr>
<td>DPA, catalyst</td>
<td>Dipropylamine</td>
<td>Sigma Aldrich</td>
</tr>
<tr>
<td>HHMP, photoinitiator</td>
<td>2-Hydroxy-4’-(2-hydroxyethoxy)-2-methylpropiophenone</td>
<td>Sigma Aldrich</td>
</tr>
<tr>
<td>Toluene, solvent</td>
<td>Toluene</td>
<td>Sigma Aldrich</td>
</tr>
</tbody>
</table>

Please see Section C.1 for further details. The following lab equipment was used in preparing the samples:

- Fume hood
- Vortex mixer (VWR 10153-834)
- Vacuum oven (VWR 89508-428) and vacuum pump (VWR 89209-762)
- HDPE molds (McMaster 8619K614, then machined by CNC)
- Hot plate (VWR 12620-978)
- Eppendorf micro-pipettes (Sigma Aldrich Z683884)
- Precision scale (VWR 75802-858)
- UV lamp (CureUV 191340), protective box, and UV-protecting glasses (ThorLabs LG3)
- Compartmented sample boxes (McMaster 4629T15)

The molds that were used to make the samples were made from 6” × 6” × 1” HDPE blocks, and we CNCed them into custom shapes in the machine shop.
We made various molds for making polydomain dogbone samples, as well as larger polydomain samples that could be clamped and stretched into monodomains. The corners of pockets are filleted, rather than manufactured with a sharp corner. The sharp corners often lead to cracks in the finished sample, since removing the cured samples is a delicate process, and the polymer can easily get caught on snags and the sample may rip.

**Synthesis procedure**

The sample preparation has been modified slightly from the original source in [64]. Note that there is a template that helps to follow the procedure, found in Section C.2. Below are instructions for following the synthesis procedure.

Record your name, date, and the time you start the synthesis procedure. Make sure you have on the following personal protective equipment (PPE): lab coat, safety glasses, gloves, closed-toe shoes, and long hair tied back. Clean the mold that you will use with isopropyl alcohol, rinse it with deionized water, and wipe it down with a Kim wipe so that it is completely dry. Prepare two glass vials of 30mL capacity, one labeled as “LCE solution,” and the other labeled as “DPA+toluene,” which will hold the diluted catalyst solution.

Set the hot plate dial level to between 7 and 7.5. Place a beaker with water on top of the hot plate, and place the glass thermometer (held by the stand) inside the beaker of water. Allow the water to heat to $\approx 80^\circ$ C.
We will follow a triple batch recipe because making a larger batch attains better accuracy than the single batch recipe from [64]. This is also the recipe for 50 mol% PETMP, but the crosslink density can be changed by modifying the amount of PETMP and EDDET accordingly. Where possible, fill out the template with the actual weight before and after transferring the chemical to the vial, and calculate the percent error between the actual weight and the expected weight.

Start by taring a weigh boat, measuring 12g of RM257 in the weigh boat, and transfer about half of the RM257 to the “LCE solution” vial. Set the rest of the RM257 aside. Add in 3000μL of toluene to the vial via micro-pipette. Place the vial on the hot plate, starting a timer. Occasionally swirl around the solution in the vial by hand.

While waiting for the solution to completely dissolve, prepare the HHMP. Fold a square piece of weigh paper in half across the diagonal, and weigh out 0.0257g of HHMP. Add the HHMP to the vial, and continue heating it on the hot plate. (Since you want to keep the scale tared to the RM257 weigh boat measurement, you’ll need to do the HHMP math by hand. Record the weight of the weigh paper, then the weight of the weigh paper plus HHMP, and finally the weight of the leftover HHMP on the weigh paper.) When the RM257+toluene mixture has dissolved enough that there is enough room in the vial to add more RM257 and toluene, finish adding the rest of the RM257 and 2549.1μL of toluene via micro-pipette. Record the weight of any leftover RM257 on the weigh boat. Place the vial back on the hot plate.

While waiting for the solution to fully dissolve again, prepare the catalyst solution in the separate vial. Make 1.5 times what the solution needs, measuring out 0.0549g of DPA (74.25μL via micro-pipette) and 2.5011g of toluene (2891.25μL via micro-pipette). Mix the catalyst solution on the Vortex mixer thoroughly.

When the LCE solution has fully dissolved and there are no solids left, record the time that it took to heat the solution. Without waiting, move forward with adding 2.1657g of PETMP (1691.64μL via micro-pipette). The PETMP is a very viscous liquid so if you choose to use the micro-pipette, you will need to wait for the solution to be fully drawn into the micro-pipette tip. Add 1.6158g of EDDET (1442.67μL via micro-pipette). Vortex mix the catalyst solution one last time and then add 1.7043g of the solution (1974μL via micro-pipette) to the LCE solution. Vortex mix the LCE solution for ≈ 20 seconds. Slightly loosen the cap on the vial and place the

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I thank Ruobing Bai for this suggestion.
vial in the vacuum oven and pull a vacuum of 20in Hg at room temperature. Keep a constant vacuum of 20in Hg for 45 seconds, then release the vacuum.

Look at the solution and see if there are any solids that have precipitated in the solution. Note the solution appearance: cloudy, clear, undissolved solids? Then if there are undissolved solids, put it back on the hot plate and record the time it takes to get a fully dissolved solution again. Take a picture of the solution and then immediately pour the solution into the mold. An example picture is shown in Figure 5.3a. Make sure the solutions are filled completely without any air bubbles and that each pocket is filled to approximately the same depth. Take a picture of the mold.

Allow the mold to rest in the fume hood at room temperature for about 12 hours before moving onto the next step, which is placing the samples into the vacuum oven.

Cleanup: Record the time that the sample preparation finished, dispose of any hazardous waste appropriately, clean utensils and dishes, turn off the scale, and turn off the hot plate.

**Placing samples into vacuum oven.** Roughly 12 hours later, put the samples into the vacuum oven. Record the date and time. Wearing gloves, place the mold into the center of the middle shelf of the vacuum oven. Pull the vacuum level to 20.5 in Hg by setting the handle to “evacuate,” turning on the vacuum pump, and when the vacuum level reads 20.5 then simultaneously turn off the vacuum pump with one hand and setting the handle to “closed” with the other hand. (We overshoot the vacuum level because the vacuum level will settle to $\approx 20$in Hg after a few hours.) Turn on the temperature switch, with the temperature dial set to 1.5. Take note of anything irregular. Record the temperature at a couple times throughout the 24 hour period.

**Taking samples out of vacuum oven.** Roughly 24 hours later, take the samples out of the vacuum oven. Record the date. Turn off the temperature switch on the vacuum oven, and release the vacuum by setting the handle to read “vent.” Take the samples out of the vacuum oven using a heat-protecting glove, and place them in the fume hood. Write down the time that the samples are taken out of the oven. Note anything irregular.
For sample storage, be sure to keep the samples stored inside a sealed box in a cool, dry cabinet, protected from ambient light and air. Whenever samples are not being prepared or used, store them in this way. You can also vacuum seal them.

About 30 minutes later, write down the time. Wear gloves and use a metal spatula to gently take each sample out of the mold. Heat each sample until it is in its isotropic state, then place it in a labeled sample box. Note any cracks, bubbles, or other irregularities in the samples.

**UV crosslinking.** Anytime after the samples have come out of the molds, you can UV crosslink them at room temperature (as long as the samples were made with HHMP, the photoinitiator). Record the date and time that you are UV crosslinking the samples, as well as the UV crosslinking equipment. You must wear UV-protecting glasses at all times during this process. Wearing gloves, clamp the sample in the desired configuration and place it inside the UV crosslinking box. Turn on the UV lamp and use a stopwatch to be able to record the time spent under the UV light. If necessary, flip sample over and repeat process until the sample is fully crosslinked in the desired configuration. Take pictures, and take note of the temperature of the box (the UV lamp may cause the box to heat up) and any other irregularities.

For instance, Figure 5.3b shows a polydomain sample that was clamped until it became optically clear, which forms a monodomain when crosslinked in that configuration.

![polydomain sample](image1)

Figure 5.3: (a) LCE solution immediately before being poured into mold. (b) Polydomain sample clamped in a stretched monodomain state, ready for UV crosslinking.
Best practices

Below are a list of items to consider when making samples:

- When solution is inside the vial, keep the lid on the vial as much as possible because liquid can evaporate and disturb any weight measurements.
- When weighing on the scale, close all doors to the scale to limit air flow that will disturb the measurements, and wait for the numbers to settle before recording the measurement.
- Never exceed the maximum scale weight, or 60g.
- Always return micro-pipettes to their maximum stated capacity for storage. (e.g. If it is a 1000μL micro-pipette, turn the reading to 1000μL.)
- Always use a new tip when using micro-pipettes. Keep the tip of the micro-pipette submerged in the liquid you are retrieving at all times. Wipe off any excess liquid from the micro-pipette tips before transferring the solution to the vial. Ensure that there are no bubbles inside the micro-pipette tip.
- The synthesis should take place roughly 12 hours before the samples go into the vacuum oven (if necessary, more than 12 hours can pass, but not less). Then another 24 hours should pass before the samples are taken out of the vacuum oven (if necessary, slightly more than 24 hours can pass, but not less). Then you can carefully take the samples out of the molds about 30 minutes after they have been taken out of the vacuum oven. You can UV crosslink anytime after they have been taken out of the molds, but testing of these samples should ideally take place within 2 days of being taken out of the molds.
5.3 Experimental setup
Design, manufacturing, and assembly

Figure 5.4: Picture of experimental setup.

Figure 5.4 shows the entire experimental system with labels, and the schematics in Figure 5.5 and 5.6 depict the connections in the test setup. The system is made up of the following subsystems: chamber assembly (chamber with windows, stationary bottom clamp, and moving pullrod with clamp), heating (heaters, RTD sensors, and temperature controller), extension (linear stage, and linear stage controller, and suspension assembly), load (load cell and external power supply), optics (optical table, lighting, microscope, camera, cross-polarizers), and data acquisition (computer and DAQ).

The setup was designed in SolidWorks (see Figure 5.1a), with metal adapters to connect the chamber to the optical table, the pullrod to the load cell, and the load cell to the linear stage. Glass windows were designed in the front and the back of the chamber to accommodate the optical setup. The pullrod and one of the adapters was originally manufactured in metal, but conversations with Sam Daly led to them being remade in machinable ceramic to prevent the load cell from overheating. We used the Jim Hall Design and Prototyping Lab (the machine shop in the Mechanical and Civil Engineering department of Caltech) to manufacture all of the metal and machinable ceramic pieces.
The data acquisition system involved soldering the wires of one of the RTDs and the load cell to the DAQ, and coding in MATLAB to obtain the temperature and load data. It also required further coding in MATLAB to acquire the extension data from the linear stage. Julia Combs, a student who worked in the lab for a summer via the Summer Undergraduate Research Fellowships (SURF) program, figured out how to obtain imaging data from the camera in MATLAB.

Clamping the sample can be difficult, depending on the stiffness and texture of the sample. We chose to use binder clips to clamp the samples because they are self-tightening. Binder clips of different sizes and materials were used depending on the sample. Clamping at high temperature is especially challenging because the
samples became susceptible to breakage at the clamping point. To avoid the sample slipping in the grips, we handle the test samples only with gloves, so that the sample does not retain any oils from skin. Additionally, sanding the insides of the binder clip can help to create friction between the sample and the clamp, or adding a piece of rubber (e.g. from a rubber glove) can also create more friction so that the samples do not slip out.

**Materials**

The various components of the setup, listed by subsystem, are:

- **Custom-made chamber assembly**
  - Chamber components (McMaster 1658A12, 8983K128, 8983K118, 2313N23)
  - Various hardware (nuts, bolts, washers, etc.)
  - Various adapters (McMaster 8489K44, 4416T47)
  - Windows and sealing (McMaster 8476K999, 92320A662)
  - Stationary bottom clamp and moving clamp on pullrod (McMaster 12755T82)
  - Moving pullrod (McMaster 8489K46)

- **Heating**
  - Two heaters (Omega OTF-102/120V)
  - Two RTD air temperature sensors (Omega RTD-805-B)
  - Temperature controller (Omega CSI32RTD-C24)

- **Extension**
  - Linear stage (Physik Instrumente M-531.EC)
  - Linear stage controller (Physik Instrumente C-863.11)
  - Linear stage z-axis mounting bracket (Physik Instrumente M-592.10)
  - Suspension system (ThorLabs VB01B)

- **Load**
  - Load cell (Omega LC101-50)
  - External 10V power supply

- **Optics**
  - Vibration-isolation table (ThorLabs T46J)
  - LED backlight (Edmund Optics 83-873)
  - Long working-distance microscope (Infinity K2/SC)
  - Camera (Edmund Optics 86-770)
  - Cross-polarizers (Infinity 991167, Edmund Optics 45-669)
  - Calibration slide (ThorLabs R1L3S5P)

- **Data acquisition**
Lab computer (Lenovo ThinkStation P330)
DAQ (Omega INET-600)
MATLAB software

**Tensile test procedure**

Below are instructions for the tensile test procedure:

Start by donning the proper PPE: put on safety glasses, gloves when handling samples, and heat-protection gloves when you will be in contact with the chamber when it is hot. On the lab computer, make a new file folder that will hold the test results (naming it with the date of the test, sample information, etc.) and also start a new file in OneNote using the test results template found in Section C.3. Copy in the MATLAB code to the folder (see the Appendix in Section D.2).

Visually check that the linear stage has no obstructions in its path, and make sure that anywhere the load cell moves will not crush it or cause damage to it. A suggestion is to put up a sign on the door to the lab saying that a test is in progress, and make sure that no wires or other objects will touch the optical table throughout the test.

Refer to the schematic in Figure 5.5 for all relevant connections. Turn on the external 10V power supply, linear stage controller, and temperature controller. Plug in the 3 USB connectors to the computer: the linear stage controller, the iNet600 data acquisition (DAQ), and the camera. For optical data, turn on the LED backlight and position the cross-polarizers. There is an in-line cross-polarizer inside the microscope, and another cross-polarizer should be between the sample and the LED backlight, as seen in Figure 5.6. Then open MATLAB, set the working directory to the current test’s folder, and open the main .m file (see Section D.2).

Wearing gloves, heat the sample you wish to test and let it cool completely. Then cut the sample into a strip using a new, sharp razor blade. With calipers, measure the width and thickness of the sample. Record these measurements in the OneNote template and in the MATLAB code, taking care to keep the units consistent.

If collecting imaging data, open the uEye Cockpit software. Make sure that the camera is facing in the correct orientation (see the red sticker on the camera). Take and save a picture of the ThorLabs calibration slide so that you can create a size scale. Ensure that polarizing filters are in place. You can set up a sequence of images to be taken, or you can capture a video. Be sure to take a picture that allows you to correlate the image to the timing of the test.
If the test is performed at a temperature higher than room temperature, then use the Omega temperature controller to set the setpoint temperature according to the instructions manual (see Figure 5.7). Note that the temperature is in Farenheit, not Celsius. Preheat the test setup to the desired temperature and then clamp the samples in to begin the test, using the preheat section of the code to record the preheat data. Put the insulation on the top of the chamber, to keep the load cell cool. Close the chamber door, and point the floor fan (on a low speed) at the load cell to keep it cool as well.

In the code, enter in the desired strain and strain rate, according to Table 5.2. Also set the position where the test will start using `startCoarsePos`. Ensure that this position will never cause the pullrod to crash into anything, which would damage the linear stage and the load cell.

<table>
<thead>
<tr>
<th>Strain Rate</th>
<th>Strain Rate Rate</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fast strain rate</td>
<td>$10^{-2}$/sec</td>
</tr>
<tr>
<td>Medium strain rate</td>
<td>$10^{-3}$/sec</td>
</tr>
<tr>
<td>Slow strain rate</td>
<td>$10^{-4}$/sec</td>
</tr>
</tbody>
</table>

Double-check that the linear stage is free to move without obstruction. Run the block of code that connects `MATLAB` to the linear stage controller and references the linear stage. This code will cause the linear stage to move to its reference position at 153mm, which is in the exact middle of the 306mm-long stage.

I thank Sam Daly for this tip about using a fan to cool the load cell.
Attach the pullrod to the load cell. With gloves on, carefully clamp the sample into the top clamp. You can finish clamping the sample using one of two methods. For the first method, if you know that the start position in \texttt{startCoarsePos} is a good location to hold the specimen in the clamps, then hold the bottom clamp open and execute the next block of code, which moves the linear stage to the start position. Skip the next block of code, which allows you to jog the linear stage up and down, without executing it. Or you may do the second method, which is to execute the block of code that moves the pullrod to \texttt{startCoarsePos}, and you can execute the block of code after that to jog the linear stage up and down to get the specimen clamped in position well. Type \texttt{u} or \texttt{d} to move the linear stage up or down in small increments until you are satisfied with the position, and you can clamp the sample in this way. Do not introduce any stresses to the specimen, and make sure the specimen is aligned well. When the specimen is fully clamped, close the chamber door.

The next block of code queries the linear stage for its position and then calculates the gauge length automatically. There is no action needed here.

The next block of code calculates the extension of the linear stage, the velocity, and the time that the test will take. This code assumes a full cycle through loading and unloading. The code also calculates the cross-sectional area. Modify the step size accordingly for various strain rates, ensuring that the linear stage will not move too fast or too slow for its specifications.

After that, the next block of code will configure the linear stage, calculating the starting position, the stopping position, and the array called \texttt{steps}. Also execute the initialization of the data acquisition system, using the correct .prf file that matches the correct load cell and temperature sensors signals. There is no action required from you here.

Finally, the \texttt{move} function is called in the next block of code, and the position, temperature, and load data are saved in their corresponding structures. Be sure not to disturb the test in any way by touching the optical table. After the test is finished, the last blocks of code can be executed to plot and save the data, which includes the stress-strain curve.

Save the files and data, complete the test template as you go, and remove the specimen from the clamps. Create a new folder for each test performed, and repeat this testing process as needed. After you complete testing, then there are cleanup procedures to follow. Save all data, exit MATLAB, and unplug the 3 USB connectors from the
computer (linear stage controller, camera, and data acquisition). Turn the power off to the linear stage, the temperature controller, and the external power supply. Cover up the camera lens and store it so that no light or dust enters the sensor. Take the pullrod off the load cell for storage so that the load cell is unloaded when not in use. Be sure to store the specimens properly (in containers, inside closed cabinets) so that they do not age as quickly in the ambient air or light.

**Best practices**

Below is a list of items to consider when using the tensile test setup:

- Don’t touch the vibration isolation table while test is running!
- It might be helpful to put a sign on the lab door saying "testing in progress" so people don’t come into the lab and disrupt the test.
- Wear safety glasses (in case of fracture).
- Make sure that when the linear stage moves, the wires and everything are not going to get caught, that the load cell won’t be jammed or overloaded.
- Shut the door of the chamber during the test, because we don’t want ambient air in the lab to affect it.
- Handle samples with gloves so that your hand’s oils don’t transfer to the sample, making it harder for the clamp to hold onto the sample.
- Use a new sharp razor blade to cut samples
- Store any samples inside the box in darkness when not being tested.
- Test samples starting on the day that the samples came out of the oven, and finish tests within 2 days.
- Use virgin samples for all the tests, unless stated otherwise. Make a note of the sample’s loading and heating history.
- I am able to leave the experiment unattended when testing is done at room temperature, with the permission from the Safety Office, who has inspected the experimental setup. However, I do not leave the experiment unattended when testing is done at high temperature because the heaters present a fire risk.
- When testing a sample in the load/unload cycle, ensure that you choose a strain that will be safe enough for the sample not to fail prematurely during the test.
**Test matrix**

Here we have the completed test matrix, with each cell denoting the date that the samples are synthesized (not tested). The “polydomain" samples are not treated to a second crosslinking and are therefore in the polydomain state at nematic temperatures, while the “monodomain $\perp$" samples are pulled in a uniform direction and treated to a second crosslinking so that they are in the monodomain state at nematic temperatures. The monodomain samples are pulled in tension perpendicular to the direction of the mesogen orientation.

**Table 5.3: Tests at nematic temperature (room temperature $\approx 22^\circ$C)**

<table>
<thead>
<tr>
<th></th>
<th>Fast strain rate</th>
<th>Medium strain rate</th>
<th>Slow strain rate</th>
</tr>
</thead>
<tbody>
<tr>
<td>Polydomain pull</td>
<td>7/29/20</td>
<td>8/12/20</td>
<td>11/4/20</td>
</tr>
<tr>
<td>till failure</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Polydomain load/unload sample 1</td>
<td>7/29/20</td>
<td>7/29/20</td>
<td>7/29/20</td>
</tr>
<tr>
<td>Polydomain load/unload sample 2</td>
<td>7/29/20</td>
<td>7/29/20</td>
<td>7/29/20</td>
</tr>
<tr>
<td>Polydomain load/unload sample 3</td>
<td>7/29/20</td>
<td>7/29/20</td>
<td>7/29/20</td>
</tr>
<tr>
<td>Monodomain $\perp$ pull till failure</td>
<td>8/12/20</td>
<td>8/12/20</td>
<td>8/12/20</td>
</tr>
<tr>
<td>load/unload sample 1</td>
<td>8/12/20</td>
<td>8/12/20</td>
<td>8/12/20</td>
</tr>
<tr>
<td>Monodomain $\perp$ load/unload sample 2</td>
<td>8/12/20</td>
<td>8/12/20</td>
<td>8/12/20</td>
</tr>
<tr>
<td>Monodomain $\perp$ load/unload sample 3</td>
<td>8/12/20</td>
<td>–</td>
<td>–</td>
</tr>
</tbody>
</table>

**Table 5.4: Tests at isotropic temperature ($\approx 130^\circ$C)**

<table>
<thead>
<tr>
<th></th>
<th>Fast strain rate</th>
<th>Medium strain rate</th>
<th>Slow strain rate</th>
</tr>
</thead>
</table>
5.4 Experimental results

In this section, all samples have 50 mol% PETMP. Each curve corresponds to distinct samples (no samples were reused). The stress is defined as force divided by the undeformed cross-sectional area.

Results for polydomain samples at room temperature

Figure 5.8a shows the results for polydomain samples pulled till failure at three different strain rates, all at room temperature. The fastest strain rate corresponds to the stiffest response, as expected. Figure 5.8b plots 3 samples tested at the fastest strain rate with their load and unload curves. Figure 5.9a shows the same for the medium strain rate, and Figure 5.9b shows the same for the slowest strain rate.

![Stress-strain curves for polydomain samples](image1)

Figure 5.8: (a) Stress-strain curve of polydomain samples that were pulled till break. (b) Load and unload stress-strain curves for polydomain samples at fast strain rate.

![Stress-strain curves for polydomain samples](image2)

Figure 5.9: (a) Load and unload stress-strain curves for polydomain samples at medium strain rate. (b) Load and unload stress-strain curves for polydomain samples at slow strain rate.
Figure 5.13 shows the stress-strain curves for all the polydomain samples at all strain rates.

![Stress-strain curves for all polydomain samples](image)

Figure 5.10: Stress-strain loading curves for all room-temperature polydomain samples at all strain rates.

**Results for monodomain samples at room temperature**

Figure 5.11a shows the results from monodomain samples that were pulled perpendicular to their crosslinked nematic pattern until failure (all at room temperature). Figures 5.11b, 5.12a, and 5.12b show the load and unload curves for samples that were pulled at the fastest, medium, and slowest strain rates, respectively.

![Stress-strain curve of monodomain samples](image)

(a) Stress-strain curve of monodomain samples that were pulled til break.

![Load and unload stress-strain curves](image)

(b) Load and unload stress-strain curves for monodomain samples at fast strain rate.
Figure 5.12: (a) Load and unload stress-strain curves for monodomain samples at medium strain rate. (b) Load and unload stress-strain curves for monodomain samples at slow strain rate.

Figure 5.13 show the stress-strain response for all the monodomain samples at all strain rates at room temperature.

Figure 5.13: Stress-strain loading curves for all monodomain samples at all strain rates.

Figure 5.14 shows the image of stripe domains that formed in the uniaxial stretch of a monodomain sample. The stretch direction is vertical, and the stripes are parallel to the direction of stretch. The width of a stripe is about 75µm.
Figure 5.14: Image of stripe domains that appeared in the uniaxial stretch of a monodomain sample pulled perpendicular to its nematic alignment direction (tested at room temperature, imaged with cross-polarizers).

**Results for polydomain samples at high temperature**

In this section, polydomain samples were tested at high temperature, \( \approx 130 \text{ degC} \). The entire testing chamber was preheated to the high temperature and then the samples were clamped in to begin the test. Figure 5.15a shows the stress-strain curves of the polydomain samples at all strain rates at high temperature, and Figure 5.15b shows the stress-strain curves of the polydomain samples at room temperature and high temperature for comparison.

![Stress-strain loading curves](image)

**Figure 5.15:** (a) Stress-strain loading curves for samples at high temperature at varying strain rates. (b) Stress-strain loading curves for all samples at low and high temperatures and varying strain rates. Note that the high-temperature curves appear very close to the origin.
The samples broke in the clamps extremely quickly at high temperature, so there was not much stress-strain data gathered in these tests. Strain rate does not have the expected effect upon the stress-strain curves in Figure 5.15a; the fastest strain rate corresponds to the softest sample, and the slowest strain rate corresponds to the stiffest sample. However, when analyzing the effect of temperature in Figure 5.15b, the three samples at high temperature are all stiffer than the three samples at low temperature, which is the expected effect.

**Cyclic testing results**

This test was conducted by Julia Combs. She tested a polydomain sample with 15 mol% PETMP at a strain rate of $10^{-2}$/sec at room temperature. She used one sample throughout the twenty cycles. For a given cycle, she clamped it, loaded it, unloaded it, heated it up in the clamp (without unclamping it) using the heat gun, waited for it to cool ($\approx$ 3 min), then repeated. The results can be seen in Figure 5.16. The stress at 200% strain decreases with the number of cycles, while the stress at which mesogen reorientation occurs increases with the number of cycles. We suspect that some cycles have a higher stress plateau than others because the sample wasn’t fully cooled down to room temperature, so the sample would be stiffer due to being in a more isotropic state.

![Cyclic Loading of Nematic Elastomers](image)

Figure 5.16: Cyclic stress-strain curves for a single sample over 20 cycles.

Repeatedly testing the same sample, or different samples from the same synthesis batch, would provide insight into the aging of the samples over time. Aging in polymers is well understood, but not well-studied in nematic elastomers. Aging
can depend upon the particular chemistry of synthesis, temperature, loading history, radiation, and more, and aging would affect nematic elastomers’ use in various long-term applications.

5.5 Conclusion
In this chapter, we have described the synthesis of nematic elastomer samples that were tested in our experimental tensile test apparatus, which features a temperature-controlled chamber and polarized light microscopy capabilities. We presented the results for the uniaxial stretch of polydomain and monodomain nematic elastomers in their isotropic and nematic temperatures. We observed stripe domains in monodomain samples pulled perpendicular to their nematic alignment, and we have also conducted cyclic loading tests. There exists much potential to build upon the current experimental setup for future research directions. For example, a future avenue of research could be obtaining full strain fields through digital image correlation (DIC). There are also some improvements that can be made to the experimental setup. The chamber design could be improved for easier clamping of samples when the setup is pre-heated, and the clamping mechanism could be improved beyond simple butterfly clamps.

Future experiments that could be conducted include uniaxial stretch of monodomain samples at varying angles with respect to director alignment, in-depth characterization of aging in samples over time, cavitation of a disk (for comparison with results from Chapter 3), and cyclic testing of samples at varying temperatures, strain rates, and director orientation.
Chapter 6

CONCLUSION AND FUTURE OUTLOOK

6.1 Summary and impact of the findings

Liquid crystal elastomers = liquid crystals + elastomer

At the heart of liquid crystal elastomers lies the interplay between two aspects of its microscopic makeup, i.e. the liquid crystal molecules embedded within a lightly crosslinked polymer backbone, and the macroscopic shape change. The microscale affects the macroscale, and vice versa. In response to an external stimulus such as change in temperature, nematic elastomers continuously deform between a high-temperature isotropic state, in which the liquid crystals are randomly oriented and there is no orientational order, and a low-temperature nematic state, in which it is energetically favorable for the rigid, rod-like liquid crystal molecules to align along a preferential direction. In return, external stretch of the material at the macroscale induces a change in the microscale, where the liquid crystal molecules reorient to align along the direction of most stretch. Furthermore, they have both liquid-like behavior (during mesogen reorientation) and solid-like behavior (during the elastic response of the elastomer chains). Finally, this unique microscopic structure leads to the ability for the material to develop fine-scale microstructure, and consequently exhibit their unusually soft behavior. These characteristics continue to inspire researchers in this field since the foundational stripe domains from the uniaxial stretch experiments of Kundler and Finkelmann [42].

Modeling nematic elastomers

This thesis utilized three different approaches to the continuum-level modeling of nematic elastomers. In Chapter 2, we introduced the classical model to characterize nematic elastomers, which is the Bladon-Terentjev-Warner (BTW) model, or “trace formula” [10]. However, because this model is an extension of the neo-Hookean model for rubbers and is therefore derived from Gaussian chain modeling, it works well to capture the material behavior at small to moderate stretches, but it fails to predict the stiffer behavior when the polymer chains are stretched sufficiently far such that Gaussian statistics are violated. For instance, the BTW model cannot capture the correct behavior at large stretch in the case of the expansion of a balloon subjected to internal pressure. Thus, we used the Ogden extension of the BTW model...
theory in Chapter 2, which allowed us to study the large-deformation behavior of a cylindrical nematic elastomer balloon and its application as a soft pump.

In Chapter 3, we introduced DeSimone and Dolzmann’s theoretical model that captures fine-scale microstructure through relaxation of the BTW theory [23]. In the model, nematic elastomers can form domains of region $L$, denoting a liquid-like behavior, region $M$, denoting the formation of fine-scale microstructure, and region $S$, denoting the solid-like behavior where there is no nematic mesogen reorientation. Again, the BTW theory is unable to capture the correct physics of nematic elastomers at large stretch, so we employed a generalized Mooney-Rivlin model, following Silhavy, Agostiniani, and DeSimone [1, 65].

Finally, Chapter 4 discussed the new constitutive relation that we developed to describe non-ideal isotropic-genesis polydomain nematic elastomers. The model featured internal variables that corresponded to the formation of microstructure, which evolved according to a dissipative relation, and viscosity associated with the polymer network.

**Cylindrical balloon and corresponding pump**

In Chapter 2, we analyzed a monodomain cylindrical balloon in which the nematic director was axially oriented in its undeformed state. We found multiple equilibrium solutions. One solution was metastable, where the nematic mesogens would not rotate from their original axial orientation. The other two solutions featured fine-scale microstructure development, where the nematic director rotated symmetrically at a positive or negative angle in opposite directions to accommodate the deformation. The balloon would form alternating domains of the two symmetric solutions, and the macroscopic behavior of the twist would also alternate between the two solutions.

Then we described how such a cylindrical balloon could form a pumping cycle through a combination of inflow, outflow, and changing temperature. We measured the ejection fraction of the pump, which described how much fluid the balloon could pump in a given cycle, and found that it was extremely high due to the softness of the material and thus the nature of the pressure-volume curve.

This work was one of the first examples of actuation from geometries beyond flat, two-dimensional sheets, and analyzed nematic elastomers under a more complex loading and condition than previously considered.
Universal deformations of nematic elastomers

In Chapter 3, we studied the four deformations of cylindrical balloon expansion, spherical balloon expansion, cavitation of a disk, and bending of a block. We discussed how the materials can form regions of $L$, $M$, and $S$ in response to the boundary conditions (e.g. for the spherical and cylindrical balloons, see Figures 3.3, 3.7 respectively). For the balloons, due to the boundary condition of internal pressure, the area of most stress was in the inner radius of the balloon, which was where the balloon formed microstructure first, before eventually the entire cross-section became enveloped by region $M$. We discussed how the deformation of the spherical balloon inflation was essentially the deformation of equibiaxial stretch, so it followed the EB curve of Figure 6.1, while the cylindrical balloon inflation with fixed axial stretch was a manifestation of planar extension, so its deformation path was the PE curve. Consequently, the spherical balloon remained in the microstructure-formation region for the rest of its deformation, whereas the cylindrical balloon stopped microstructure formation at a certain point and had a solid-like response at high stretch.

![Diagram](image)

Figure 6.1: Regions of $L$, $M$, and $S$ in the phase diagram of $(s,t)$. Also shown are common deformations and their paths through the three regions: equibiaxial stretch (EB), unequal biaxial stretch (UB) with various strain ratios ($5/3$, $5/2$, $5/1$), planar extension (PE), and uniaxial stretch (U).
Furthermore, there was a spontaneous deformation associated with the temperature-induced phase transition, in which the surrounding polymer chains stretch parallel to the preferred direction and contract in the transverse directions. This phenomenon, which occurs at zero stress, was observed in both Chapters 2 and 3. In the case of the balloons subjected to internal pressure, for anisotropy parameters greater than 1, the pressure-stretch curves did not begin inflation at the point \((\lambda_0, p) = (1, 0)\). It was also seen clearly in the plot of moment vs. stretch for the deformation of bending. These spontaneous deformations associated with the unique thermo-mechanical coupling in nematic elastomers exemplify why these materials are so compelling to study.

We also discussed the cavitation of a disk, which was originally studied for ordinary rubber by Gent and Lindley [30], and the bending of a block. We compared the results between ordinary rubber response and the nematic elastomer response. Interestingly, in the bending deformation, we found that there were multiple equilibrium solutions as the deformation progressed, similar to the multiple equilibrium solutions found in Chapter 2.

Previous work in the field of nematic elastomers has focused on studying homogeneous deformations, so this chapter is a significant step towards understanding nematic elastomers under various inhomogeneous deformations.

**Computational characterization of non-ideal, isotropic-genesis polydomain nematic elastomers**

In Chapter 4, we developed an entirely new constitutive model to describe non-ideal, isotropic-genesis polydomain nematic elastomers. We implemented the model in the commercial finite-element software ABAQUS by writing a custom UMAT. We verified the ABAQUS code against the one-dimensional theory in MATLAB for various biaxial extension tests (uniaxial extension, planar extension, equibiaxial extension, and unequal biaxial extension), then validated the results from the model against experimental results for the same biaxial extension tests. We found great agreement between the simulation and the experiments, which were conducted by Tokumoto *et al.* [71]. We then studied cylinder torsion at various height-to-diameter ratios and anisotropy parameters. We compared the results between ordinary rubber and nematic elastomer and discussed the kinking instability.

Though nematic elastomers have been studied through multiple decades, there are still barriers in realizing these materials in engineering applications, unlike materials
such as shape memory alloys, which are commercially available and accepted in industry [11, 51]. The finite element model using our constitutive relation from Chapter 4 is a significant step towards a tool that can study nematic elastomers as an engineering material.

**Experimental characterization of rate dependence and temperature dependence in polydomain and monodomain nematic elastomers**

Finally, Chapter 5 described the synthesis of nematic elastomer samples that were tested in our experimental tensile test apparatus, which featured a temperature-controlled chamber. We presented the results for the uniaxial stretch of polydomain and monodomain nematic elastomers tested at their isotropic and nematic temperatures. The thermo-mechanical coupling inherent in nematic elastomers is responsible for the characteristic stress-strain curves presented in this chapter, for example in the polydomain-to-monodomain transition and in the monodomain-pull-perpendicular experiments that yielded the stripe domain patterns. We could clearly identify the distinct regimes in the stress-strain plots where the liquid crystal mesogen reorientation dominates vs. where the elasticity of the polymer backbone dominates.

The experimental setup also featured polarized light microscopy capabilities. Because of nematic elastomers’ optical properties derived from the underlying liquid crystals, we were able to gain valuable information from viewing samples under cross-polarizers. We observed the formation of fine-scale microstructure, exhibited by stripe domains in monodomain samples pulled perpendicular to their nematic alignment. Finally, we also studied viscoelastic effects by studying the hysteresis at various strain rates and conducting a cyclic loading test.

### 6.2 Future outlook

Following the work presented in this thesis, there are many exciting future avenues of research within the field of nematic elastomers, and more broadly, active materials.

**Theoretical characterization**

The natural next steps in characterizing these materials theoretically are to use the relaxed generalized Mooney-Rivlin model to obtain analytical solutions to the remaining families of universal deformations that we did not cover in Chapter 3. Additionally, the cylindrical balloon deformation in Chapter 2 had multiple stable solutions, as did the bending deformation in Chapter 3. There is a lot of potential
for designing structures with multiple stable solutions for energy-efficiency reasons, so that the only energy expended is moving between the various stable states. The snap-through instability of Chapter 2 and the kinking instability of Chapter 4 are both interesting phenomena that result from finite elasticity at large deformation, and it would be interesting to continue studying other such instabilities in nematic elastomers.

**Computational characterization**

To build upon the computational work of Chapter 4, one could perform simulations of the universal deformations in ABAQUS using the constitutive relation for non-ideal polydomains, and investigating the effects of viscoelasticity, for instance plotting the hysteresis between loading and unloading. We also saw that interesting instabilities can arise, for instance in the torsion of a cylinder, opening the door to further investigation regarding the onset of such instabilities in nematic elastomers, the dependence of the instability upon material parameters and geometry, and more. With the UMAT built, further finite element simulations answering such questions are straightforward to run and analyze.

**Experimental characterization**

The natural next step in experimentally characterizing these materials is to expand upon the experimental results of Chapter 5 to build a complete set of material parameters to match an Ogden model for nematic elastomers for the ideal Bladon-Terentjev-Warner model with the non-ideality in Chapter 2, the relaxed generalized Mooney-Rivlin model of Chapter 3, and the constitutive relation for non-ideal polydomains for Chapter 4. Then, one can perform experiments on the expansion of a monodomain nematic elastomer balloon and quantitatively compare the expansion and twist parameters from Section 2.3, as well as manufacturing and testing a pump made from this monodomain balloon to construct pressure-volume curves at different anisotropy parameters, as described in Section 2.4. The physical size of such a pump could be on the order of centimeters, such as the balloon found in [34]. Furthermore, one can perform experiments within various classes of universal deformations, e.g. the bending of a polydomain block, inflation of polydomain balloons (spherical and cylindrical), and cavitation of a polydomain disk, and match the experimental results with the theoretical results of Chapter 3. Other experiments that were traditionally performed on thin films, such as the bulge and blister tests, would also be useful avenues of exploration to characterize the material.
Nematic elastomers and other active materials

Within the field of active materials, there are exciting paths forward leading towards the multifunctional, the adaptable, and the autonomous. We can think of the integration of active materials with origami/kirigami for shape-morphing applications, as well as designing adaptable features such as roughness and stickiness for bio-inspired soft robotics applications.

Thermotropic nematic elastomers are quick to heat, but the cooling time can be slow in ambient air based on the temperature differential and the geometry of the sample. Actuating within a bath increases response times but can be limiting depending on the application. Phototropic nematic elastomers, for instance, have better response times, although there are other issues associated with penetration depth of the light. The combination of such nematic elastomers responding to multiple stimuli, or the combination of various active materials responding to multiple stimuli, can create multifunctional structures in which the order and extent of the responses can be controlled and tuned for the desired actuation. Additionally, composites of nematic elastomers can be designed for one or more desired properties, e.g. fiber-reinforced elastomers for augmented mechanical behavior, stretchable wiring for augmented electrical capabilities. Composites of active materials can be optimized for various loading configurations using topology optimization.

As mentioned previously, we observe three distinct length scales in nematic elastomers: the nematic mesogens (order of nanometers), domains of nematic alignment (order of microns), and the macroscale (on the order of centimeters). However, if a desired application is of a different macroscopic length scale than this, perhaps designing artificial nematic elastomers, featuring a fundamental phase transition occurring at the smallest length scale fully coupled with shape change at the macro-scale, might be a fruitful area of exploration.
BIBLIOGRAPHY


Appendix A

Supplementary Information in Developing the Generalized Mooney-Rivlin Model

A.1 Principal stretches

Recall that the cofactor of a second-order tensor $F$ is $\text{cof} \, F = (\det F) \, F^{-T}$ when $\det F \neq 0$. Let $\lambda_{\text{max}} \geq \lambda_{\text{mid}} \geq \lambda_{\text{min}}$ be the singular values or principal stretches of $F$, and let $\lambda_{\text{max}}^2 \geq \lambda_{\text{mid}}^2 \geq \lambda_{\text{min}}^2$ be the eigenvalues of $FF^T$. Let $s = \lambda_{\text{max}}$ be the largest singular value of $F$, and let $t = \lambda_{\text{max}} \lambda_{\text{mid}}$ be the largest singular value of $\text{cof} \, F$. We can summarize the principal stretches in terms of $s$ and $t$:

$$\lambda_{\text{max}} = s, \quad \lambda_{\text{mid}} = \frac{t}{s}, \quad \lambda_{\text{min}} = \frac{1}{t},$$

where we have used incompressibility for the last relation ($\lambda_{\text{max}} \lambda_{\text{mid}} \lambda_{\text{min}} = 1$).

A.2 Minimization of the energy with respect to the nematic director

Energy based on the first invariant

The energy for nematic elastomers based on $F$ is as follows:

$$\tilde{W}_1 = \min_{n \, \text{s.t.} \, |n| = 1} \left\{ c \left| \text{tr} \left( \ell_n^{-1} b \right) - 3 \right|^{\frac{1}{p}} \right\}. \quad (A.2)$$

Because $I_1$ is an invariant, its value is the same no matter what basis is chosen. Therefore, let us choose the principal basis:

$$b = \begin{pmatrix} \lambda_{\text{max}}^2 \\ \lambda_{\text{mid}}^2 \\ \lambda_{\text{min}}^2 \end{pmatrix}. \quad (A.3)$$

Minimizing the energy over the director $n$ yields the result that $n$ is parallel to the eigenvector corresponding to the largest eigenvalue, $\lambda_{\text{max}}^2$. That is, $n = (1, 0, 0)^T$. Thus, the step-length tensor should be as follows:

$$\ell_n^{-1} = r^{1/3} \left( I + \left( \frac{1}{r} - 1 \right) n \otimes n \right)$$

$$= \begin{pmatrix} r^{-2/3} \\ r^{-2/3} \\ r^{-2/3} \end{pmatrix}. \quad (A.4)$$

$$= \begin{pmatrix} r^{1/3} \\ r^{1/3} \\ r^{1/3} \end{pmatrix}. \quad (A.5)$$
Therefore, we have

\[
\hat{W}_1(s, t) = c \left| \text{tr} \left( \ell_n^{-1} b \right) - 3 \right|^p
\]

\[
= c \left| r^{1/3} \left( \frac{\lambda_{\text{max}}^2}{r} + \lambda_{\text{mid}}^2 + \lambda_{\text{min}}^2 \right) - 3 \right|^p
\]

\[
= c \left| r^{1/3} \left( \frac{s^2}{r} + \frac{t^2}{s^2} + \frac{1}{t^2} \right) - 3 \right|^p, \tag{A.7}
\]

where in the last equality we have used the fact that \( s = \lambda_{\text{max}} \) and \( t = \lambda_{\text{max}} \lambda_{\text{mid}} \).

### A.3 Energy based on the second invariant

The energy for nematic elastomers based on cof \( \mathbf{F} \) is as follows:

\[
\hat{W}_2 = \min_{n \text{ s.t. } |n|=1} \left\{ d \left| \text{tr} \left[ \left( \mathbf{F}^T \ell_n^{-1} \mathbf{F} \right)^{-1} \right] - 3 \right|^q \right\}. \tag{A.9}
\]

As in Section A.2, we will choose to work in the principal basis because \( I_2 \) is an invariant, which is independent of basis:

\[
b = \begin{pmatrix}
\lambda_{\text{max}}^2 \\
\lambda_{\text{mid}}^2 \\
\lambda_{\text{min}}^2
\end{pmatrix}.
\]

Similarly to the case of \( W_1 \), minimizing the energy over the director \( n \) yields the result that \( n \) is parallel to the eigenvector corresponding to the largest eigenvalue, \( \lambda_{\text{max}}^2 \), making \( n = (1, 0, 0)^T \). Thus, the step-length tensor should be as follows:

\[
\ell_n^{-1} = r^{1/3} \left( I + \left( \frac{1}{r} - 1 \right) n \otimes n \right)
\]

\[
= \begin{pmatrix}
r^{-2/3} \\
r^{1/3} \\
r^{1/3}
\end{pmatrix}.
\]

Therefore, we have

\[
\hat{W}_2(s, t) = d \left| \text{tr} \left[ \left( \ell_n^{-1} b \right)^{-1} \right] - 3 \right|^q
\]

\[
= d \left| r^{-1/3} \left( \frac{r}{\lambda_{\text{max}}^2} + \frac{1}{\lambda_{\text{mid}}^2} + \frac{1}{\lambda_{\text{min}}^2} \right) - 3 \right|^q
\]

\[
= d \left| r^{-1/3} \left( \frac{r}{s^2} + \frac{s^2}{r^2} + \frac{1}{t^2} \right) - 3 \right|^q. \tag{A.15}
\]

Again, in the last equality we have used the fact that \( s = \lambda_{\text{max}} \) and \( t = \lambda_{\text{max}} \lambda_{\text{mid}} \).
A.4 Simplification of the regions

The regions are defined by the following constraints:

\[
L : \{(s,t) : t \leq s^2, t \geq \sqrt{s}, t \leq r^{1/6}\}
\]
\[
M : \{(s,t) : t \geq r^{1/6}, t \leq s^2, t \geq r^{-1/2}s^2\}
\]
\[
S : \{(s,t) : t \geq \sqrt{s}, t \leq r^{-1/2}s^2\}
\]

We can rewrite these constraints in terms of the principal stretches of \(F\), \(\lambda_{\text{max}} \geq \lambda_{\text{mid}} \geq \lambda_{\text{min}}\), by using the relations found in Equation A.1. For region \(L\), the first constraint is:

\[
t \leq s^2
\]  
\[
\lambda_{\text{max}}\lambda_{\text{mid}} \leq \lambda_{\text{max}}^2
\]  
\[
\lambda_{\text{mid}} \leq \lambda_{\text{max}}
\]

This is true always, so it is not necessary to state this as a constraint. Similarly, the second constraint simplifies to:

\[
t \geq \sqrt{s}
\]  
\[
\lambda_{\text{max}}\lambda_{\text{mid}} \geq \sqrt{\lambda_{\text{max}}}
\]  
\[
\lambda_{\text{max}}^2\lambda_{\text{mid}}^2 \geq \lambda_{\text{max}}
\]  
\[
\lambda_{\text{max}}\lambda_{\text{mid}}^2 \geq 1
\]  
\[
\frac{1}{\lambda_{\text{min}}} \lambda_{\text{mid}} \geq 1
\]  
\[
\lambda_{\text{mid}} \geq \lambda_{\text{min}}
\]

Again, this relation is true always. Thus, the last constraint in region \(L\) becomes

\[
t \leq r^{1/6}
\]  
\[
\lambda_{\text{max}}\lambda_{\text{mid}} \leq r^{1/6}
\]

In region \(M\), the first constraint is

\[
t \geq r^{1/6}
\]  
\[
\lambda_{\text{max}}\lambda_{\text{mid}} \geq r^{1/6}
\]

The second constraint in \(M\) is the same as the first constraint in region \(L\). The third constraint in \(M\) is

\[
t \geq r^{-1/2}s^2
\]  
\[
\lambda_{\text{max}}\lambda_{\text{mid}} \geq r^{-1/2}\lambda_{\text{max}}^2
\]  
\[
\frac{\lambda_{\text{max}}}{\lambda_{\text{mid}}} \leq r^{1/2}
\]
In region $S$, the first constraint is the same as the second constraint in region $L$. The second constraint in $S$ is similar to the third constraint in $M$:

\[ t \leq r^{-1/2}s^2 \]  
\[ \frac{\lambda_{\text{max}}}{\lambda_{\text{mid}}} \geq r^{1/2} \]  

(A.33)  

(A.34)

Thus, we have:

\[ L : \{ \lambda_{\text{max}} \lambda_{\text{mid}} \leq r^{1/6} \} \]

\[ M : \{ \lambda_{\text{max}} \lambda_{\text{mid}} \geq r^{1/6}, \frac{\lambda_{\text{max}}}{\lambda_{\text{mid}}} \leq \sqrt{r} \} \]  

(A.35)

\[ S : \{ \frac{\lambda_{\text{max}}}{\lambda_{\text{mid}}} \geq \sqrt{r} \} \]
Appendix B

DERIVING DDSDDE FOR THE UMAT

B.1 Useful items for deriving the material Jacobian

Here is a useful derivative:

\[ j = \frac{\partial (\det F')}{\partial F_{ij}} \frac{\partial F_{ij}}{\partial t} \]  \hspace{1cm} (B.1)

\[ = \dot{F}^T F \dot{F} \]  \hspace{1cm} (B.2)

\[ = \dot{F} F \dot{F}^{-1} \]  \hspace{1cm} (B.3)

\[ = J \text{tr} \ell \]  \hspace{1cm} (B.4)

Additionally, we can prove that

\[ \text{tr} \ell = \text{tr} d \]  \hspace{1cm} (B.5)

by

\[ \text{tr} d = \text{tr} \left[ \frac{1}{2} \left( \ell + \ell^\top \right) \right] \]  \hspace{1cm} (B.6)

\[ = \frac{1}{2} \left[ \text{tr}(\ell) + \text{tr}(\ell^\top) \right] \]  \hspace{1cm} (B.7)

\[ = \frac{1}{2} \left[ \text{tr}(\ell) + \text{tr}(\ell) \right] \]  \hspace{1cm} (B.8)

\[ = \text{tr} \ell \]  \hspace{1cm} (B.9)

or the fact that

\[ \text{tr}(\ell) = \text{tr}(d + w) = \text{tr}(d) \]  \hspace{1cm} (B.10)

since \( \text{tr}(w) = 0 \).

B.2 Auxiliary remark for the material Jacobian

The time derivative of the right Cauchy-Green tensor can be related to the rate of deformation tensor:

\[ \dot{C} = \dot{F}^T F \]  \hspace{1cm} (B.11)

\[ = \dot{F}^T F + F^T \dot{F} \]  \hspace{1cm} (B.12)

\[ = F^T F^{-1} \dot{F} + F^T \dot{F} F^{-1} \]  \hspace{1cm} (B.13)

\[ = F^T \ell F + F^T \ell F \]  \hspace{1cm} (B.14)

\[ = 2 F^T dF \]  \hspace{1cm} (B.15)
Thus,

\[
\text{tr} \left( \tilde{G}^{-1} \dot{C} \right) = \text{tr} \left( 2 \tilde{G}^{-1} F^T dF \right) \\
= 2 \text{tr} \left( F \tilde{G}^{-1} F^T \dot{d} \right) \\
= 2 \left( F \tilde{G}^{-1} F^T \right)_{ij} d_{ji}
\]  

(B.16)  

(B.17)  

(B.18)

B.3 Auxiliary remark for the material Jacobian

Given \( A_{ij} = A_{ji}, d_{ij} = d_{ji} \), find \( C_{ijkl} = C_{klij} = C_{ijlk} \) s.t. \( A_{ij} = C_{ijkl}d_{kl} \).

Remark: There are an infinity of solutions. Add any symmetric \( C \) with eigenvectors perpendicular to \( d \).

One possible solution is

\[
C_{ijkl} = a_1(A_{ij}d_{kl} + d_{ij}A_{kl}) + a_2d_{ij}d_{kl}. 
\]

(B.19)

Then,

\[
C_{ijkl}d_{kl} = a_1 A_{ij} |d|^2 + d_{ij}(a_1 A_{kl}d_{kl} + a_2 |d|^2).
\]

(B.20)

Pick: \( a_1 = \frac{1}{|d|^2} \) and \( a_2 = -a_1 \frac{\Delta_{kl}d_{kl}}{|d|^2} = -\frac{\Delta_{kl}d_{kl}}{|d|^4} \) so that we have

\[
C_{ijkl}d_{kl} = A_{ij}.
\]

(B.21)

This means that

\[
C_{ijkl} = \frac{1}{|d|^2} \left( A_{ij}d_{kl} + d_{ij}A_{kl} \right) - \frac{1}{|d|^4} A_{mn}d_{mn}d_{ij}d_{kl}.
\]

(B.22)
SYNTHESIS AND TESTING

C.1 Chemical details

<table>
<thead>
<tr>
<th>Item</th>
<th>Density (g/mL)</th>
<th>Molar weight (g/mol)</th>
<th>Molecular formula</th>
</tr>
</thead>
<tbody>
<tr>
<td>RM257</td>
<td>N/A</td>
<td>588.6</td>
<td>$C_{33}H_{32}O_{10}$</td>
</tr>
<tr>
<td>Toluene</td>
<td>0.865</td>
<td>92.14</td>
<td>$C_7H_8$</td>
</tr>
<tr>
<td>HHMP</td>
<td>N/A</td>
<td>224.25</td>
<td>$C_{12}H_{16}O_4$</td>
</tr>
<tr>
<td>PETMP</td>
<td>1.28</td>
<td>488.66</td>
<td>$C_{17}H_{28}O_8S_4$</td>
</tr>
<tr>
<td>EDDET</td>
<td>1.12</td>
<td>182.3</td>
<td>$C_{6}H_{14}O_2S_2$</td>
</tr>
<tr>
<td>DPA</td>
<td>0.738</td>
<td>101.19</td>
<td>$C_{6}H_{15}N$</td>
</tr>
</tbody>
</table>

C.2 Synthesis template

Synthesis
Name: _______________
Date: _______________
Time start: _______________
Time finish: _______________
Sample type: _______________
Molds used: _______________
mol% PETMP: _______________

To do:

- [ ] Wear PPE
  - [ ] Lab coat
  - [ ] Safety glasses
  - [ ] Gloves
  - [ ] Closed-toe shoes
  - [ ] Tie long hair back
- [ ] Clean mold with IPA, DI water, Kim wipe
- [ ] Prepare 2 vials, labeled as “LCE solution” and “DPA+toluene”
- [ ] Prepare any glass pipettes
Hot plate dial level: ________________
Hot plate temperature: ________________

*Note: This is the triple batch recipe (a larger batch attains better accuracy than the single batch recipe from [64]).*

<table>
<thead>
<tr>
<th>Item</th>
<th>Expected weight (g)</th>
<th>Micropipette volume (μL)</th>
<th>Actual weight, subtotal (g)</th>
<th>Leftover on weigh boat or paper (g)</th>
<th>Actual weight, final (g)</th>
<th>% error</th>
</tr>
</thead>
<tbody>
<tr>
<td>RM257</td>
<td>12 g</td>
<td>—</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Toluene</td>
<td>4.8g</td>
<td>5549.1μL</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>HHMP</td>
<td>0.0257g</td>
<td>—</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Time on the hot plate:
Make catalyst solution (DPA+toluene) in separate vial:

<table>
<thead>
<tr>
<th>Item</th>
<th>Expected weight (g)</th>
<th>Micropipette volume (μL)</th>
<th>Actual weight, subtotal (g)</th>
<th>Leftover on weigh boat or paper (g)</th>
<th>Actual weight, final (g)</th>
<th>% error</th>
</tr>
</thead>
<tbody>
<tr>
<td>DPA</td>
<td>0.0549 g</td>
<td>74.25 μL</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Toluene</td>
<td>0.0549 g</td>
<td>74.25 μL</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Mix catalyst solution on vortex mixer

<table>
<thead>
<tr>
<th>Item</th>
<th>Expected weight (g)</th>
<th>Micropipette volume (μL)</th>
<th>Actual weight, subtotal (g)</th>
<th>Leftover on weigh boat or paper (g)</th>
<th>Actual weight, final (g)</th>
<th>% error</th>
</tr>
</thead>
<tbody>
<tr>
<td>PETMP (50 mol%)</td>
<td>2.1657 g</td>
<td>1691.64 μL</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>EDDET</td>
<td>1.6158g</td>
<td>1442.67μL</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Catalyst solution</td>
<td>1.7043g</td>
<td>1974μL</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Time on vortex mixer: 20 sec
Time in vacuum oven at 20in Hg at room temperature: 45 sec

Did sample recrystallize and need a 2nd time on hot plate? ________________
Solution appearance (cloudy, clear, undissolved solids?): ________________
Picture of sample in mold:

To do:

- [ ] Dispose of all hazardous waste inside plastic bag
- [ ] Clean beakers
- [ ] Clean utensils
- [ ] Turn off scale, hot plate, lights
Notes:

**Placing samples into vacuum oven**
Date: ________________
Time into oven: ________________

To do:

- □ Pull vacuum level to 20.5 in Hg
- □ Set temperature dial to 1.5

Notes:

**Taking samples out of vacuum oven**
Date: ________________
Time out of oven: ________________

Time of transferring samples from mold to sample boxes (should be ≈ 30 min later): ________________

Notes:

**UV Crosslinking**
Date: ________________
Time: ________________

UV crosslinking equipment used: ________________

Amount of time hit by UV light per side: ________________

Pictures:

Notes:

C.3 **Tensile test template**
Name: ________________
Date of test: ________________
Time of test: ________________

Test type (load only, lo ad/unload, pull til failure): ________________

Temperature: ________________
Strain rate: ________________
Velocity: ________________

Linear stage extension: ________________
Total time of test: ________________
Sample details:
Sample type (polydomain/monodomain): ____________
Sample identifier: ____________
Date of synthesis: ____________
Amount of time after taken out of oven: ____________
Dimensions of sample:
Width: ____________
Thickness: ____________
Gauge length: ____________
Notes on heating/cooling history: ____________
Notes on loading history: ____________
Microscope/video?: ____________
How did the sample break: ____________
Pictures and data (stress-strain curve, temperature data, images of sample before/after test, etc.):
Notes:
Appendix D

CODE

D.1 UMAT code for simulations of nematic elastomers in ABAQUS

This section contains the FORTRAN code for the user material defined by Chapter 4, implemented as a UMAT in ABAQUS. Some portions of code are reproduced from [15, 44, 60], explicitly denoted within comments in the code.

```fortran
subroutine umat(stress, statev, ddsdde, sse, spd, scd,
+ rpl, ddsddt, drpdlde, drpdlde,
+ stran, dstran, time, dttime, temp, dtemp, predef, dpred, cmname,
+ ndf, nshr, ntens, nstatv, props, nprops, coords, drot, pnewdt,
+ cort, dfgrd[0], dfgrd[1], noel, npt, layer, kspt, kstep, kinc)
!
! include 'aba_param.inc'
!
! character*80 cmname

dimension stress(ntens), statev(nstatv),
+ ddsdde(ntens,ntens), ddsddt(ntens), drpdlde(ntens),
+ stran(ntens), dstran(ntens), time(2), predef(1), dpred(1),
+ props(nprops), coords(3), drot(3,3), dfgrd[0](3,3), dfgrd[1](3,3)
!
! variables defined and used in the umat:
!
! external myfunc, myconstraint1, myconstraint2

type i,j,k, lx, ly, lz, np

real*8 iden(3,3), material_jacobian(3,3,3,3), stress_3x3(3,3), dPdF(3,3,3,3)
real*8 F_3x3(3,3), detF, trb, b_3x3(3,3), Finv(3,3), C_3x3(3,3), detC
real*8 mu1, mu2, p_prop, r_prop, c_prop, k_prop, alpha_delta, alpha_lambda,
+ kappa, lambda, delta, new_lambda, new_delta
real*8 G_diag(3,3), G_3x3(3,3), Ginv(3,3), Finvtranspose(3,3), trGinvb,
Pstress(3,3)
real*8 eigvalC(3), eigvec_mat(3,3), tol_val, alpha_const
real*8 eigvalb(3), eigvec_mat_b(3,3), dstran_3x3(3,3), prev_F(3,3)
real*8 prevFinv(3,3), 1_3x3(3,3), d_3x3(3,3), trd
real*8 Gtilde_3x3(3,3), Gtildeinv(3,3), trGtildeinvC, testscalar, dnorm,
+ Gtildeinvdot(3,3), prev_Gtildeinv(3,3)
real*8 A_3x3(3,3), AA_3x3(3,3), trA, trAA, trAAd, Gtildedot(3,3), Fdot(3,3),
+ prev_Gtilde(3,3)
real*8 Lambda_3x3(3,3), U_3x3(3,3), _3x3_inv(3,3), R_3x3(3,3), F_3x3_check
(3,3), Gdiainv(3,3), helper(3,3), oldA_3x3(3,3)
real*8 beta, w_3x3(3,3), temp_mat(6)
real*8 zero, one, two, three, four, five, six, nine, third, half
parameter(zero=0.d0, one=1.d0, two=2.d0, three=3.d0, four=4.d0, five=5.d0, six =6.
d0, nine=9.d0, third=1.d0/3.d0, half=1.d0/2.d0)
real*8 x(2), minf, lb(2), ub(2), cldata, c2data, fdata(22)
```

```fortran
end subroutine umat
```
real*8 ps(3), anps(3,3)
integer*8 opt
integer ires ! ires is an integer return value which is positive on success
and negative on failure
include 'nlopt.f'
!
mu1 = props(1)
mu2 = props(2)
kappa = props(3)
r_prop = props(4)
c_prop = props(5)
k_prop = props(6)
alpha_delta = props(7)
alpha_lambda = alpha_delta/100.d0
p_prop = props(8)
!
tol_val = 5.d-5
alpha_const = 5.d-6
beta = 1.d6
!
F_3x3 = dfgrd1
statev(3) = F_3x3(1,1)
statev(4) = F_3x3(1,2)
statev(5) = F_3x3(1,3)
statev(6) = F_3x3(2,1)
statev(7) = F_3x3(2,2)
statev(8) = F_3x3(2,3)
statev(9) = F_3x3(3,1)
statev(10) = F_3x3(3,2)
statev(11) = F_3x3(3,3)
!
identity matrix
call onem(iden)
!
! Compute the relative volume change
call mdet(F_3x3, detF)
!
! Compute the inverse of the deformation gradient
call m3inv(F_3x3, Finv)
!
Finvtranspose = transpose(Finv)
!
! compute the left & right Cauchy-Green tensor
b_3x3 = matmul(F_3x3, transpose(F_3x3)) ! matrix multiplication
C_3x3 = matmul(transpose(F_3x3), F_3x3) ! matrix multiplication
call mdet(C_3x3, detC)
!
! get the eigenvalues and eigenvectors of C_3x3 (in descending order)
write(*,"(f13.15)" 'in main code'
call zerom(temp_mat)
temp_mat(1) = C_3x3(1,1)
temp_mat(2) = C_3x3(2,2)
temp_mat(3) = C_3x3(3,3)
temp_mat(4) = C_3x3(1,2)
temp_mat(5) = C_3x3(1,3)
temp_mat(6) = C_3x3(2,3)
np = 3
call sprints(temp_mat, eigvalC, eigvec_mat, 1, ndi, nshr)
eigvec_mat = transpose(eigvec_mat)
call eigsrt(eigvalC, eigvec_mat, 3, np)
call m3inv(dfgrd0, prevFinv)
l_3x3 = one/dtime*(matmul(F_3x3, prevFinv)-iden)
d_3x3 = half*(l_3x3 + transpose(l_3x3))
w_3x3 = half*(l_3x3 - transpose(l_3x3))
call tracem(d_3x3, trd)
call tracem(matmul(d_3x3, transpose(d_3x3)), dnorm)

! at the start of an abaqus calculation, the state
! variables are passed into umat with zero values.
! initialize the state variables. at this point,
! the time total_time and step_time both have a value
! equal to zero and the step counter, kstep, is
! equal to 1.
if ((time(1).eq.zero).and.(kstep.eq.1)) then
  lambda = one
  statev(1) = lambda
  delta = one
  statev(2) = delta
  stress_3x3 = zero
  !
end if

! store the values of the state variables at the beginning of the time step
lambda = statev(1) ! at the previous time step
delta = statev(2) ! at the previous time step
!
!! UPDATE LAMBDA AND DELTA USING NLOpt
opt = 0.d0
!call nlo_create(opt, NLOPT_LN_COBYLA, 2)
call nlo_create(opt, NLOPT_LD_MMA, 2) ! 2nd argument is the name of the
optimization algorithm
lb(1) = 1.d0
lb(2) = 1.d0
ub(1) = r_prop**(1.d0/3.d0)
ub(2) = r_prop**(1.d0/6.d0)
call nlo_set_lower_bounds(ires, opt, lb)
call nlo_set_upper_bounds(ires, opt, ub)
fdata(1) = mu1
fdata(2) = mu2
fdata(3) = detF
fdata(4) = kappa
fdata(5) = c_prop
fdata(6) = r_prop
fdata(7) = k_prop
fdata(8) = dtime
fdata(9) = alpha_lambda
fdata(10) = alpha_delta
fdata(11) = lambda
fdata(12) = delta
fdata(13) = p_prop
fdata(14) = F_3x3(1,1)
fdata(15) = F_3x3(1,2)
fdata(16) = F_3x3(1,3)
fdata(17) = F_3x3(2,1)
fdata(18) = F_3x3(2,2)
fdata(19) = F_3x3(2,3)
fdata(20) = F_3x3(3,1)
fdata(21) = F_3x3(3,2)
fdata(22) = F_3x3(3,3)
call nlo_set_min_objective(ires , opt , myfunc , fdata)
call nlo_add_inequality_constraint(ires , opt , myconstraint1 , c1data , 1.d-8)
call nlo_add_inequality_constraint(ires , opt , myconstraint2 , c2data , 1.d-8)
call nlo_set_xtol_rel(ires , opt , 1.d-8)
! initialize w/ previous values
x(1) = lambda
x(2) = delta
!
call nlo_optimize(ires , opt , x, minf)
!if (ires.lt.0) then
!  write(*,*) 'kstep =', kstep , 'increment num = ', kinc , 'total time = ',
time(2)
!  write(*,*) 'nlopt failed! ires = ', ires
!else
!  write(*,*) 'kstep =', kstep , 'increment num = ', kinc , 'total time = ',
time(2)
!  write(*,*) x(1) , x(2) , F_3x3(1,1)
!  write(*,*) 'min val = ', minf , 'and ires =', ires
!endif
call nlo_destroy(opt)
! end of the Nlopt scheme
!
new_lambda = x(1)
new_delta = x(2)
!
statev(1) = new_lambda
statev(2) = new_delta
!
! calculate G_diag
call zerom(G_diag)
G_diag(1,1) = new_lambda**two ! double asterisk ** is for raising to a power
G_diag(2,2) = new_delta**two/(new_lambda**two)
G_diag(3,3) = one/(new_delta**two)
call m3inv(G_diag ,Gdiaginv)
!
Gtilde_3x3 = matmul(eigvec_mat ,matmul(G_diag ,transpose(eigvec_mat))))
call m3inv(Gtilde_3x3,Gtildeinv)
A_3x3 = matmul(F_3x3,matmul(Gtildeinv,transpose(F_3x3))))
call tracem(A_3x3 , trA)
!
! 3x3 cauchy stress
stress_3x3 = (mu1*detF**(-two/three)*(A_3x3 -(one/three)*trA*iden)
  + kappa*dlog(detF)*iden)/detF + beta*d_3x3
transform 3x3 stress into voigt notation

stress(1) = stress_3x3(1,1)
stress(2) = stress_3x3(2,2)
stress(3) = stress_3x3(3,3)
stress(4) = stress_3x3(1,2)
stress(5) = stress_3x3(1,3)
stress(6) = stress_3x3(2,3)

! calculate the material jacobian

material_jacobian = zero
if (dnorm.lt.tol_val) then
  do i=1,3
    do j=1,3
      do k=1,3
        do l=1,3
          material_jacobian(i,j,k,l) = material_jacobian(i,j,k,l)
          + (two/three)*mu1*detF**(-five/three)*(A_3x3(i,j)*iden(k,l)+A_3x3(k,l)*iden(i,j))
          + (two/nine)*mu1*detF**(-five/three)*trA*iden(i,j)*iden(k,l)
          + kappa/detF*iden(i,j)*iden(k,l)
          + mu1/two*detF**(-five/three)*(A_3x3(1,1)*iden(i,k)+A_3x3(k,1)*iden(i,1))
          + mull/two*detF**(-five/three)*(A_3x3(1,1)*iden(j,k)+A_3x3(j,1)*iden(j,1))
          + beta/two*trd*(iden(k,l)*iden(j,l)+iden(j,l)*iden(k,l))
          + beta/(two*dtime)*(iden(i,j)*iden(j,l)+iden(j,l)*iden(i,j))
          - two*mu1*alpha_const*detF**(-five/three)*
            (one/(dnorm**two)*(d_3x3(i,j)*d_3x3(k,l)-third*trd*(iden(i,j)*d_3x3(k,l))
            +d_3x3(i,j)*iden(k,l))
          + one/(dnorm**four)*third*(trd**two)*d_3x3(i,j)*d_3x3(k,l))
          + beta/two*trd*(iden(i,k)*iden(j,l)+iden(j,k)*iden(i,l))
        enddo
      enddo
    enddo
  enddo
else
  do i=1,3
    do j=1,3
      do k=1,3
        do l=1,3
          material_jacobian(i,j,k,l) = material_jacobian(i,j,k,l)
          + (two/three)*mu1*detF**(-five/three)*(A_3x3(i,j)*iden(k,l)+A_3x3(k,l)*iden(i,j))
          + (two/nine)*mu1*detF**(-five/three)*trA*iden(i,j)*iden(k,l)
          + kappa/detF*iden(i,j)*iden(k,l)
          + mu1/two*detF**(-five/three)*(A_3x3(1,1)*iden(i,k)+A_3x3(k,1)*iden(i,1))
          + mull/two*detF**(-five/three)*(A_3x3(1,1)*iden(j,k)+A_3x3(j,1)*iden(j,1))
          - two*mu1*alpha_const*detF**(-five/three)*
            (one/(dnorm**two)*(d_3x3(i,j)*d_3x3(k,l)-third*trd*(iden(i,j)*d_3x3(k,l))
            +d_3x3(i,j)*iden(k,l))
          + one/(dnorm**four)*third*(trd**two)*d_3x3(i,j)*d_3x3(k,l))
          + beta/two*trd*(iden(i,k)*iden(j,l)+iden(j,k)*iden(i,l))
        enddo
      enddo
    enddo
  enddo
end if
+ beta/two*iden(i,k)*w_3x3(1,j)+iden(i,l)*w_3x3(k,j)+w_3x3(k,i)*iden(1,l)+w_3x3(1,i)*iden(j,k))
+ beta/(two*dtime)*(iden(i,k)*iden(j,l)+iden(j,k)*iden(i,l))
enddo
enddo
enddo
end if

!**********************************************************************
! transform the tangent matrix into voigt notation. code from
! Alexander K. Landauer, Xiuqi Li, Christian Franck, and David L. Henann.
! Experimental characterization and hyperelastic constitutive modeling of
! open-cell elastomeric foams. Journal of the Mechanics and Physics of
! Solids
!**********************************************************************

ddsdde = zero
!
do i=1,ndi
do j=1,ndi
  ddsdde(1,j) = material_jacobian(i,i,j,j)
enddo
enddo
!
if (nshr.ne.0) then
do i=1,ndi
  ddsdde(i,ndi+1) = material_jacobian(i,i,1,2)
  ddsdde(ndi+1,i) = material_jacobian(1,2,i,i)
enddo

if (nshr.ne.1) then
  ddsdde(ndi+1,ndi+1) = material_jacobian(1,2,1,2)
  ddsdde(1,ndi+2) = material_jacobian(i,i,1,3)
  ddsdde(ndi+2,i) = material_jacobian(1,3,i,i)
enddo

if (nshr.ne.2) then
  ddsdde(ndi+2,ndi+2) = material_jacobian(1,3,1,3)
  ddsdde(ndi+2,ndi+1) = material_jacobian(1,3,1,2)
  ddsdde(1,ndi+3) = material_jacobian(i,i,2,3)
  ddsdde(ndi+3,i) = material_jacobian(2,3,i,i)
enddo

end if
end if
return
end subroutine umat
!

!**********************************************************************
subroutine myfunc(func_val, n, x, grad, need_gradient, fdata)

double precision x(n), grad(n), fdata(22)

  real*8 func_val
  integer i, j, n, need_gradient, nrot, np, ndi, nshr
  real*8 mu1, mu2, detF, kappa, c_prop, r_prop, k_prop, dtime, alpha_lambda,
  alpha_delta, lambda, delta, p_prop
  real*8 F_3x3(3,3), temp1(3,3), temp2(3,3), trtemp1, trtemp2
  real*8 C_3x3(3,3), eigvalC(3), eigvec_mat(3,3), G_diag(3,3), Gtilde_3x3(3,3),
  Gtildeinv(3,3), A_3x3(3,3), trA
  real*8 dGtildedlambda(3,3), dGtildeddelta(3,3),term1(3,3), trterm1, term2(3,3)
, trterm2, dDdlambda, dDddelta, dfddelta
  real*8 W_val, f_val, diss_val, temp_mat(6)

parameter(zero=0.d0, one=1.d0, two=2.d0, three=3.d0, four=4.d0, five=5.d0, six =6.
, d0, nine=9.d0, third=1.d0/3.d0, half=1.d0/2.d0)

mu1 = fdata(1)
mu2 = fdata(2)
detF = fdata(3)
kappa = fdata(4)
c_prop = fdata(5)
r_prop = fdata(6)
k_prop = fdata(7)
dtime = fdata(8)
alpha_lambda = fdata(9)
alpha_delta = fdata(10)
lambda = fdata(11)
delta = fdata(12)
p_prop = fdata(13)
F_3x3(1,1) = fdata(14)
F_3x3(1,2) = fdata(15)
F_3x3(1,3) = fdata(16)
F_3x3(2,1) = fdata(17)
F_3x3(2,2) = fdata(18)
F_3x3(2,3) = fdata(19)
F_3x3(3,1) = fdata(20)
F_3x3(3,2) = fdata(21)
F_3x3(3,3) = fdata(22)
!
!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!! func_val
!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!

np = 3
ndi = 3
nshr = 3
C_3x3 = matmul(transpose(F_3x3), F_3x3)
call zerom(temp_mat)
temp_mat(1) = C_3x3(1,1)
temp_mat(2) = C_3x3(2,2)
temp_mat(3) = C_3x3(3,3)
temp_mat(4) = C_3x3(1,2)
temp_mat(5) = C_3x3(1,3)
temp_mat(6) = C_3x3(2,3)
call sprind(temp_mat, eigvalC, eigvec_mat, 1, ndi, nshr)
eigvec_mat = transpose(eigvec_mat)
call eigsrt(eigvalC, eigvec_mat, 3, np)
call zerom(G_diag)
G_diag(1,1) = x(1)**two ! double asterisk ** is for raising to a power
G_diag(2,2) = x(2)**two/(x(1)**two)
G_diag(3,3) = one/(x(2)**two)
Gtilde_3x3 = matmul(eigvec_mat, matmul(G_diag, transpose(eigvec_mat)))
call m3inv(Gtilde_3x3, Gtildeinv)
A_3x3 = matmul(F_3x3, matmul(Gtildeinv, transpose(F_3x3)))
call tracem(A_3x3, trA)
W_val = (mu1/two)*(detF**(-two/three)*trA-three) + (kappa/two)*(dlog(detF))**two
f_val = c_prop*(x(2)-one)/((r_prop**(one/six)-x(2))**k_prop)
diss_val = (alpha_lambda/two)*((x(1)-lambda)/dtime)**p_prop + (alpha_delta/two)*((x(2)-delta)/dtime)**p_prop
func_val = W_val + f_val + dtime*diss_val
!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!! gradient
!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!
call zerom(dGtildedlambda)
dGtildedlambda(1,1) = two*x(1)
dGtildedlambda(2,2) = -two*x(2)**two/(x(1)**three)
dGtildedlambda = matmul(eigvec_mat, matmul(dGtildedlambda, transpose(eigvec_mat)))
call zerom(dGtildeddelta)
dGtildeddelta(2,2) = two*x(2)/(x(1)**two)
dGtildeddelta(3,3) = -two/(x(2)**three)
dGtildeddelta = matmul(eigvec_mat, matmul(dGtildeddelta, transpose(eigvec_mat)))
call tracem(term1, trterm1)
term1 = matmul(Gtildeinv, matmul(dGtildedlambda, matmul(Gtildeinv, C_3x3)))
call tracem(term2, trterm2)
term2 = matmul(Gtildeinv, matmul(dGtildeddelta, matmul(Gtildeinv, C_3x3)))
call tracem(term2, trterm2)
!
ddlambda = -(mu1/two)*detF**(-two/three)*trterm1
dddlambda = -(mu1/two)*detF**(-two/three)*trterm2
!
ddlambda = (alpha_lambda/two)*(p_prop/dtime)*((x(1)-lambda)/dtime)**(p_prop-one)
dddlambda = (alpha_delta/two)*(p_prop/dtime)*((x(2)-delta)/dtime)**(p_prop-one)
!
ddfdelta = c_prop*((r_prop**(one/six)-x(2)+(x(2)-one)*k_prop)/((r_prop**(one/six)-x(2))**(k_prop+one)))
if (needGradient.ne.0) then
  grad(1) = dWdlambda + dtime*ddlambda
  grad(2) = dWddelta + dffdelta + dtime*ddddelta
endif
end
!
subroutine myconstraint1(func_val, n, x, grad, needGradient, d)
integer need_gradient
real*8 func_val, x(n), grad(n)
if (need_gradient.ne.0) then
  grad(1) = 0.5d0/dsqrt(x(1))
  grad(2) = -1.d0
endif
func_val = dsqrt(x(1)) - x(2)
end

!***********************************************************
!
subroutine myconstraint2(func_val , n, x, grad , need_gradient , d)
integer need_gradient
real*8 func_val , x(n), grad(n)
if (need_gradient.ne.0) then
  grad(1) = -2.d0*x(1)
  grad(2) = 1.d0
endif
func_val = x(2) - x(1)**2.d0
end

!****************************************************************************
! utility subroutines
!****************************************************************************
!****************************************************************************
!
subroutine onem(a)
!
! this subroutine initializes the 3x3 matrix a as the identity matrix
!
implicit none
!
integer i,j
!
real*8 a(3,3)
!
do i=1,3
do j=1,3
  if (i .eq. j) then
    a(i,j) = 1.d0
  else
    a(i,j) = 0.d0
  end if
end do
end do
!
return
end subroutine onem

!****************************************************************************
! this subroutine sets all entries of a 3 by 3 matrix to 0.d0.
!****************************************************************************
!
subroutine zerom(a)
real*8 a(3,3)
!
return
end subroutine zerom
do 1 i=1,3
  do 1 j=1,3
    a(i,j) = 0.d0
1       continue
!
return
end

!*********************************************************************
! this subroutine calculates the trace of a 3 by 3 matrix [a]
! and stores the result in the scalar tra
!*********************************************************************
subroutine tracem(a,tra)
  real*8 a(3,3),tra
  tra = a(1,1) + a(2,2) + a(3,3)
  return
end

!*********************************************************************
! this subroutine calculates the determinant
! of a 3 by 3 matrix [a].
!*********************************************************************
subroutine mdet(a, det)
  real*8 a(3,3), det
  det =   a(1,1)*a(2,2)*a(3,3)
     +   a(1,1)*a(2,3)*a(3,2)
     +   a(1,2)*a(2,3)*a(3,1)
     +   a(2,1)*a(2,3)*a(3,1)
     -   a(3,1)*a(2,2)*a(1,3)
     -   a(3,1)*a(2,3)*a(1,1)
     -   a(3,2)*a(2,1)*a(1,2)
  return
end

!*********************************************************************
! this subroutine calculates the transpose of a 3 by 3
! matrix [a], and places the result in atrans.
!*********************************************************************
subroutine mtrans(a,atrans)
  real*8 a(3,3), atrans(3,3)
  do 1 i=1,3
    do 1 j=1,3
      atrans(j,i) = a(i,j)
1       continue
  return
end
**Subroutine m3inv**

Given a 3x3 matrix \( a \) with determinant \( \det \), this subroutine calculates its inverse \( a^{-1} \) and places the result in \( a^{-1} \).

- **Input**:
  - \( a(3,3) \) -- the matrix whose inverse is desired.
  - \( \det \) -- the computed determinant of \( a \).
  - \( \text{acoefac}(3,3) \) -- the matrix of cofactors of \( a(i,j) \).
  - \( \text{aadj}(3,3) \) -- the adjoint of \( a \). It is the matrix obtained by replacing each element of \( a \) by its cofactor and then taking the transpose of the resulting matrix.

- **Output**:
  - \( \text{ainv}(3,3) \) -- returned as inverse of \( a \).
  - \( [\text{ainv}] = [\text{aadj}] / \det \).

**Algorithm**:

1. Call `mdet` to calculate the determinant of \( a \).
2. If \( \det = 0 \), print an error message and stop.
3. Call `mcofac` to calculate the cofactors of \( a \).
4. Call `mtrans` to transpose the cofactor matrix.
5. For each element \( a_{ij} \) in \( a \):
   - Calculate \( \text{ainv}_{ij} = \text{aadj}_{ij} / \det \).

**Source Code**:

```fortran
subroutine m3inv(a,ainv)
    real*8 a(3,3), ainv(3,3), det , acofac(3,3), aadj(3,3)
    call mdet(a,det)
    if ( det .eq. 0.d0 ) then
        write(*,10)
        stop
    endif
    call mcofac(a,acofac)
    call mtrans(acofac,aadj)
    do 1 i = 1,3
        do 1 j = 1,3
            ainv(i,j) = aadj(i,j)/det
        1 continue
    10 format(5x,'--error in m3inv --- the matrix is singular',/,
           + 10x,'program terminated')
    return
end
```

---

**Subroutine mcofac**

This subroutine calculates the cofactor of a 3x3 matrix \( a \), and places the result in \( \text{acoefac} \).

**Algorithm**:

1. Calculate the cofactor for each element of \( a \).

**Source Code**:

```fortran
subroutine mcofac(a,acofac)
    real*8 a(3,3), acofac(3,3)
    acofac(1,1) = a(2,2)*a(3,3) - a(3,2)*a(2,3)
    acofac(1,2) = -(a(2,1)*a(3,3) - a(3,1)*a(2,3))
    acofac(1,3) = a(2,1)*a(3,2) - a(3,1)*a(2,2)
    acofac(2,1) = -(a(1,2)*a(3,3) - a(3,2)*a(1,3))
    acofac(2,2) = a(1,1)*a(3,3) - a(3,1)*a(1,3)
    acofac(2,3) = -(a(1,1)*a(3,2) - a(3,1)*a(1,2))
    acofac(3,1) = a(1,2)*a(2,3) - a(2,2)*a(1,3)
    acofac(3,2) = -(a(1,1)*a(2,3) - a(2,1)*a(1,3))
    acofac(3,3) = a(1,1)*a(2,2) - a(2,1)*a(1,2)
```

---

**Subroutine mdet**

This subroutine calculates the determinant of a 3x3 matrix \( a \).

**Algorithm**:

1. Calculate the determinant using the formula for a 3x3 matrix.

**Source Code**:

```fortran
subroutine mdet(a,det)
    real*8 a(3,3), det
    det = a(1,1)*(a(2,2)*a(3,3) - a(3,2)*a(2,3))
       - a(1,2)*(a(2,1)*a(3,3) - a(3,1)*a(2,3))
       + a(1,3)*(a(2,1)*a(3,2) - a(3,1)*a(2,2))
```
**Program 4.4**

```fortran
!**********************************************************************
! Computes all eigenvalues and eigenvectors of a real symmetric matrix a, which is of size n
! by n, stored in a physical np by np array. On output, elements of a above
! the diagonal are
! destroyed. d returns the eigenvalues of a in its first n elements. v is a
! matrix with the same
! logical and physical dimensions as a, whose columns contain, on output,
! the normalized
! eigenvectors of a. nrot returns the number of Jacobi rotations that were
! required.
!-----------------------------------------------------------

subroutine Jacobi(a,n,np,d,v,nrot)
!
! input ...
! a(n,n): real symmetric matrix
! np:
! d(n): eigenvalues of a
! v(n,n): columns contain normalized eigenvectors of a
! nrot: # jacobi rotations that were required
!
! Recipes in F77. Numerical Methods for Fortran 77: The Art of Scientific
!
implicit none ! there's no "implicit none" in the numerical recipes

INTEGER n,np,nrot,NMAX
REAL*8 a(np,np),d(np),v(np,np)
PARAMETER (NMAX=500)
INTEGER i,ip,iq,j
REAL c,g,h,s,m,t,tau,theta,tresh,b(NMAX),z(NMAX)
real*8 zero,one,two,three,four,five,six,nine,third,half
parameter(zero=0.d0,one=1.d0,two=2.d0,three=3.d0,four=4.d0,five=5.d0,six=6.d0,nine=9.d0,third=1.d0/3.d0,half=1.d0/2.d0)
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!
do ip=1,n-1 !Sum off-diagonal elements.
   do iq=ip+1,n
      sm=sm+abs(a(ip,iq))
   enddo
enddo
if(sm.eq.0.d0) return ! The normal return, which relies on quadratic conver
if(i.lt.4) then ! gence to machine underflow.
   tresh=0.2d0*sm/n**2.d0 !...on the first three sweeps.
else
   tresh=0.d0 !...thereafter.
endif
! do ip=1,n-1
   do iq=ip+1,n
      g=100.d0*abs(a(ip,iq)) !After four sweeps, skip the rotation if the
      ! off-diagonal element is small.
      if((i.gt.4).and.(abs(d(ip))+g.eq.abs(d(ip))).and.(abs(d(iq))+g.eq.abs(d(iq))))then
         a(ip,iq)=0.d0
      else if(abs(a(ip,iq)).gt.tresh)then
         h=d(iq)-d(ip)
         if(abs(h)+g.eq.abs(h))then
            t=a(ip,iq)/h ! t = 1/(2theta)
         else
            theta=0.5d0*h/a(ip,iq) !Equation (11.1.10).
            t=1.d0/(abs(theta)+sqrt(1.d0+theta**2.d0))
         endif
         c=1.d0/sqrt(1.d0+t**2.d0)
         s=t*c
         tau=s/(1.d0+c)
         h=t*a(ip,iq)
         z(ip)=z[ip]-h
         z(iq)=z(iq)+h
         d(ip)=d(ip)-h
         d(iq)=d(iq)+h
         a(ip,iq)=0.d0
      do j=1,ip-1 !Case of rotations 1 \leq j \leq p.
         g=a(j,ip)
         h=a(j,iq)
         a(j,ip)=g-s*(h+g*tau)
         a(j,iq)=h+s*(g-h*tau)
      enddo
      do j=ip+1,iq-1 !Case of rotations p<j<q.
         g=a(ip,j)
         h=a(j,iq)
         a(ip,j)=g-s*(h+g*tau)
         a(j,iq)=h+s*(g-h*tau)
      enddo
      do j=iq+1,n !Case of rotations q<j \leq n.
         g=a(ip,j)
         h=a(iq,j)
         a(ip,j)=g-s*(h+g*tau)
         a(iq,j)=h+s*(g-h*tau)
QVR
VUU
enddo
VUV
do j=1,n
VUW
g=v(j,ip)
VUX
h=v(j,iq)
VUY
v(j,ip)=g-s*(h+g*tau)
VVP
v(j,iq)=h+s*(g-h*tau)
VVQ
enddo
VVV
nrot=nrot+1
VVR
endif
VVS
enddo
VVU
enddo
VVV

!**********************************************************************
VWW
subroutine eigsrt(d,v,n,np)
VWX
! Given the eigenvalues d and eigenvectors v as output from jacobi (11.1) or
tqli (11.3),
VWY
! this routine sorts the eigenvalues into descending order, and rearranges
the columns of v correspondingly.
VXW
! The method is straight insertion.
VYX
!
VZV
! Press, William H, Saul A Teukolsky, William T Vetterling, and Brian P
Flannery. Numerical
VZV
! Recipes in F77. Numerical Methods for Fortran 77: The Art of Scientific
VZV
implicit none
VZV
INTEGER n,np
VZV
REAL*8 d(np),v(np,np)
VZV
INTEGER i,j,k
VZV
REAL p
VZV
do i=1,n-1
VZV
k=i
VZV
p=d(i)
VZV
do j=i+1,n
VZV
if(d(j).ge.p)then
VZV
  k=j
VZV
  p=d(j)
VZV
endif
VZV
endif
VZV
if(k.ne.i)then
VZV
  d(k)=d(i)
VZV
  d(i)=p
VZV
do j=1,n
VZV
  p=v(j,i)
VZV
  v(j,i)=v(j,k)
VZV
  v(j,k)=p
VZV
endo
! this is a helper function, uncomment as needed to see eigenvalues, eigenvectors

subroutine eighelp(A_3x3, eigvalA, eigvec_mat_A, n, np, nrot)
  INTEGER n, np, nrot
  REAL*8 A_3x3(np, np), eigvalA(np), eigvec_mat_A(np, np)
  real*8 temp_mat(3,3), LHS(3), RHS(3), vec1(3), vec2(3), vec3(3)
  real*8 temp_eigvec(3)
  INTEGER i, j, k

  !write(*,*) 'original matrix'
  !write(*,*) A_3x3(1,1), A_3x3(1,2), A_3x3(1,3)
  !write(*,*) A_3x3(2,1), A_3x3(2,2), A_3x3(2,3)
  !write(*,*) A_3x3(3,1), A_3x3(3,2), A_3x3(3,3)
  !write (*,202) ! eigenvalues
  !write(*,*) eigvalA(1), eigvalA(2), eigvalA(3)
  !write (*,203) ! eigenvectors
  !write (*,*) eigvec_mat_A(1,1), eigvec_mat_A(1,2), eigvec_mat_A(1,3)
  !write (*,*) eigvec_mat_A(2,1), eigvec_mat_A(2,2), eigvec_mat_A(2,3)
  !write (*,*) eigvec_mat_A(3,1), eigvec_mat_A(3,2), eigvec_mat_A(3,3)
  ! rename the eigenvectors to vector vec1, vec2, and vec3
  do j = 1,3
    vec1(j) = eigvec_mat_A(j,1)
    vec2(j) = eigvec_mat_A(j,2)
    vec3(j) = eigvec_mat_A(j,3)
  end do
  !write (*,*) vec1(1), vec2(1), vec3(1)
  !write (*,*) vec1(2), vec2(2), vec3(2)
  !write (*,*) vec1(3), vec2(3), vec3(3)
  call zerom(temp_mat)
  temp_mat(1,1) = eigvalA(1)
  temp_mat(2,2) = eigvalA(2)
  temp_mat(3,3) = eigvalA(3)
  temp_mat = A_3x3 - matmul(eigvec_mat_A, matmul(temp_mat, transpose( eigvec_mat_A)))
  !write(*,*) 'original matrix minus reconstructed matrix'
  !write(*,*) temp_mat(1,1), temp_mat(1,2), temp_mat(1,3)
  !write(*,*) temp_mat(2,1), temp_mat(2,2), temp_mat(2,3)
  !write(*,*) temp_mat(3,1), temp_mat(3,2), temp_mat(3,3)
  201 format (6f12.6)
  202 format (/,' Eigenvalues')
  203 format (/,' Eigenvectors')

  !write(*,*) 'lets check A_{ij}*v_j = eigval_i*v_i'
  !write(*,*) 'eigvalA(1)'
  LHS = 0.D0
  do i=1,3
    do j = 1,3
      LHS(i) = LHS(i) + A_3x3(i,j)*vec1(j)
    enddo
  enddo
```matlab
endo
RHS(i) = eigvalA(1)*vec1(i)

tempeigvec(i) = LHS(i) - RHS(i)
endo
!write(*,*) 'LHS - RHS vector =', (tempeigvec(i),i=1,3)
!
!write(*,*) 'eigvalA(2)'
LHS = 0.D0
do i=1,3
do j = 1,3
   LHS(i) = LHS(i) + A_3x3(i,j)*vec2(j)
endo
RHS(i) = eigvalA(2)*vec2(i)
tempeigvec(i) = LHS(i) - RHS(i)
endo
!write(*,*) 'LHS - RHS vector =', (tempeigvec(i),i=1,3)
!
!write(*,*) 'eigvalA(3)'
LHS = 0.D0
do i=1,3
do j = 1,3
   LHS(i) = LHS(i) + A_3x3(i,j)*vec3(j)
endo
RHS(i) = eigvalA(3)*vec3(i)
tempeigvec(i) = LHS(i) - RHS(i)
endo
!write(*,*) 'LHS - RHS vector =', (tempeigvec(i),i=1,3)
!
return
end
```

D.2 MATLAB code for the thermo-mechanical characterization experiments

We include the code to run a uniaxial test using our custom setup, as discussed in Chapter 5. Note that the code is specific to the type of linear stage and controller, the load cell, data acquisition system, and camera. See Section 5.3 for the equipment list. We include the following code:

- **main.m**: the main code with instructions for setting up, executing, and cleaning up a test
- **test_parameters.m**: the function that defines test parameters
- **init_load.m**: the function that initiates the data acquisition
- **get_load.m**: the function that obtains data from the DAQ
- **PI_move.m**: the function that controls movement of the linear stage
The main.m code:

```matlab
%% CONSTANTS

% imput sample geometry here
width = .280; % inches, sample width
thickness = .051; % inches, sample thickness
eps = 500; % [%] desired strain
epsdot = 4.5*10^-3; %[1/sec]

% global variables
global pos_data;
global temp_data;
global load_data;
global time_data;
global datetime_starttest;
startCoarsePos = 249;
jog_step = 1; %mm
jog_velocity = 4; % mm/sec

% plotting constants
linewidth = 1.5;
fontsize = 16;
fontsizemult = 1.2;

%% Connect to controller and reference stage (without sample connected)
% the stage will move to reference position of 102mm (the middle of the stage)
% this is for a single axis that is connected via USB.
% if you are using a different controller & stage, please set up in
% MikroMove first, then edit accordingly in connect_and_reference_loaner().
if ~exist('PIdevice')
    [PIdevice, axis, Controller] = connect_and_reference_loaner();
end

disp('')
disp('linear stage is done moving to reference position. position in mm =')
disp(PIdevice.qPOS(axis))

%% put sample into top clamp
% Move to coarse start position and attach pullrod
PIdevice.VEL(axis, jog_velocity);
PIdevice.MOV(axis, startCoarsePos);
while(abs(PIdevice.qPOS(axis) - startCoarsePos) > 0.001)
    continue
end

%% Jogging: move up/down to get specimen clamped in position
PIdevice.VEL(axis, jog_velocity);
u = 'u';
d = 'd';
while (1)
    dir = input('Move u or d? (hit enter when finished jogging.) ');
```
if (dir == 'u')
    PIdevice.MOV(axis, PIdevice.qPOS(axis)-jog_step);
else if (dir == 'd')
    PIdevice.MOV(axis, PIdevice.qPOS(axis)+jog_step);
else
    break
end

clear u d

%% calculate gauge length
currPos = PIdevice.qPOS(axis); % mm
gaugelength_theoretical_mm = 273.6978 - currPos; %mm
gaugelength_theoretical_in = gaugelength_theoretical_mm * 0.0393701;
% measure and input the gauge length!
gaugelength_measured_in = .375; % inches, sample gauge length
gaugelength_measured_mm = SI(gaugelength_measured_in, 'in') * 10^3; %mm

abs(gaugelength_theoretical_in - gaugelength_measured_in)
gaugelength = gaugelength_theoretical_mm;
gaugelength_in = gaugelength_theoretical_in;

%% calculate test parameters
[extension, test_velocity, tmax, epsdot] = test_parameters(eps, gaugelength_theoretical_in, 'in', epsdot);
fprintf('epsdot = %.3e
', epsdot)
fprintf('test_velocity = %.3e
', test_velocity) % test_velocity in mm/sec
% extension: mm, this is how much total distance you want stage to
% move between startPos & stopPos
% this is positive if you want stage to physically move up
% (towards 0mm)
crosssecarea = width*thickness*(25.4)^2; %mm^2
sample_period = 1; % sec (must be >= 0.5)
step = test_velocity*sample_period; % mm

% Configure linear stage
startPos = PIdevice.qPOS(axis);
stopPos = startPos - extension;
if (startPos > stopPos)
    steps = [startPos:-step:stopPos, (stopPos+step):step:startPos];
else
    steps = [startPos:step:stopPos, (stopPos-step):-step:startPos];
end
max(steps)
min(steps)

%% Initialize inet + collect preheating data
if ~exist('inet')
    inet = init_load(false, 'tori_RTD_louisaload_jun252020'); % prf file
end
numb = 100000;
preheatload=NaN*zeros(size(numb));
preheattemp=NaN*zeros(size(numb));

for i = 1:numb
    pause(2) % wait every 2 sec
    [preheattemp(i), preheatload(i)] = get_load(inet);
end

% type ctrl+c when preheating is finished

endnum = 410; % input the last index when you stop the preheating
preheattime = linspace(1,endnum*2,endnum);
figure;
subplot(2,1,1)
plot(preheattime ,preheattemp(1:endnum))
xlabel('time (s)')
ylabel('temp (deg F)')
subplot(2,1,2)
plot(preheattime ,preheatload)
xlabel('time (s)')
ylabel('load (lbs)')

% save('preheat_data.mat','preheattime','preheatload','preheattemp')

% MOVE
% Move routine

% extension data is saved in this variable pos_data
% even if you interrupt the script using ctrl+C, it will save the data in
% this variable, also saved in a .mat file

% create vector of zeros to fill to speed up data collection
pos_data = NaN*zeros(size(steps));
temp_data = NaN*zeros(size(steps));
load_data = NaN*zeros(size(steps));
time_data = NaN*zeros(size(steps));
PIdevice.VEL(axis , test_velocity);
move(startPos , stopPos , step , steps , PIdevice , axis , inet)

% calculate the actual test_velocity
actual_test_velocity = diff(pos_data)./diff(time_data);
output_data(1,1) = test_velocity;
output_data(1,2) = mean(actual_test_velocity(actual_test_velocity>0));
output_data(1,3) = std(actual_test_velocity(actual_test_velocity>0));
output_data(1,:) % print out these items in the command window

% Plot raw data
N = length(pos_data);
new_pos_data = zeros(N,1);
for i=1:N
    new_pos_data(i) = pos_data(1)-pos_data(i);
end

halfway=floor(N/2);

figure;
subplot(2,1,1)
hold on;
plot(new_pos_data(1:halfway), load_data(1:halfway), '.-');
plot(new_pos_data(halfway+1:end), load_data(halfway+1:end), '.-');
plot0a=plot(new_pos_data, load_data, '.-');
xlabel('extension, mm');
ylabel('load, lbs');

subplot(2,1,2)
plot0b=plot(pos_data, temp_data, '.-');
xlabel('extension, mm');
ylabel('temperature, degrees');
saveas(plot0a, 'load_extension', 'png'); %saves it in the folder where this .m file is
saveas(plot0b, 'temp_extension', 'png'); %saves it in the folder where this .m file is

% Plot stress-strain
% this is for load_data being in lbs
stress_data = load_data*.4536*9.81/(crosssecarea/1000^2); %N/m^2=Pa
stress_data_zeroed = stress_data(:)-stress_data(1);
strain_data = new_pos_data/gaugelength;

figure;
plot1=plot(strain_data*100, stress_data_zeroed/1000, '.-');
ax=gca;
xlabel('Strain [%]');
ylabel('Stress [kPa]');
yt=ax.YAxis;
yt.FontSize=fontsize;
xt=ax.XAxis;
xt.FontSize=fontsize;
set(gca, 'linewidth', linewidth);
set(plot1, 'LineWidth', linewidth);
saveas(plot1, 'stress_strain', 'emf'); %saves it in the folder where this .m file is
saveas(plot1, 'stress_strain', 'png'); %saves it in the folder where this .m file is

save('stress_data.mat', 'strain_data', 'stress_data_zeroed', 'width', 'thickness', 'gaugelength_in', 'gaugelength', 'crosssecarea')

% If you want to close the connection
% PIdevice.CloseConnection();
%
% If you want to unload the dll and destroy the class object
% Controller.Destroy();
% clear Controller;
% clear PIdevice;

The test_parameters.m code:

```matlab
function [dl, ldot, t, epsdot_actual] = test_parameters(eps, lo, unit, epsdot)  

time = 0.0254*lo*10^3;  % [mm] convert the gauge length from inches into mm

dl = lo*eps;  % [mm] total elongation needed for sample

%assert is vector

eps = eps/100;  % not percentage of strain

% make sure that everything is in the appropriate units

lo = 0.0254*lo*10^3;  % [mm] convert the gauge length from inches into mm

dl = lo*eps;  % [mm] total elongation needed for sample

%epsdot = 10e-2;  % strain rate

ldot_desired = lo*epsdot;  % [mm/s] velocity of sample

ldot = round(ldot_desired /.0005) *.0005;

epsdot_actual = ldot/lo;

epsdot_min = 0.0025/lo;

if ldot < .0025
    warning(sprintf('Test velocity too small (must be >= 0.0025 mm/s). For given
gauge length, epsdot must be >=%.3e', epsdot_min))
end

warning(sprintf('WARNING: Actual epsdot changed from %.3e to %.3e due to test
velocity quantization.
', epsdot, epsdot_actual))

t = 2*dl/ldot;  % [s] time that the test will take to run

t = t/60;  % [min] convert time to minutes

end
```

The init_load.m code:

```matlab
function inet = init_load(varargin)

%---------------------------------------------------------
% First optional argument 'simulate_i555_instead_of_connecting_to_hdwr':
% if true, simulate i555 hardware instead of working with real hardware.

if (length(varargin) >= 1)
    simulate_i555_instead_of_connecting_to_hdwr = varargin{1};
else
    simulate_i555_instead_of_connecting_to_hdwr = false;
end

%---------------------------------------------------------
```
% Second optional argument 'instrunet_prf_settings_filename':
% name of .prf settings file to load, pass '' if you don't want to load one.

if (length(varargin) >= 2)
instrunet_prf_settings_filename = varargin{2};
else
    instrunet_prf_settings_filename = '';
end

% Initialize instrunet and get one object that describes it called 'inet'. This is like running instrunet world software. You can only have one of these running at a time, and you cannot run instrunet world application software outside matlab while this object is alive.
% You need to dispose of it after you use it or stop running matlab and that will dispose of it. To dispose of it, call 'Delete/inet'.

inet = instruNet_Class( ...
instrunet_prf_settings_filename, ... % name of .prf settings file to load, pass '' if you don't want to load one
    simulate_i555_instead_of_connecting_to_hdwr, ... % if true, simulate i555 hardware instead of working with real hardware
        false, ... % show_alert_before_loading_settings_file, set to true if you want to show an alert before loading prf file
        true, ... % showAlertIfPrfIsNotFound, set to true if you want to show an alert if prf is not found
        false, ... % turn_off_all_alert_dialogs_when_initialize, set to true if you don't want any alerts during initialization
        false, ... % showNetworkPageAfterLoadDriver, set to true if you want to show instrunet world NETWORK page after loading
        false); % showRecordPageAfterLoadDriver, set to true if you want to show instrunet world RECORD page after loading

% Get the quantity of differential analog input channel connected to MATLAB.

getNumDiVinChannels = inet.ain_di.qty;

% Useful variables stored inside instruNet_Class object:

% inet.system.initializedSuccessfully - true if instruNet initialized ok
% inet.system.weAreSimulating_i555 - true if simulating i555
% inet.system.loadedPrfOk - true if loaded .prf settings file without error
% inet.system.triedToLoadedPrfFile - true if tried to loaded .prf settings file
% inet.system.prf_fileandpath_name - name of loaded .prf settings file, if one was loaded
% inet.ain_di.qty - number of differential analog input channels
% inet.ain_se.qty - number of single-ended analog input channels
% inet.aout.qty - number of analog output channels
% Run example code that reads/writes scalar values to/from instrunet hardware. This includes channels (e.g. analog input) and parameters (e.g. sensorType).

%instruNet_Class.ShowAlert('TORI: How to read a scalar (i.e. one value) read/write with instrunet hardware. Press OK to continue.');

if (inet.ain_diqty <= 0)
  %instruNet_Class.ShowAlert('TORI: We cannot run instrunet_scalar_io() due to not having hardware channels. Does Matlab have iNet driver open in another instance? If so, you need to close it (try Exiting Matlab and come back in).');
  iNetErr = 1;
  return;
end

%scalar reading
% Load iNet channel address (numNum , devNum , modNum , chNum) into structs 'sAin1' and 'sDio30'.
loadcell1 = Load_Channel_Address(inet, 1,1,1, 13);
% Get loadcell1's channel's value given channel address stored in loadcell1 struct
[ain_valueIn , iNetErr] = GetChannelValue(inet, loadcell1);
ain_valueIn; %this should be the load in Newtons at the point when we run that line

% try reading more data points of the load!
tic
[ain_valueIn2 , iNetErr] = GetChannelValue(inet, loadcell1);
toc
ain_valueIn2;
[ain_valueIn3 , iNetErr] = GetChannelValue(inet, loadcell1);
ain_valueIn3;
[ain_valueIn4 , iNetErr] = GetChannelValue(inet, loadcell1);
ain_valueIn4;

% digitizing
% ------------------------
% End user sets this to 'true' to open instrunet world RECORD page after we digitize,
% to see your data and check your set up.
open_record_page_to_see_data_after_we_finish_digitizing = true;
% ------------------------
% End user sets this to 'true' to plot digitized data in new window
plot_digitized_data_in_new_window = true;
% ------------------------
% End user sets this to 'true' to calculate average value of each
% digitized point for each channel.
calculate_average_value = false;

% ------------------------
% End user sets this to 'true' if you want to process complete buffers (scans) at a
% time. Alternatively, set to false to process little segments
% of data as they become available.

processOneScanAtATime = true;

% ------------------------
% End user sets this to 'true' if you want mouse down to stop digitization
allowMouseDownToStopDigitize = false;

% make sure we are not digitizing and get ready to start digitizing
Stop_iNet_Digitizing(inet);

% ------------------------
% Get 'numOfDigitizeChannels' and 'ptsPerScan' values from iNet32.dll
% and also create zero'ed ram buffer to hold digitize data in
% matlab 2x2 array: 'inet.digitize.data.value(1:ptsPerScan, 1:
numOfDigitizeChannels)'.

Digitize__Allocate_2x2_Data_Matrix(inet);
end

The get_load.m code:

function [temp_val, load_val] = get_load(inet)
    load_vals = [NaN, NaN];

    % Collect Digitized Data
    %
    % It is ABSOLUTELY NECESSARY to let your computer's processor service
    % instruNet digitizing buffers periodically 10 to .3 times per second)
    % while digitizing. This is done by continuously calling
    % DigitizeListOfChannels() or Service_All_iNet_Digitize_Buffers() in a loop.
    allowMouseDownToStopDigitize = false;
    processOneScanAtATime = true;
    check_for_controllerNotRespondingTimeout = true;
    [ iNetErr, ...
      last_scanNum_base1_tsfred_inFull_oneCtlr_int64, ...
      we_stopped_digitizing_machine_or_it_finished_on_its_own, ...
      we_started_digitizing_machine_since_it_was_not_on, ...
      totalNumOfPtsAccessed_AllInputChannels_ThisRoutine_int64, ...
      endUser_sampleRate_PointsPerSecondPerChannel_actual_double, ...
      total_time_for_entire_digitization_Secs_actual_double, ...]
total_time_for_each_scan_Secs_actual, ...
user_pressed_mouse_button_to_stop_early ...

] = DigitizeListOfChannels(inet, ...
allowMouseDownToStopDigitize, ...
processOneScanAtATime, ...
check_for_controllerNotRespondingTimeout);

if (iNetErr ~= 0)
  return;
end

% Update these timing parameters (from internal inet32.dll memory):
% endUser_sampleRate
if (1) %weAreDoingOurFirstAccessOfData == true)
  [iNetErr] = Digitize__UpdateTimingParameters(inet);
end

% scan thru all channels enabled for high speed digitize
for digiChNum_base1 = 1 : inet.digitize.p.numOfChannels

  % READ DIGITIZED DATA
  %
  % find out how much data is in digitize channel # 'digiChNum_base1'
  % data buffer, and move it into matlab 2x2 data array
  % 'inet.digitize.data.value(1:ptsPerScan, 1:numOfDigitizeChannels').
  %
  %  first_PointOfSegment_Indexbase1 = pointToPullindex_base1
  %  last_PointOfSegment_Indexbase1 = pointToPullindex_base1 +
  numPointsToPull - 1
  %
  %  first_PointOfSegment = inet.digitize.data.value(
  %  first_PointOfSegment_Indexbase1, digiChNum_base1)
  %  last_PointOfSegment = inet.digitize.data.value(
  %  last_PointOfSegment_Indexbase1, digiChNum_base1)
  %
  %  data segment = inet.digitize.data.value(
  %  first_PointOfSegment_Indexbase1 : last_PointOfSegment_Indexbase1, 
  %  digiChNum_base1)

  [ iNetErr, ...
    src_pointToPullindex_base1, ... % (1... numPtsPerScan)
    src_numPointsToPull, ... % # of points to to pull
    out of the buffer
    src_scanNumberToPull_base1 ... % scan number {1...
    numScans} of the scan that we are currently pulling (base 1)
  ] = GetDigitizedSegment(inet, digiChNum_base1);

  if (iNetErr ~= 0)
    return;
  end
% if we have new data from this channel (i.e. number of new points
% is specified by variable 'src_numPointsToPull')...
if (src_numPointsToPull > 0)
    % Calculate the total # of pts that has been digitized since we
    % started digitizing
    totalPtsPulledSoFar(digiChNum_base1) = totalPtsPulledSoFar(
        digiChNum_base1) + src_numPointsToPull;
    % Calculate where segment is w.r.t. our ram buffer
    % data segment = inet.digitize.data.value(
        first_PointOfSegment_Indexbase1 : last_PointOfSegment_Indexbase1,
        digiChNum_base1)
    first_PointOfSegment_BufferIndexBase1 = src_pointToPullIndex_base1;
    last_PointOfSegment_BufferIndexBase1 = src_pointToPullIndex_base1 +
        src_numPointsToPull - 1;
    % Calculate the time of the first and last points of the segment,
    % units Seconds, relative to when we started digitizing.
    first_PointOfSegment_Secs = ( double(
        first_PointOfSegment_BufferIndexBase1-1) * inet.digitize.p.
        endUser_samplePeriodSecs ) ...
        + ( double(src_scanNumberToPull_base1
            -1) * inet.digitize.p.secsPerScan );
    last_PointOfSegment_Secs = ( double(first_PointOfSegment_Secs)
        ) ...
        + ( double(src_numPointsToPull -1) * inet.digitize.p.endUser_samplePeriodSecs);
    % true if segment includes the last point in the ram buffer
    segmentIncludesLastPointInRamBuffer = ( 
        last_PointOfSegment_BufferIndexBase1 == inet.digitize.p.ptsPerScan);
    % true if segment includes the last point in the ram buffer
    segmentIncludesLastPointInEntireDigitization = 
        segmentIncludesLastPointInRamBuffer && (src_scanNumberToPull_base1 == inet.
        digitize.p.noOfScans);
    load_vals(digiChNum_base1) = inet.digitize.data.value(
        first_PointOfSegment_BufferIndexBase1 : last_PointOfSegment_BufferIndexBase1,
        digiChNum_base1);
end
end

The PI_move.m code:

function PI_move(startPos, stopPos, step, steps, PIdevice, axis, inet)
global pos_data;
global temp_data;
global load_data;
global time_data;
global datetime_starttest;
fid = fopen('log.txt', 'a+');
finishup = onCleanup(@() myCleanUpFun(PIdevice, axis, fid));
logging = true;

%%% this following code is in the move function
% assume that referencing stuff has made it so that when this function is
called, the PIdevice is located at startPos

pos_data(1) = PIdevice.qPOS(axis);
start = tic;
time_data(1) = toc(start);
[temp_data(1), load_data(1)] = get_load(inet);
datetime_starttest = datetime;
waiting = true;
for i = 2:length(steps)
    % Move the device
    step_size = steps(i) - steps(i-1);
    PIdevice.MOV(axis, steps(i) + step_size/2);
    % Record current data
    pos_data(i) = PIdevice.qPOS(axis);
    [temp_data(i), load_data(i)] = get_load(inet);
    time_data(i) = toc(start);
    v = [time_data(i), pos_data(i), load_data(i), temp_data(i)]
    if logging
        try
            fprintf(fid, '%f,%f,%f,%f\n', v);
        catch
            logging = false;
            disp('logging failed')
        end
    end
    if ~waiting
        disp('logging took too long')
    end
    % Wait until it's moved to the new location
    step_sign = sign(step_size);
    waiting = false;
    while(step_sign*PIdevice.qPOS(axis) < step_sign*steps(i))
        waiting = true;
        continue;
    end
end

function myCleanUpFun(PIdevice, axis, fid)
    PIdvce.MOV(axis, PIdvce.qPOS(axis));
global temp_data
global time_data
global load_data
global datetime_starttest
save('extension_data.mat','time_data','pos_data','temp_data','load_data','
datetime_starttest')

try
close(fid)
catch
end
end