Dynamic Deformables:
Implementation and Production Practicalities

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Abstract

Simulating dynamic deformation has been an integral component of Pixar’s storytelling since Boo’s shirt in *Monsters, Inc.* (2001). Recently, several key transformations have been applied to Pixar’s core simulator *Fizt* that improve its speed, robustness, and generality. Starting with *Coco* (2017), improved collision detection and response were incorporated into the cloth solver, then with *Cars 3* (2017) 3D solids were introduced, and in *Onward* (2020) clothing is allowed to interact with a character’s body with two-way coupling.

The 3D solids are based on a fast, compact, and powerful new formulation that we have published over the last few years at SIGGRAPH. Under this formulation, the construction and eigendecomposition of the force gradient, long considered the most onerous part of the implementation, becomes fast and simple. We provide a detailed, self-contained, and unified treatment here that is not available in the technical papers.

This new formulation is only a starting point for creating a simulator that is up challenges of a production environment. One challenge is performance: we discuss our current best practices for accelerating system assembly and solver performance. Another challenge that requires considerable attention is robust collision detection and response. Much has been written about collision detection approaches such as proximity-queries, continuous collisions and global intersection analysis. We discuss our strategies for using these techniques, which provides us with valuable information that is needed to handle challenging scenarios.
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Chapter 1

Introduction

1.1 The Goals of This Course

Fizt\textsuperscript{1} has been the core simulator at Pixar ever since *Monsters, Inc.* in 2001. While many of the basics of the battle-tested Baraff and Witkin (1998) model that it is based on have remained the same over the years, a variety of key improvements and refinements have also been introduced. Many of these improvements have been spread across technical papers and talks presented the last two years at SIGGRAPH:

- **Better Collisions and Faster Cloth for Pixar’s Coco**, David Eberle (Talk presented at SIGGRAPH 2018)
- **Robust Skin Simulation in Incredibles 2**, Ryan Kautzman, Gordon Cameron, Theodore Kim (Talk presented at SIGGRAPH 2018)

Together, these improvements constitute the next major version of the simulator, which is referred to internally as Fizt\textsuperscript{2}. One of the goals of this course is to give a unified overview of these developments in a way that no single paper or talk has been able to before.

\textsuperscript{1}Pronounced “fizz” as in soda and “tea” as in Earl Grey, hot.
In addition to these, we have also include a new introduction to the fundamentals of deformation mechanics (Chapter 2), based on a university course taught by one of the authors over the last ten years. This treatment specifically focuses on introducing the specific tensor-based formulation that was used to obtain the results in the aforementioned technical papers. Matlab and C++ implementations of several subtle pieces of code have been inlined as well, such as the $\frac{\partial F}{\partial x}$ matrix for tetrahedra (Fig. E.1) and shape derivative matrices (Fig. D.3) for hexahedra.

1.2 Author Biographies

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David Eberle is the Simulation Tools R&D Lead at Pixar Animation Studios. He has worked on numerous simulators used for animation and visual effects at Disney, Havok, PDI/Dreamworks and Pixar. He holds a B.S. in Mathematics from the University of South Carolina and an M.S. in Mathematics from Texas A&M University. His film credits include Reign of Fire, Shrek 2, Shrek 3, Madagascar, Megamind, Inside Out, Finding Dory, Cars 3, Coco and Onward.

1.3 Acknowledgements

These notes are an ongoing draft, so the authors would like to thank the following people for finding errors and omissions in the text: J.C. Chang, Towaki Takikawa, Peng Yu, and especially Shinjiro Sueda. If you find other errors (there are sure to be many), feel free to reach out to the authors.
Chapter 2

Deformation Fundamentals

2.1 Do I Need to Read This Chapter?

If you are brand-new to deformation simulation, or are coming back to it after some time away, the answer is: yes.

There are several existing introductions to deformation geared towards computer graphics, including the classic Witkin and Baraff (1997) as well as the more modern Sifakis and Barbic (2012) or Bargteil and Shinar (2018). If you are a practitioner that is already familiar with those materials, you can probably skip most of this chapter. The one caveat is that many of the fast, compact structures we show later on will depend on the notation for higher order tensors from Golub and Van Loan (2013). This does not show up that often in computer graphics, so you may still want to read Chapter 3.

Figure 2.1.: The behavior of a hyperelastic bunny. Even when it is crushed by a 100 lb. weight, it recovers its original shape afterwards.
2.2 What Kind of Squashing Are We Talking About?

We will be looking at a specific form of solid deformation known as hyperelastic deformation. Say we take a bunny and squash it with a 100 lb. weight (Fig. 2.1). When we remove the weight, it will sproing back to its exact original shape. This is the essence of hyperelasticity: when all external forces are removed, a hyperelastic object returns to its original shape. It does not matter if the bunny was stretched to a million times its original length, or crushed down into a pancake. Once the forces are removed, it bounces back to its original bunny shape.\(^1\)

The original shape is thus considered quite special, as it is what the object “remembers” and always “tries” to return to, and is assigned a special name: the rest shape. In fact, when we squash a bunny with a 100 lb. weight, it is taking on the best possible or closest possible shape to the rest shape that it can muster under the current conditions.

But what do we mean by best or closest? Best compared to what, or closest with respect to what? This is really two questions.

1. How squashed am I? Alternately, how far am I from my original shape? Or: how badly deformed am I compared to my original shape? Most concretely: can I compute a quantitative deformation score that tells me exactly how stretched or squashed I am, relative to my original shape?

2. How much should I push back? Once I know how exactly how deformed I am, what should I do with this knowledge? Is being really deformed considered extra-bad, so I should push extra-hard to return to my original shape?

Let’s look at these questions in turn.

\[ \text{Figure 2.2.: On the left is the original, rest-shape triangle. On the right we (slightly) deform the triangle by moving } x_1 \text{ downwards.} \]

2.3 How Squashed Am I?

First, let’s look at an obvious way to measure deformation that will turn out to be wrong. From there, we can think about what went wrong and come up with something better.

\(^1\)Clearly, this does not always happen in reality. If you stretch a rubber bunny too much, the fibers inside will slightly tear. When you let go, it will “remember” its rowdy treatment, and instead return to a slightly-stretched-out-looking-bunny shape. This is called plastic deformation, and can be layered on top of hyperelastic deformation, so we put it aside for the moment.
2.3.1 Measuring The Wrong Way

Let’s look at a triangle (Fig. 2.2). We will denote its vertices in the rest shape using an overbar as \( \bar{x}_0, \bar{x}_1, \bar{x}_2 \). After the triangle has been squashed away from its original shape, we denote its deformed shape\(^2\) as \( x_0, x_1, x_2 \). One obvious way to compute a score for how deformed we are is then:

\[
\Psi_{\text{wrong}} = \sum_{i=0}^{2} \| \bar{x}_i - x_i \|_2.
\] (2.1)

Here, we use \( \| \cdot \|_2 \) to denote the 2-norm. Don’t fret if you haven’t seen this notation before; we have a description of it in Appendix C.1.1. It is a generalized shorthand for denoting “distance”. We will also interchangeably refer to scoring functions as energy functions. For our purposes, scoring and energy functions have the exact same meaning.

The scoring function \( \Psi_{\text{wrong}} \) certainly looks reasonable. As you squash \( x_1 \) downwards more, or stretch it upwards more, the score (energy) definitely increases. We definitely want the score function to equal zero when no deformation is occurring, and some big number when lots of deformation is occurring. If that’s all that is required, \( \Psi_{\text{wrong}} \) certainly seems to get the job done.

Figure 2.3.: We translate the original triangle (in black) to the upper left (in red). Does this triangle count as “deformed”?

Or does it? The problem is that a bunch of things that we would not intuitively consider deformation also gets lumped in with the score. In Fig. 2.3, we have translated the triangle a small distance to the upper right. If we use \( \Psi_{\text{wrong}} \) on this new triangle, then it will show up as a non-zero score.

However, by any reasonable definition, this triangle is not deformed. Instead, it has been moved, or translated. Fig. 2.4 shows another problem case. If we rotate the triangle, then again, \( \Psi_{\text{wrong}} \) will return a non-zero score, mistakenly regarding this triangle as “deformed”. However, by any reasonable definition, the triangle has been spun or rotated. It has not been deformed.

\(^2\)Other texts may use the terms material and world coordinates, so if you ever come across these, just remember that rest = material and deformed = world.
What we want is a scoring function $\Psi$ that is translation and rotation invariant. If a triangle has been merely translated or rotated, its deformation score should show up as zero. Let’s see if we can compute a score that will accomplish this.

2.3.2 Removing the Translation (Easy)

One way to encode the transformations that a triangle can undergo to use an affine map:

$$\phi(x) = Fx + t$$

(2.2)

Here, $t \in \mathbb{R}^2$ and $F \in \mathbb{R}^{2 \times 2}$ in 2D, and $t \in \mathbb{R}^3$ and $F \in \mathbb{R}^{3 \times 3}$ in 3D (see the symbol table in Appendix A). There are only a few things that an affine transformation can encode:

- Rotation
- Scaling
- Reflection
- Translation.\(^3\)

We saw that rotation and translation kept showing up and polluting our deformation score, even though we don’t intuitively consider those as “deformations”. However, scaling matches our notion of deformation quite nicely. Can we just isolate and measure that?

As a first step, let’s see if we can carve away the translation, which is relatively easy. The translation lives entirely in $t$, so if we perform some manipulation that cuts away that

\(^3\)You may ask: what about shearing? Wasn’t that in linear algebra class too? That can be written in terms of rotations and scalings, so we can omit it here.
component, we are done. The canonical way of doing this is to take the derivative with respect to the rest shape:

\[
\frac{\partial \phi(\bar{x})}{\partial \bar{x}} = \frac{\partial}{\partial \bar{x}} (F\bar{x} + t) = F.
\] (2.3)

Since we are taking the derivative (gradient) of the affine map (deformation), the matrix \( F \) that pops out is called the deformation gradient. This matrix now contains rotation, scaling, and reflection, but \textit{not translation}. If we base some sort of deformation score on \( F \), there is no way that translation can now pollute the result.

\subsection*{2.3.3 Deformation Gradient: Really Important}

\textbf{It is hard to overstate how important the deformation gradient is.} Almost everything we do for the rest of this chapter will deal with \( F \). The variable has an unwieldy name, and everybody (Bonet and Wood (2008); Bower (2009); Marsden and Hughes (1994); Belytschko et al. (2013)) uses \( F \) to denote it, even though neither “deformation” nor “gradient” starts with an ‘f’. Ideally the variable would have a snappy name and an intuitive symbol, but that is not the world we live in.\footnote{In geometry processing, it is sometimes denoted with \( J \), presumably because it is the deformation Jacobian. Better, but still not great.} We are stuck with \( F \), so just remember that \( F \) is important.

You may be asking yourself: if \( F \) is so important, are you going to tell me how to compute it? Absolutely. It is a little involved, and takes a bit of space, so it has its own section in Appendix D. Taking for granted that we know \textit{how} to compute this quantity, let’s examine \textit{what it means}.

\subsection*{2.3.4 Removing the Rotation (Not So Easy)}

We now have \( F \), where translation has been mercifully subtracted off, but scaling, rotation and reflection remain entangled. How can we remove the rotation? This will turn out to be more difficult than removing the translation, and there will actually be more than one answer.

Let’s try to build a simple score function out of \( F \). The simplest way to boil a matrix down to a scalar score is to take its \textit{ squared Frobenius norm}, denoted \( \| \cdot \|_F \). As with the 2-norm in §2.3.1, don’t worry if you’re not intimately familiar with this norm. We document it in detail in Appendix C.1.2, but since this is the first time it is appearing, we will write it out explicitly,

\[
\Psi_{\text{Dirichlet}} = \| F \|^2_F = \sum_{i=0}^{2} \sum_{j=0}^{2} f_{ij}^2.
\] (2.4)
where we have numbered the entries of $F$ thusly:

$$F = \begin{bmatrix} f_{00} & f_{01} & f_{02} \\ f_{10} & f_{11} & f_{12} \\ f_{20} & f_{21} & f_{22} \end{bmatrix}.$$  

(2.5)

This score function is also known as the Dirichlet energy, and it does not get the job done. Remember, we want the deformation score to return zero when there has been zero deformation. However, the Dirichlet energy will return zero when all the entries of $F$ are zero, i.e. $F = 0$.

If we plug $F = 0$ into our original deformation function $\phi(\bar{x}) = F\bar{x} + t$, it corresponds to the case where all the points $\bar{x}$ are crushed into a zero-volume black hole centered at $t$. As a scoring function, the Dirichlet energy might be a great black hole detector, but it’s a crummy deformation measure.

### 2.3.4.1 A First (Wrong) Attempt

The Dirichlet energy returns a zero when $F = 0$. What does an energy (score) that returns zero under zero deformation even look like? Well, if $F = I$, then certainly no deformation is occurring. In that case, $\phi(\bar{x}) = F\bar{x} + t = I\bar{x} + t = \bar{x} + t$, which is purely a translation.

Can we design a function that returns zero when $F = I$? Well, if $F = I$, then $\|F\|^2_F = 3$. So, let’s try the scoring function:

$$\Psi_{\text{Neo-Hookean}} = \|F\|^2_F - 3.$$  

(2.6)

This is not a good idea. While it is true that when $F = I$, the score will equal zero, we also want zero to be the smallest score possible. Otherwise, the score becomes difficult to interpret. A zero score means zero deformation, a score greater than zero means some deformation, but what does a score less than zero mean? Less than zero deformation is happening?

In the case of $\Psi_{\text{Neo-Hookean}}$, if $F = 0$, then $\Psi_{\text{Neo-Hookean}} = -3$. So, it’s a pretty bad deformation measure. Things get worse if

$$F = \begin{bmatrix} 0 & 0 & 0 \\ 0 & \sqrt{3}/2 & 0 \\ 0 & 0 & \sqrt{3}/2 \end{bmatrix}.$$  

(2.7)

This refers to a Wile-E-Coyote-style pancake state, which is definitely not a zero deformation state. And yet, the $\Psi_{\text{Neo-Hookean}}$ score returns a zero. Apparently there are many sneaky configurations that can fool this scoring function into thinking that no deformation has occurred. We could try to apply a squaring Band-Aid, e.g. $\Psi_{\text{Neo-Hookean,Band-Aid}} = \ldots$
((∥F∥_F^2 - 3)^2, but this would only address the negative scores problem. The scoring function would still return zero for the same sneaky deformed states.\(^5\)

However, for some reason it has the fancy name “Neo-Hookean”, so it must be important. It has been studied extensively in mechanical engineering and material science for the last 80 years (Mooney (1940); Rivlin (1948)), because we will see later that if you add a few important components, it becomes a very interesting deformation measure. But, it’s not very good in its current form, so let’s put it aside for now.

### 2.3.4.2 A Second (Still Wrong) Attempt

Next, let’s try something capricious and stupid-looking (C&SL), which I promise will eventually lead somewhere. Let’s just subtract \( F \) and \( I \) and take its norm. Then, we are at least guaranteed that when \( F = I \), our scoring function will equal zero:

\[
\Psi_{C&SL} = ∥F - I∥_F^2. \tag{2.8}
\]

Additionally, since the \( I \) is inside the norm, everything gets squared, so there is no possibility of our function producing meaningless negative scores. We can use identity B.16 from Appendix B to expand this out to:

\[
\Psi_{C&SL} = ∥F - I∥_F^2 = ∥F∥_F^2 + ∥I∥_F^2 - 2 \text{tr} F = ∥F∥_F^2 + 3 - 2 \text{tr} F.
\]

I have assumed that we’re looking at the 3D case, which is why \( ∥I∥_F^2 = 3 \). Looking at the expansion, the score still isn’t great. The first term is the black-hole favoring Dirichlet energy, which doesn’t do us any favors. Then, we have the constant of 3, which looks suspiciously similar to our not-great Neo-Hookean energy, followed by a new \( \text{tr} F \) term. If you haven’t seen a matrix trace in a while, there is a refresher in Appendix C.2. Since this is its first appearance here, we will write it out explicitly:

\[
\text{tr} F = \sum_{i=0}^{2} f_{ii} = f_{00} + f_{11} + f_{22}. \tag{2.9}
\]

This scoring function will equal zero when \( F = I \), but when \( F \neq I \), weird things happen. For example, if \( F \) happens to be some rotation matrix,

\[
F(\theta) = \begin{bmatrix}
1 & 0 & 0 \\
0 & \cos \theta & -\sin \theta \\
0 & \sin \theta & \cos \theta
\end{bmatrix}, \tag{2.10}
\]

\(^5\)Fun fact: This squaring is equivalent to the (yet-to-be-seen) volume preservation term from the St. Venant Kirchhoff model. It is still not terribly useful all on its own though.
then $\Psi_{C&SL}$ will return a non-zero score whenever $\theta$ is not zero or some multiple of $2\pi$. But, these would still clearly correspond to deformation-free states! It’s better than a black-hole detector, but not by much.

Overall, it’s weird that $\text{tr} \ F$ is shows up at all in this formula. Since $F$ can be an arbitrary, non-symmetric matrix, it is not clear that its trace is terribly meaningful outside of cancelling off $\|F\|_F^2 + 3$ in the overly-specific case of $F = I$.

### 2.3.4.3 A Third Attempt: St. Venant Kirchhoff

How can we make a version of $\Psi_{C&SL}$ that instead equals zero whenever $F$ is a pure rotation? One way to do this is to exploit a nice linear algebra identity. If $F$ is a pure rotation, then we know that it must be the case that $F^T F = I$. This follows from the fact that $F^{-1} = F^T$ when $F$ is orthonormal.

Well, why don’t we compute $F^T F$, subtract off the rotation part, i.e. the $I$ part, and see what’s left over? These leftovers probably characterize the deformation in some reasonable way. We can write this down as:

$$\Psi_{\text{StVK, stretch}} = \frac{1}{2}\|F^T F - I\|_F^2.$$ (2.11)

This is a pretty good idea! The idea is good enough that people have given it a special name: The St. Venant-Kirchhoff stretching energy, or “StVK, stretch”, for short.\(^6\) The individual components of $F^T F - I$ are popular enough that they have special names and symbols:

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>$C = F^T F$</td>
<td>The right Cauchy-Green tensor</td>
</tr>
<tr>
<td>$E = \frac{1}{2} \left( F^T F - I \right)$</td>
<td>Green’s strain</td>
</tr>
</tbody>
</table>

Using these, the energy can be written in a highly compact form:

$$\Psi_{\text{StVK, stretch}} = \|E\|_F^2.$$ (2.12)

Again, this is a decent way to measure deformation. We’re using $F$, so the translation has been subtracted off, and we are using an $F^T F$ identity to factor off rotation. If we were going to complain about one thing though, it would be that the scoring function is overly non-linear. Performing the same expansion as we did for $\Psi_{C&SL}$, we can get:

$$\Psi_{\text{StVK, stretch}} = \|F^T F - I\|_F^2 = \|F^T F\|_F^2 + \text{tr} I - 2 \text{tr}(F^T F).$$ (2.13)

The leading term, $\|F^T F\|_F^2$, is 4\(^{th}\)-order, or quartic in $F$. The $F$ gets squared by $F^T F$, and then squared again by the norm, $\| \cdot \|_F^2$, which makes it 4\(^{th}\) order overall. We will see later that this 4\(^{th}\)-order-ness adds additional computational complexity, but in a way that is not actually desirable.

\(^6\) Actually, I call it the StVK stretching energy, because it’s the first term (the stretching term) in the StVK energy. So if you look up “St. Venant-Kirchhoff”, you’ll see this term, plus another one. We’ll take a look at that other one later.
2.3.4.4 A Fourth Attempt: As-Rigid-As-Possible

We took advantage of the $F^T F = I$ identity, but are there other identities that we could have used? If we’re specifically concerned with teasing apart the rotation and non-rotation component of a matrix, the polar decomposition is helpful:

$$F = RS. \quad (2.14)$$

Here, the matrix $R$ is the rotational part of $F$, and $S$ is the non-rotational (scaling) component of $F$. This decomposition seems custom-built to solve exactly the problem we are interested in.

However, you need to be a little careful about using the polar decomposition, because its classic definition is slightly different from the version we want. Usually $R$ is only defined as a unitary matrix, not a rotation matrix, so it only needs to satisfy $R^T R = I$. This means that there could be a reflection lurking somewhere in $R$, whereas we want $R$ to be a pure rotation. In our case, if a reflection has to lurk somewhere, we would prefer that it do so in $S$. We will call this specific flavor the rotation variant of the polar decomposition. Details on how to compute this are in Appendix F.

In short: if you end up calling the polar decomposition code in some library, make sure it is computing the rotation variant, not the traditional version. If it is not, you will need to write a small wrapper for the library call in the style of Figs. F.1 and F.2.

With that out of the way, let’s assume you can compute the rotation-variant polar decomposition, and now have $R$ and $S$ in hand. What can you do with them? In StVK, we took $F^T F$ and subtracted off its rotational component, $I$. Well, now we have a different representation for the rotation, $R$, but can we do the same thing? Subtract the rotation component off of $F$, and assume that the leftovers must characterize the deformation somehow?

The most basic, gee-I-wonder-if-this-will-work way of writing this down is:

$$\Psi_{ARAP} = \|F - R\|_F^2. \quad (2.15)$$

There are a few worrying-looking things about this formulation. Usually when you have two matrices, multiplying them together is a first thing to try, because you can see what the multiplication is really doing by peeking at the SVD or eigendecomposition. But, instead of multiplying, we’re subtracting. Can we really just subtract two arbitrary-looking matrices from each other and get something that isn’t total garbage? They are related via the polar decomposition, but does that really mean we can just start subtracting these entries from each other?

Surprisingly, the answer is yes. In fact, this energy function is so indispensable that it has been given a special name in geometry processing: the As-Rigid-As-Possible (ARAP) energy (Sorkine and Alexa (2007)). We’ll do a little more analysis of this energy a little later to see how it is actually more clever than it first appears.
This energy does not have the “overly non-linear” problem of StVK. If we apply identity B.16, we get:

\[
\Psi_{\text{ARAP}} = \| \mathbf{F} - \mathbf{R} \|^2_F \\
= \| \mathbf{F} \|^2_F + \| \mathbf{R} \|^2_F - 2 \text{tr}(\mathbf{F}^T \mathbf{R}) \\
= \| \mathbf{F} \|^2_F + 3 - 2 \text{tr}(\mathbf{S}).
\] (2.16)

The energy is at most quadratic in \( \mathbf{F} \). We will see later that the introduction of \( \mathbf{R} \) and \( \mathbf{S} \) creates some computational challenges, but for the time being, it is undeniable that \( \Psi_{\text{ARAP}} \) is appealingly less non-linear than \( \Psi_{\text{StVK,stretch}} \).

In physical simulation, the ARAP energy is actually the first term in the popular co-rotational energy. This energy has been known in mechanical engineering since at least the 1980s (Rankin and Brogan (1986)), and the ARAP energy corresponds to the specific mechanical case where Poisson’s ratio is set to zero. This energy was rediscovered by several graphics researchers in the early 2000s (Müller et al. (2002); Etzmuss et al. (2003); Irving et al. (2004)), which created some confusion because they each gave it a different name, such as “stiffness warping” or “rotated linear”. However, these models are all equivalent, and nowadays everybody seems to have settled on “co-rotational” as the name for this energy.

2.3.4.5 Summary of Energy (Scoring) Functions

We have now seen a bunch of different energy functions. Let’s summarize their pluses and minuses:

<table>
<thead>
<tr>
<th>Energy</th>
<th>Pros</th>
<th>Cons</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dirichlet</td>
<td>Not many. A good black hole detector?</td>
<td>Doesn’t measure deformation effectively.</td>
</tr>
<tr>
<td>Neo-Hookean</td>
<td>Returns zero when ( \mathbf{F} = \mathbf{I} )</td>
<td>Returns zero for other configurations too, and can also return negative scores.</td>
</tr>
<tr>
<td>C&amp;SL</td>
<td>Returns zero when ( \mathbf{F} = \mathbf{I} ), and does not return negative scores.</td>
<td>Does not return zero under pure rotations.</td>
</tr>
<tr>
<td>StVK</td>
<td>Returns zero when ( \mathbf{F} ) is a pure rotation, and reasonable scores otherwise.</td>
<td>Overly non-linear (quartic).</td>
</tr>
<tr>
<td>ARAP</td>
<td>Returns zero when ( \mathbf{F} ) is a rotation, reasonable scores otherwise, and is quadratic</td>
<td>That ( \mathbf{R} ) term is going to cause trouble.</td>
</tr>
</tbody>
</table>

As I said at the top, removing the rotation from the score is not easy, and there is more than one strategy. Now that we have a bunch of options, what should we do with them?
2.4 Force Computation: How Much Should I Push Back?

Now that we can measure how deformed we are, we can return to the second question from §2.2: *How much should I push back?*

The deformation measure directly generates forces. Surprise! When we were muddling over different ways to measure deformation, we were actually trying out different elastic energy potentials. The two are completely equivalent for our purposes. The deformation measure directly dictates the force response. If you want to design a new material model, come up with a different way to measure deformation.

To get the forces $f$ on the nodes of a triangle in 2D, we stack the vertices of our triangle ($x_0$, $x_1$ and $x_2$) into a big vector,

$$
{x} = \begin{bmatrix} x_0 \\ x_1 \\ x_2 \end{bmatrix} = \begin{bmatrix} x_0 \\ y_0 \\ x_1 \\ y_1 \\ x_2 \\ y_2 \end{bmatrix} \in \mathbb{R}^6,
$$

and then take the gradient:

$$
f = -a \frac{\partial \Psi}{\partial x}. \tag{2.18}
$$

Here, $a$ is the area of the original triangle at rest. A really tiny deformed triangle the size of an ant is not going to exert the same force as a deformed triangle the size of the Empire State Building, so we should scale it.

That’s all well and good, and Eqn. 2.18 looks straightforward, except that all the energies we looked at in §2.3.4.5 were written in terms of $F$, not $x$. In fact, we made a big deal out of the fact that $F$ doesn’t have all these junky measurement-polluting properties of $x$, like translation, and that it was clearly the better primary variable. Having gone down that path, how are we supposed to take the gradient with respect to $x$ now?

2.4.1 Computing $\frac{\partial \Psi}{\partial x}$

We pushed the computation of $F$ to Appendix D, but we’re going to pull the final result of that discussion back up here. For a triangle, we pack its original $\bar{x}_i$ and deformed $x_i$ vertices into two matrices, $D_m$ and $D_s$, thusly:

$$
D_m = \begin{bmatrix} \bar{x}_1 - \bar{x}_0 \\ \bar{x}_2 - \bar{x}_0 \end{bmatrix} \quad \quad \quad D_s = \begin{bmatrix} x_1 - x_0 \\ x_2 - x_0 \end{bmatrix}.
$$

The deformation gradient is then a composition of these two:

$$
F = D_s D_m^{-1}. \tag{2.20}
$$
First the good news: we need the gradient with respect to \( x \), not \( \bar{x} \). If we instead needed \( \frac{\partial \Psi}{\partial \bar{x}} \), things would get ugly real fast, because \( \bar{x} \) appears in \( D_m \), which is inside an inverse, \( D_m^{-1} \), so we’d need to figure out the derivative of a matrix inverse.

Now the bad news: it’s still pretty ugly. Let’s take a really simple energy, \( \Psi_{\text{Dirichlet}} \) as an example:

\[
\Psi_{\text{Dirichlet}} = \| F \|_F^2.
\]  

(2.21)

Here’s what we’re going to have to do:

- Plug \( F = D \bar{s} D_m^{-1} \) into \( \Psi_{\text{Dirichlet}} \).
- Multiply everything through to get a massive one-line equation for \( \| F \|_F^2 \).
- Take the derivative of this massive equation six times, once for each entry in \( x \), and stack the results into a force vector.
- Hope you didn’t make a mistake in your derivation somewhere. Port the expressions into C++, and hope you don’t make a mistake transferring these equations from paper to code.

### 2.4.2 Don’t Do It This Way

Let’s gaze at this ugliness for just a minute longer. Just a minute though. After that I have to look away, and tell you there’s a better way. To start, we can tag each entry in \( D_m^{-1} \) thusly,

\[
D_m^{-1} = \begin{bmatrix} m_0 & m_2 \\ m_1 & m_3 \end{bmatrix}
\]  

(2.22)

then we’ll try to multiply out the matrix \( F^T F \) in anticipation of cramming everything together for \( \| F \|_F^2 \).

\[
F^T F = \begin{bmatrix} m_0 & m_1 \\ m_2 & m_3 \end{bmatrix} \begin{bmatrix} x_1 - x_0 \\ 0 \end{bmatrix} \begin{bmatrix} 0 \\ x_2 - x_0 \end{bmatrix} + \begin{bmatrix} m_1 & m_0 \\ m_3 & m_2 \end{bmatrix} \begin{bmatrix} x_1 - x_0 \\ x_2 - x_0 \end{bmatrix} \begin{bmatrix} x_1 - x_0 \\ x_2 - x_0 \end{bmatrix} = \begin{bmatrix} d_0 & d_1 \\ d_2 & d_3 \end{bmatrix}.
\]  

(2.23)

Just to clean things up, let’s introduce more temporary variables:

\[
\begin{bmatrix} (x_1 - x_0)^T (x_1 - x_0) & (x_2 - x_0)^T (x_1 - x_0) \\ (x_2 - x_0)^T (x_1 - x_0) & (x_2 - x_0)^T (x_2 - x_0) \end{bmatrix} = \begin{bmatrix} d_0 & d_1 \\ d_1 & d_2 \end{bmatrix}.
\]  

(Note it’s symmetric)
Now let’s forge ahead further,

\[
F^T F = \begin{bmatrix}
  m_0 & m_1 & d_0 & d_1 \\
  m_2 & m_3 & d_1 & d_2 \\
  m_0 & m_2 & d_0 & d_1 \\
  m_1 & m_3 & d_1 & d_2
\end{bmatrix}
\begin{bmatrix}
  m_0 \\
  m_1 \\
  m_2 \\
  m_3
\end{bmatrix}
\]

and then finally smash everything together with the Frobenius norm:

\[
\|F\|^2_F = (m_0^2 d_0 + 2m_0 m_1 d_1 + m_1^2 d_2)^2 + (m_0 m_2 d_0 + (m_0 m_3 + m_1 m_2) d_1 + m_1 m_3 d_2)^2 + (m_2^2 d_0 + 2m_2 m_3 d_1 + m_3^2 d_2)^2.
\]

Okay, I have to stop now. Cripes, it hurts to even look at this. The expression is big enough that it barely fits on two lines. I probably made a mistake in there somewhere; good luck finding it. Next we’ll have to substitute back in \(d_0 = (x_1 - x_0)^T (x_1 - x_0)\) and from there, \(x_0 = [x_0 \ y_0]^T\), as well as \(d_1, d_2, x_1\) and \(x_2\), and then take multiple derivatives of the resulting hideousness. Also, things get worse in 3D, and the Dirichlet energy isn’t even that complicated an energy! Things sink to even deeper depths of hideousness for more complex material models.

You could use a computer algebra system like Mathematica or Matlab’s Symbolic Toolkit to handle all this. That would just offload the ugliness generation to the computer, and the final result would still be ugly. Surely there is a better way.

### 2.4.3 What Now?

There is a better way, which we will describe in the next chapter. When we do, you will see that the forces are now cleaner to compute, but the force gradients (a.k.a the energy Hessians) are still ugly.

You may be tempted to take the coward’s way out here. All of this ugliness started because I insisted that we use \(F\) to compute our deformation measure instead of \(x\). If we had just stuck to \(x\), we wouldn’t be in this mess, because computing \(\frac{\partial \Psi}{\partial x}\) when \(\Psi\) is written purely in terms of \(x\) is much easier. Just look at this gorgeous-looking spring-mass energy:

\[
\Psi = \frac{\mu}{2} ||x - x_0||^2\ (2.24)
\]

\[
\frac{\partial \Psi}{\partial x} = \mu(x - x_0).\ (2.25)
\]

SO EASY. Don’t be seduced though. If allow ourselves be sucked down into this Charybdis, we’ll be limiting ourselves to the world of spring-mass models, and the richer universes of squishing and squashing volumetric responses will be inaccessible. Let’s not do that; let’s see how far we can take this \(F\) thing.
Chapter 3

Computing Forces the Tensor Way

If you’re just joining us after skipping Chapter 1, the following is our current predicament. We are trying to compute forces, \( f = a \frac{\partial \Psi}{\partial x} \), but it turns out that directly taking the \( x \) derivative when your energy \( \Psi \) is based on the deformation gradient \( F \) is extremely ugly. It’s ugly enough that we are starting to question whether using an \( F \)-based energy is even a good idea at all. Now you’re all caught up, so on with the show.

We will clean things up using an application of the chain rule that separates \( \frac{\partial \Psi}{\partial x} \) into two components:

\[
\frac{\partial \Psi}{\partial x} = \frac{\partial \Psi}{\partial F} \frac{\partial F}{\partial x}.
\]

Each piece, \( \frac{\partial \Psi}{\partial F} \) and \( \frac{\partial F}{\partial x} \), is relatively clean in isolation. The ugliness we saw in the previous chapter appears if you try to bite off more than you can chew and handle both pieces simultaneously. However, if you haven’t worked with higher-order tensors before\(^1\), you’ll run into trouble the moment you start trying to work out these terms. The whole thing will look like a dirty calculus trick that some professor played on you where he\(^2\) left all the difficult details out. Let’s look at the details now.

3.1 Thinking About 3\(^{rd}\)-Order Tensors

The first term in Eqn. 3.1, the \( \frac{\partial \Psi}{\partial F} \) term, is relatively easy. Since \( \Psi \in \mathbb{R} \) and \( F \in \mathbb{R}^{3 \times 3} \) the derivative is just a matrix:

\[
\frac{\partial \Psi}{\partial F} = \begin{bmatrix}
\frac{\partial \Psi}{\partial f_{00}} & \frac{\partial \Psi}{\partial f_{01}} & \frac{\partial \Psi}{\partial f_{02}} \\
\frac{\partial \Psi}{\partial f_{10}} & \frac{\partial \Psi}{\partial f_{11}} & \frac{\partial \Psi}{\partial f_{12}} \\
\frac{\partial \Psi}{\partial f_{20}} & \frac{\partial \Psi}{\partial f_{21}} & \frac{\partial \Psi}{\partial f_{22}}
\end{bmatrix} \in \mathbb{R}^{3 \times 3}.
\]

\(^1\) Running a few TensorFlow Python scripts definitely doesn’t count. If you have actually worked with tensors before, feel free to skip ahead to §3.4.

\(^2\) That would be me.
The $f_{ij}$ scalars are the entries of $F$, as described in Appendix A. The result is clearly a $3 \times 3$ matrix, also known as a 2nd-order tensor.

The confusion begins when we try to compute $\frac{\partial F}{\partial x}$. Since $F \in \mathbb{R}^{3 \times 3}$ is a matrix and $x \in \mathbb{R}^{12}$ is a vector, what form should the derivative take? Let’s look at $x_0$, the first entry of $x$. We can take the derivative of $F$ with respect to that:

$$\frac{\partial F}{\partial x_0} = \begin{bmatrix} \frac{\partial f_{00}}{\partial x_0} & \frac{\partial f_{10}}{\partial x_0} & \frac{\partial f_{20}}{\partial x_0} \\ \frac{\partial f_{01}}{\partial x_0} & \frac{\partial f_{11}}{\partial x_0} & \frac{\partial f_{21}}{\partial x_0} \\ \frac{\partial f_{02}}{\partial x_0} & \frac{\partial f_{12}}{\partial x_0} & \frac{\partial f_{22}}{\partial x_0} \end{bmatrix} \in \mathbb{R}^{3 \times 3}. \quad (3.3)$$

Already, the result is a matrix. What are we supposed to do with the next derivative, $\frac{\partial F}{\partial x_1}$, and the one after that, and the one after that? Eventually I’ll have a pile of 12 matrices, but what am I supposed to do with them? Is there some standard way of arranging these matrices together?

The answer is no. The problem is that we now have a 3rd-order tensor, which usually doesn’t show up in an introductory linear algebra texts. If you look in texts on higher-order tensor manipulation (e.g. Golub and Van Loan (2013); Simmonds (2012); Kolda and Bader (2009)), they describe a variety of ways of arranging this pile of matrices, some of which have important-sounding names like Einstein notation. Surely that’s the one? If it’s good enough for Albert Einstein, it’s good enough for us? In fact, it’s not; we should use the right tool for the right situation, and we’re not looking at relativity right now.

Figure 3.1.: We could take each matrix along the top and stack them all into a cube. For the full $\frac{\partial F}{\partial x}$ there are 12 slices, but hopefully you get the picture with just 4. We’re not going to do it this way though.

---

3Some of your colleagues may dispute my claim and say that there is definitely a standard notation for tensors that everybody who’s anybody uses. I would recommend being suspicious of any colleague who tries to make such an intimidation-based argument.
The dimension of the derivative is \( \frac{\partial F}{\partial x} \in \mathbb{R}^{3 \times 3 \times 12} \), where the third dimension is the big giveaway that we’re looking at a 3rd-order tensor. Many people think of this as a cube of matrices. We can take our pile of 12 matrices, and stack them like a deck of cards into a cube (Fig. 3.1). Just to sow further confusion, if you look at my paper, Kim and James (2012), you’ll see that I decided to use this mental representation in that paper too.\(^4\)

Figure 3.2.: We prefer to think of 3rd-order tensors as a vector of matrices. It’s like a long cutting board on a kitchen counter, with the matrices stacked along it, like individually-wrapped slices of American cheese. There should be 12 matrices in total, but I only show 4 because if I show all 12 they’d be so small you couldn’t read the writing on them.

We’re not going to take that view here. Instead, we’re going to think of 3rd-order tensors as a vector of matrices. Think of the \( \frac{\partial F}{\partial x} \) tensor like a big wooden cutting board on your kitchen counter. Each matrix \( \frac{\partial F}{\partial x_i} \) is then placed in order, along a line, down the board (Fig. 3.2), like slices of individually-wrapped American cheese. When we write out a 3rd-order tensor explicitly, we’ll use the square brackets to imitate our mental picture of a cutting-board-with-cheese. For example:

\[
\begin{bmatrix}
a & c \\
b & d \\
\vdots & \vdots \\
i & k \\
j & l \\
\end{bmatrix}
= 
\begin{bmatrix}
A \\
B \\
C \\
\end{bmatrix}.
\]

(3.4)

On the right hand side, we wrapped each matrix in a [ ] as well. This is to distinguish the 3rd-order tensor from the more familiar 2nd-order block matrix.

\(^4\)Cubes are a good way to think about things when you’re doing model reduction, but that’s not what we’re doing right now.
3.2 Multiplication With 3rd-Order Tensors

Once we have a mental picture of the 3rd-order tensor in place, how should we think of stuff like multiplication? This is pretty important, since once of our initial claims was that the chaining \( \frac{\partial \Psi}{\partial \mathbf{x}} = \frac{\partial \Psi}{\partial \mathbf{F}} \frac{\partial \mathbf{F}}{\partial \mathbf{x}} \) would make things easier, and so far it sure doesn’t look easier. What’s the product of \( \frac{\partial \Psi}{\partial \mathbf{F}} \), a matrix, and \( \frac{\partial \mathbf{F}}{\partial \mathbf{x}} \), a cutting board?

This is the part where you get annoyed with the professor for gliding past an important detail. If we’re writing down rock-solid, 100%-corresponds to the actual computation you’ll be doing we should really write:

\[
\frac{\partial \Psi}{\partial \mathbf{x}} = \frac{\partial \mathbf{F}}{\partial \mathbf{x}} : \frac{\partial \Psi}{\partial \mathbf{F}},
\]

where the ‘:’ is a double-contraction. If you have not seen double-contractions before, we review what it means to contract two matrices together in Appendix C.3. The key takeaway from that section is:

\[
\mathbf{A} : \mathbf{B} = \begin{bmatrix}
    a_0 & a_2 \\
    a_1 & a_3 \\
\end{bmatrix}
\begin{bmatrix}
    b_0 & b_2 \\
    b_1 & b_3 \\
\end{bmatrix} = a_0b_0 + a_1b_1 + a_2b_2 + a_3b_3.
\]

However, \( \frac{\partial \mathbf{F}}{\partial \mathbf{x}} \) is a 3rd-order tensor, not a matrix. What does a 3rd vs. 2nd double-contraction mean? We define it as:

\[
\mathbf{A} : \mathbf{B} = \begin{bmatrix}
    a_0 & a_2 & a_4 & a_6 & a_8 \\
    a_1 & a_3 & a_5 & a_7 & a_9 \\
\end{bmatrix}
\begin{bmatrix}
    b_0 & b_2 \\
    b_1 & b_3 \\
\end{bmatrix} = \begin{bmatrix}
    a_0b_0 + a_1b_1 + a_2b_2 + a_3b_3 \\
    a_4b_0 + a_5b_1 + a_6b_2 + a_7b_3 \\
    a_8b_0 + a_9b_1 + a_{10}b_2 + a_{11}b_3 \\
\end{bmatrix}.
\]

We have now used the blackboard font \( \mathbf{A} \) to denote a higher-order tensor for the first time. We will use these to denote tensors that are 3rd-order and above; the order should be (I hope) evident from the context.

To form another mental picture, \( \frac{\partial \mathbf{F}}{\partial \mathbf{x}} \) is a cheese slice off to the side that has the exact same size as those on our cutting board, \( \frac{\partial \mathbf{F}}{\partial \mathbf{x}} \). We then perform a double-contraction of this new slice against all of those on the cutting board (Fig. 3.3). The result is then a good ole’ vector.

3.3 Multiplication With Flattened Tensors

Just to drive you crazy, there’s a second, more conventional way to write all this down. Any tensor, be it 3rd-order, 4th-order, or 100th order, can always be flattened or vectorized...
Figure 3.3.: Double-contraction of a 3\textsuperscript{rd}-order tensor against a 2\textsuperscript{nd}-order tensor is like taking a cheese slice (right) and double-contracting it against each slice on the cutting board (left). The result here would be an $\mathbb{R}^4$ vector.

back into the familiar 2\textsuperscript{nd}-order matrix form. We’re more-or-less going to follow the convention of Golub and Van Loan (2013) here.

Unfortunately, it will turn out that both the tensor and matrix representations are useful. Just like you had to learn both radians and degrees for trigonometric functions, you’re going to have to learn both the tensor and flattened versions here. It will be worth it, I promise.

We introduce the vectorization operator $\text{vec} (\cdot)$ to convert any matrix to a vector, and any higher-order tensor to a matrix. First up, this is how it converts a matrix to a vector:

$$
\text{vec} (A) = \begin{bmatrix}
  a_0 & a_2 \\
  a_1 & a_3
\end{bmatrix}
$$

The important pattern to notice is that we stacked the columns on top of each other. We did not stack the rows, though we very well could have. This is merely the convention that we have found most useful. Getting this convention right is sufficiently important that I’m just going to go ahead and give you the Matlab and C++ code that does this the right way (Fig. 3.4). If you’re using Octave, it has $\text{vec}$ built-in already, and it follows the same convention we do.

Next up, we define the following convention for 3\textsuperscript{rd}-order tensors:

$$
\text{vec} (A) = \begin{bmatrix}
  [A] \\
  [B] \\
  [C]
\end{bmatrix} = \begin{bmatrix}
  \text{vec} (A) & \text{vec} (B) & \text{vec} (C)
\end{bmatrix}.
$$

First we unfurl the matrices in the 3\textsuperscript{rd}-order tensor into a row, and then vectorize them.
function [x] = vec(A)
    [rows cols] = size(A);
    x = reshape(A, rows * cols, 1);
end

static Vector9 flatten(const Matrix3x3& A)
{
    Vector9 flattened;
    int index = 0;
    for (int y = 0; y < 3; y++)
        for (int x = 0; x < 3; x++, index++)
            flattened[index] = A(x, y);
    return flattened;
}

Figure 3.4.: Our matrix flattening convention, in Matlab and C++.

all in turn. Here’s a concrete example:

\[
\begin{bmatrix}
  a_0 & a_2 \\
  a_1 & a_3 \\
  a_4 & a_6 \\
  a_5 & a_7 \\
  a_8 & a_{10} \\
  a_9 & a_{11}
\end{bmatrix}
= \begin{bmatrix}
  \text{vec} \begin{bmatrix} a_0 & a_2 \end{bmatrix} & \text{vec} \begin{bmatrix} a_4 & a_6 \end{bmatrix} & \text{vec} \begin{bmatrix} a_8 & a_{10} \end{bmatrix}
\end{bmatrix}
= \begin{bmatrix}
  a_0 & a_4 & a_8 \\
  a_1 & a_5 & a_9 \\
  a_2 & a_6 & a_{10} \\
  a_3 & a_7 & a_{11}
\end{bmatrix}
\]

(3.10)

It may seem a little antithetical to the convention that we don’t just stack everything into one long vector. All the matrix multiplies wouldn’t work out in that case though, which would mess up your day even worser.

Finally, let’s look at the flattened version of that $A : B$ double-contraction we saw in
Eqn. 3.7:

\[
(\text{vec } A)^T \text{vec } B = \left( \begin{bmatrix}
 a_0 & a_2 \\
 a_1 & a_3 \\
 a_4 & a_6 \\
 a_5 & a_7 \\
 a_8 & a_{10} \\
 a_9 & a_{11}
\end{bmatrix} \right)^T \begin{bmatrix}
 b_0 & b_2 \\
 b_1 & b_3
\end{bmatrix}
\]  \hspace{1cm} (3.11)

\[
= \begin{bmatrix}
 a_0 & a_4 & a_8 \\
 a_1 & a_5 & a_9 \\
 a_2 & a_6 & a_{10} \\
 a_3 & a_7 & a_{11}
\end{bmatrix}
\begin{bmatrix}
 b_0 \\
 b_1 \\
 b_2 \\
 b_3
\end{bmatrix}
\]  \hspace{1cm} (3.12)

\[
= \begin{bmatrix}
 a_0 b_0 + a_1 b_1 + a_2 b_2 + a_3 b_3 \\
 a_4 b_0 + a_5 b_1 + a_6 b_2 + a_7 b_3 \\
 a_8 b_0 + a_9 b_1 + a_{10} b_2 + a_{11} b_3
\end{bmatrix}
\]  \hspace{1cm} (3.13)

The result is the same as Eqn. 3.7, but arrived at (at least in the final step) through a conventional matrix multiply. It works!

3.4 Computing Forces (Finally)

With all this tensor stuff in place, we can finally compute some forces. Once again, the force is \( f = -a \frac{\partial \Psi}{\partial \mathbf{x}} \), so the equation we need to compute is:

\[
\frac{\partial \Psi}{\partial \mathbf{x}} = \frac{\partial \mathbf{F}}{\partial \mathbf{x}} : \frac{\partial \Psi}{\partial \mathbf{F}}.
\]  \hspace{1cm} (3.14)

The trick is that the 3rd-order tensor, \( \frac{\partial \mathbf{F}}{\partial \mathbf{x}} \), has a static, simple, and energy-independent structure that we can code up once and forget about. The other term, \( \frac{\partial \Psi}{\partial \mathbf{F}} \), will have a simple and clean structure. If you want to implement a new energy, all you need to do is derive a new \( \frac{\partial \Psi}{\partial \mathbf{F}} \).

3.4.1 Computing the \( \frac{\partial \mathbf{F}}{\partial \mathbf{x}} \) Tensor

To see where \( \mathbf{x} \) appears in \( \mathbf{F} \), we’re going to use one of the expressions we derived in Appendix D:

\[
\mathbf{F} = \mathbf{D}_x \mathbf{D}_{x_0}^{-1}
\]  \hspace{1cm} (3.15)

\[
= \begin{bmatrix}
 \mathbf{x}_1 - \mathbf{x}_0 & \mathbf{x}_2 - \mathbf{x}_0 & \mathbf{x}_3 - \mathbf{x}_0
\end{bmatrix}
\begin{bmatrix}
 \mathbf{x}_1 - \mathbf{x}_0 & \mathbf{x}_2 - \mathbf{x}_0 & \mathbf{x}_3 - \mathbf{x}_0
\end{bmatrix}^{-1}
\]  \hspace{1cm} (3.16)
Fortunately we’re differentiating with respect to \( x = \begin{bmatrix} x_0 \\ x_1 \\ x_2 \\ x_3 \end{bmatrix} \) and not \( \bar{x} = \begin{bmatrix} \bar{x}_0 \\ \bar{x}_1 \\ \bar{x}_2 \\ \bar{x}_3 \end{bmatrix} \), so we don’t have to figure out how to take the derivative of a matrix inverse (yuck). In the end, we’re going to get a bunch of expressions that look like:

\[
\frac{\partial F}{\partial x_i} = \frac{\partial D_s}{\partial x_i} D_m^{-1}.
\]

(3.17)

We already have \( D_m^{-1} \) in hand, and \( \frac{\partial D_s}{\partial x_i} \) will work out to a bunch of simple matrices. There will be a lot of them (twelve!), but they will be simple. It takes a lot of space to list these, so instead I’ve stashed it all in Appendix E.

We could compute things the purely matrix way as well,

\[
\frac{\partial \Psi}{\partial x} = \text{vec} \left( \frac{\partial F}{\partial x} \right)^T \text{vec} \left( \frac{\partial \Psi}{\partial F} \right).
\]

(3.18)

There’s C++ code in Fig. E.1 that explicitly lays out the flattened version of \( \text{vec} \left( \frac{\partial F}{\partial x} \right) \) for you. I’ve had to derive this matrix several times already in my lifetime, and I’m getting a little tired of it, so I’m caching it here for posterity. You’re welcome, Future Me.

### 3.5 Dirichlet Forces: So Easy

Back in §2.4.2, we tried to compute forces for the Dirichlet energy,

\[
\Psi_{\text{Dirichlet}} = \|F\|_F^2,
\]

but gave up because the expressions rolled out of control the moment you tried to differentiate an \( F \)-based energy using \( x \). We’ve now skimmed off all the difficulty and distilled it into \( \frac{\partial F}{\partial x} \). All that’s left is to take \( \frac{\partial \Psi}{\partial F} \), and this is probably pretty easy for an \( F \)-based energy, right?

The matrix \( \frac{\partial \Psi}{\partial F} \) is also known as the first Piola-Kirchhoff stress tensor, which is an unwieldy name, so we abbreviate it to the PK1, and sometimes write it as \( P(F) \). The PK1 of the Dirichlet energy is:

\[
\frac{\partial \Psi_{\text{Dirichlet}}}{\partial F} = P(F) = 2F.
\]

(3.19)

SO EASY. It’s almost as easy as computing the forces of the spring-mass system in Eqn. 2.25. In fact, it looks like an intro-to-calculus scalar derivative. The \( F^T F \) expression kind of looks like the matrix version of \( F^2 \), and then we took the derivative to get \( 2F \). There are indeed a bunch of matrix calculus identities that look like this, and a few useful ones are listed in Appendix B. Now to compute the forces, all we do is call the C++ code I gave you in Fig. E.1, and multiply it by \( \text{vec}(2F) \).
3.6 Other Forces: Still Pretty Easy

If we want to compute the forces for some of the other energies we looked at, it becomes straightforward.

3.6.1 St. Venant Kirchhoff, Stretching Only

Let’s look at the stretching term for the St. Venant-Kirchhoff model from before:

\[ \Psi_{\text{StVK, stretch}} = \| \mathbf{E} \|^2_F, \tag{3.20} \]

where \( \mathbf{E} = \frac{1}{2} (\mathbf{F}^T \mathbf{F} - \mathbf{I}) \). If we learned any lesson before, it’s that we should take the derivative with respect to whatever basic variable the energy is written in terms of (\( \mathbf{F} \), for the Dirichlet case), and then bake out all the difficult change-of-variable stuff into some other tensor, \( \frac{\partial \mathbf{F}}{\partial \mathbf{x}} \).

The basic variable for this energy is sure seems like \( \mathbf{E} \), not \( \mathbf{F} \). We could indeed write things out as \( \frac{\partial \Psi}{\partial \mathbf{x}} = \frac{\partial \Psi}{\partial \mathbf{E}} \frac{\partial \mathbf{E}}{\partial \mathbf{x}} \). Then we’d have to derive a new change-of-basis tensor, \( \frac{\partial \mathbf{E}}{\partial \mathbf{x}} \). But, once we had that in hand, computing \( \frac{\partial \Psi}{\partial \mathbf{E}} \), otherwise known as the second Piola-Kirchhoff stress tensor (PK2) would be really easy:

\[ \frac{\partial \Psi_{\text{StVK, stretch}}}{\partial \mathbf{E}} = \mathbf{E}. \tag{3.21} \]

Before we go down the PK2 road though, you should know that StVK is the only energy that really uses \( \mathbf{E} \). All the other energies\(^5\) use \( \mathbf{F} \). So, before we build a customized \( \frac{\partial \mathbf{E}}{\partial \mathbf{x}} \) tensor for StVK, let’s see how difficult it is to go down the \( \mathbf{F} \) road.

\[ \Psi_{\text{StVK, stretch}} = \frac{1}{4} \| \mathbf{F}^T \mathbf{F} - \mathbf{I} \|^2_F, \]

\[ = \frac{1}{4} \left( \| \mathbf{F}^T \mathbf{F} \|^2_F + \text{tr} \mathbf{I} - 2 \text{tr}(\mathbf{F}^T \mathbf{F}) \right) \]

\[ = \frac{1}{4} \left( \| \mathbf{F}^T \mathbf{F} \|^2_F + \text{tr} \mathbf{I} - 2 \| \mathbf{F} \|^2_F \right). \tag{3.22} \]

We already know how to take the derivative of \( \| \mathbf{F} \|^2_F \), since it’s just the Dirichlet energy, and the derivative of \( \| \mathbf{F}^T \mathbf{F} \|^2_F \) is in Appendix B.

\[ \mathbf{P}_{\text{StVK, stretch}}(\mathbf{F}) = \frac{1}{4} \left( 4 \mathbf{F}^T \mathbf{F} - 4 \mathbf{F} \right) \]

\[ = \mathbf{F} \left( \mathbf{F}^T \mathbf{F} - \mathbf{I} \right) \]

\[ = \mathbf{F} \mathbf{E} \tag{3.23} \]

STILL PRETTY EASY. The pattern holds here where \( \| \mathbf{F}^T \mathbf{F} \|^2_F \) looks suspiciously like a matrix version of \( \mathbf{F}^4 \), and its derivative \( 4 \mathbf{F} \mathbf{F}^T \mathbf{F} \) looks an awful lot like \( 4 \mathbf{F}^3 \).

\(^5\)The ones I care about, anyway.

30
3.6.2 The Complete St. Venant Kirchhoff

Up until now, we’ve only looked at the stretching term from StVK, but now is a good moment to look at the complete energy:

$$\Psi_{StVK} = \mu \|E\|_F^2 + \frac{\lambda}{2} (\text{tr } E)^2$$  \hspace{1cm} (3.24)

The $\|E\|_F^2$ term we’ve been examining is there of course, but a new volume preservation term $(\text{tr } E)^2$ has been added\(^6\). The constants $\mu$ and $\lambda$ are present to you can tell the model how much relative stretching resistance vs. volume preservation you want. The PK1 is then:

$$P_{StVK}(F) = \mu FE + \lambda (\text{tr } E) F.$$ \hspace{1cm} (3.25)

To arrive at this, we used identities B.12 and B.30 from Appendix B. Some minor gymnastics are required if you’re not used to this sort of thing, but the final expression is still short and easy to digest.

3.6.3 As-Rigid-As-Possible

The As-Rigid-As-Possible (ARAP) energy can be expanded to:

$$\Psi_{ARAP} = \frac{\mu}{2} \|F - R\|_F^2$$ \hspace{1cm} (3.26)

$$= \frac{\mu}{2} \left( \|F\|_F^2 + \|R\|_F^2 - 2 \text{tr } (F^T R) \right)$$ \hspace{1cm} (3.27)

$$= \frac{\mu}{2} \left( \|F\|_F^2 + \|R\|_F^2 - 2 \text{tr } S \right)$$ \hspace{1cm} (3.28)

We used identity B.16 and the polar decomposition $F = RS \implies S = F^T R$ (since $S$ is symmetric) to arrive at this form. The PK1 is then:

$$P_{ARAP}(F) = \mu (F - R)$$ \hspace{1cm} (3.29)

We used identities B.17 and B.32 to arrive at this expression. This one looks quite appealing! We just extract the expression inside the Frobenius norm, and that becomes the PK1. This is extremely similar to what we saw in §2.4.3 with the spring-mass system, where we just extracted the vector from inside the 2-norm. This is not a coincidence, because as we’ll see later, ARAP has quite a strong claim on the title of “Most-Spring-Mass-Like in F-based World”.

3.6.4 The Many Forms of Neo-Hookean

Back in §2.3.4.1, we looked at the Neo-Hookean energy:

$$\Psi_{Neo-Hookean} = \|F\|_F^2 - 3.$$ \hspace{1cm} (3.30)

\(^6\text{We’ll see later that it doesn’t actually preserve volume that well if you deform the object too much.}\)
If we get the PK1 of this energy,

$$P_{\text{Neo-Hookean}}(F) = 2F,$$  \(3.31\)

it’s exactly the same PK1 as the Dirichlet energy. The only difference between this energy and Dirichlet was the minus three, and that gets burned off under differentiation. From any practical force-generating standpoint, this energy is equivalent to Dirichlet.

Why does it even exist then? In the original paper by Mooney (1940), an additional equation appeared in the form of a hard constraint, \(\det F = 1\). We’re not going to look at methods like this, because this constraint is too severe for the kinds of deformations we see in computer animation.

There are many energies that call themselves “Neo-Hookean”, so we’ll look at one of the more popular ones from Bonet and Wood (2008):

$$\Psi_{\text{BW08}} = \mu \left( \| F \|_F^2 - 3 \right) - \mu \log(J) + \lambda \frac{\left( \log(J) \right)^2}{2}. \quad (3.32)$$

I’ve put this off as long as possible, but here we are starting to use the shorthand of \(J = \det F\). This \(\det F\) term is extremely important because it tells us how much volume has or hasn’t been preserved. As we’ll see later, any energy that claims to be “volume preserving” but doesn’t contain \(\det F\) is just a sham pretender to the throne.

Since \(\det F\) is going to start showing up a lot, it’s time we started using a symbol that’s simpler to read, and \(J\) is extremely common elsewhere in the literature (e.g. Bonet and Wood (2008), Marsden and Hughes (1994), Bower (2009)).

The PK1 of \(\Psi_{\text{BW08}}\) is then:

$$P_{\text{BW08}}(F) = \mu \left( F - \frac{1}{J} \frac{\partial J}{\partial F} \right) + \lambda \frac{\log J}{J} \frac{\partial J}{\partial F}. \quad (3.33)$$

The new term that appears is \(\frac{\partial J}{\partial F}\), the gradient of \(J\). In 3D, this term can be written in terms of the columns of \(F\),

$$F = \begin{bmatrix} f_0 & f_1 & f_2 \end{bmatrix}. \quad (3.34)$$

The gradient of \(J\) is then the pair-wise cross-products of these columns,

$$\frac{\partial J}{\partial F} = \begin{bmatrix} f_1 \times f_2 & f_2 \times f_0 & f_0 \times f_1 \end{bmatrix}. \quad (3.35)$$

This identity is important enough that we stacked it into Appendix B, so when you need it later, you can find it there too. Once you have this identity in hand, deriving the PK1 for an energy that contains a \(\det F = J\) term is not a big deal. Take scalar derivatives like usual, and then you finally hit a \(\frac{\partial J}{\partial F}\) you can stop, because we’ve got an identity for that.

As we’ll see later, the \(\frac{\partial J}{\partial F}\) term appears in lots of places. Just to underscore how important it is, I’m giving you a Matlab implementation in Fig. 3.5
function [final] = gradientJ(F)
f0 = F(:,1);
f1 = F(:,2);
f2 = F(:,3);
final = [cross(f1,f2) cross(f2,f0) cross(f0,f1)];
end

Figure 3.5.: Matlab/Octave code for the 3D version of $\frac{\partial J}{\partial F}$ from Eqn. 5.6.

3.7 What Now?

We’ve now seen how to compute forces the tensor way. This is enough to get a simulator off the ground that does explicit time integration, but that’s not good enough. We want implicit timestepping, since the Fizt integrator from Baraff and Witkin (1998), and the large timesteps it enables, is our target deployment.

For that, we need force gradients, i.e. $\frac{\partial f}{\partial x}$. All the tensor stuff we just saw made force computation easier and prettier, but all the ulcer-inducing ugliness comes rushing back the moment we try to take a force gradient. It’ll take some more work to make force gradient computation equivalently easy and pretty. Let’s take a look at that next.
Chapter 4

Computing Force Gradients the Tensor Way

If you’re just joining us, we’re looking at Baraff and Witkin (1998)-style implicit integration and how it uses force gradients to enable large, stable timesteps. If you thought that computing forces directly from the strain energy was painful,

$$ f = -v \frac{\partial \Psi}{\partial x}, \quad (4.1) $$

then, hoo nelly, are these force gradients excruciating:

$$ \frac{\partial f}{\partial x} = -v \frac{\partial^2 \Psi}{\partial x^2}. \quad (4.2) $$

The way we cleaned everything up in Chapter 3 was to apply the chain rule,

$$ \frac{\partial \Psi}{\partial x} = \frac{\partial F}{\partial x} : \frac{\partial \Psi}{\partial F}, \quad (4.3) $$

which baked all the difficulty into an energy-agnostic tensor $\frac{\partial F}{\partial x}$ that we only have to derive once. Then for each energy model, we only have to deal with the relatively easier $\frac{\partial \Psi}{\partial F}$, a.k.a the first Piola-Kirchhoff stress tensor (PK1).

Can we do something similar for the force gradient? Something like

$$ \frac{\partial^2 \Psi}{\partial x^2} = \frac{\partial F^T}{\partial x} \frac{\partial^2 \Psi}{\partial F} \frac{\partial F}{\partial x} \quad (4.4) $$

sure seems promising. But just like in Chapter 3, you’ll discover that the whole thing starts looking like a clear-as-mud, dirty professor trick as soon as you try to apply this equation. Let’s look at it in more detail, because it’ll take some work to actually use it.
4.1 4th-order Tensors

The first observation that will make you curse the sky is that $\frac{\partial^2 \psi}{\partial F^2}$ is a 4th-order tensor. To see this more clearly, recall that we started writing $\frac{\partial \psi}{\partial F}$ as the matrix $P(F) \in \mathbb{R}^{3 \times 3}$. We can write $\frac{\partial^2 \psi}{\partial F^2}$ as $\frac{\partial P}{\partial F}$, which is the derivative of a matrix with respect to a matrix. We can write this out explicitly as:

$$\frac{\partial P}{\partial F} = \begin{bmatrix}
\frac{\partial P}{\partial f_{00}} & \frac{\partial P}{\partial f_{01}} & \frac{\partial P}{\partial f_{02}} \\
\frac{\partial P}{\partial f_{10}} & \frac{\partial P}{\partial f_{11}} & \frac{\partial P}{\partial f_{12}} \\
\frac{\partial P}{\partial f_{20}} & \frac{\partial P}{\partial f_{21}} & \frac{\partial P}{\partial f_{22}}
\end{bmatrix}.$$  \hspace{1cm} (4.5)

As in the 3rd-order case, we put brackets around each $\frac{\partial P}{\partial f_{ij}}$ to emphasize that they are in the entries in a higher-order tensor, not the blocks in a traditional 2nd-order matrix.

Just like the 3rd-order tensor was a vector of matrices, a 4th-order tensor is a matrix of matrices. Instead of a cutting board containing a line of cheese slices, the cutting board now has cheese slices arranged in a grid (Fig. 4.1).

![Figure 4.1. A 4th-order tensor: like a cutting board covered with a grid of individually wrapped slices of American cheese. A matrix of matrices.](image)

A brief digression: this mental picture does indeed generalize to even higher orders. If you ever needed the next derivative $\frac{\partial^3 \psi}{\partial F^3}$, you would need to compute a 6th-order tensor, i.e. a matrix of matrices of matrices.3 You can just stack another layer on top (Fig. 4.2). For 8th-order, you would take four copies and arrange them in a grid again. And so on to $\infty$th-order, like a giant fractal pagoda.

---

1The $(F)$ argument of $P$ has been dropped for brevity.
2We won’t cover it here, but I have needed this before, so you may encounter a case where you do too.
Figure 4.2.: A 6th-order tensor: we take the cutting boards of cheese from Fig. 4.1, make four copies, get an even bigger cutting board for underneath, and arrange them all into another grid. So much cheddar.

Now back to the 4th-order case. Double-contraction looks similar to the 3rd-order case,

\[
\begin{bmatrix}
  a_0 & a_2 \\
  a_1 & a_3 \\
  a_4 & a_6 \\
  a_5 & a_7
\end{bmatrix}
\begin{bmatrix}
  a_8 & a_{10} \\
  a_9 & a_{11} \\
  a_{12} & a_{14} \\
  a_{13} & a_{15}
\end{bmatrix}
\begin{bmatrix}
  b_0 & b_2 \\
  b_1 & b_3
\end{bmatrix}
= \begin{bmatrix}
  (a_0b_0 + a_1b_1 + a_2b_2 + a_3b_3) \\
  (a_4b_0 + a_5b_1 + a_6b_2 + a_7b_3) \\
  (a_8b_0 + a_9b_1 + a_{10}b_2 + a_{11}b_3)
\end{bmatrix} \begin{bmatrix}
  (a_0b_0 + a_1b_1 + a_2b_2 + a_3b_3) \\
  (a_4b_0 + a_5b_1 + a_6b_2 + a_7b_3) \\
  (a_8b_0 + a_9b_1 + a_{10}b_2 + a_{11}b_3)
\end{bmatrix},
\]

(4.6)

except instead of producing a vector, the results are arranged into a matrix.

If we want to flatten the following tensor,

\[
C = \begin{bmatrix}
  1 & 3 \\
  2 & 4 \\
  5 & 7 \\
  6 & 8
\end{bmatrix}
\begin{bmatrix}
  9 & 11 \\
  10 & 12 \\
  13 & 15 \\
  14 & 16
\end{bmatrix}
\]

(4.7)

then we first flatten in column-wise order, and then each matrix in turn:

\[
\text{vec}(C) = \left[ \begin{array}{c|c|c|c}
\text{vec}(C_{00}) & \text{vec}(C_{10}) & \text{vec}(C_{01}) & \text{vec}(C_{11}) \\
\end{array} \right]
\]

(4.9)

\[
\begin{bmatrix}
  1 & 5 & 9 & 13 \\
  2 & 6 & 10 & 14 \\
  3 & 7 & 11 & 15 \\
  4 & 8 & 12 & 16
\end{bmatrix}
\]

(4.10)
One thing that we have not defined is how to multiply a 4th-order tensor with a 3rd-order tensor. Rather than do that, we are instead going to flatten everything so that it all reduces to a matrix-matrix multiply. Not only will this be easier to think about, but we will see some interesting structures appear.

4.2 Computing Force Gradients

We now have enough operators in hand that we can write the useful-looking chain rule that we saw before in more concrete terms:

$$\frac{\partial^2 \Psi}{\partial \mathbf{x}^T} = \text{vec} \left( \frac{\partial \mathbf{F}}{\partial \mathbf{x}} \right)^T \text{vec} \left( \frac{\partial^2 \Psi}{\partial \mathbf{F}^T} \right) \text{vec} \left( \frac{\partial \mathbf{F}}{\partial \mathbf{x}} \right).$$  \hspace{1cm} (4.11)

We already know how to compute \(\text{vec} \left( \frac{\partial \mathbf{F}}{\partial \mathbf{x}} \right)\) from Appendix E, and in fact have the C++ code for it Fig. E.1. The only term that remains to be derived is \(\text{vec} \left( \frac{\partial^2 \Psi}{\partial \mathbf{F}^T} \right)\). We will abbreviate this matrix as \(\mathbf{H}\), because it represents the energy Hessian.

4.2.1 Dirichlet Hessian: So Easy

Let’s try it out on the simplest energy we know, the Dirichlet energy:

$$\Psi_{\text{Dirichlet}} = \| \mathbf{F} \|_F^2$$

$$\frac{\partial \Psi_{\text{Dirichlet}}}{\partial \mathbf{F}} = \mathbf{P} = 2\mathbf{F}.$$  \hspace{1cm} (4.12)

Each entry \(\frac{\partial \mathbf{P}}{\partial f_{ij}}\) in the 4th-order tensor \(\frac{\partial \mathbf{P}}{\partial \mathbf{F}}\) is then:

$$\frac{\partial \mathbf{P}}{\partial f_{ij}} = 2 \frac{\partial \mathbf{F}}{\partial f_{ij}}.$$  \hspace{1cm} (4.12)

What does each \(\frac{\partial \mathbf{F}}{\partial f_{ij}}\) look like? There’s a lot of them (nine), but they are all quite simple:
Each just has a one at the corresponding entry \((i, j)\). Here’s what we get when we flatten this out:

\[
\text{vec} \left( \frac{\partial \mathbf{F}}{\partial \mathbf{F}} \right) = \begin{bmatrix}
1 & 1 & 1 & 1 \\
1 & 1 & 1 & 1 \\
1 & 1 & 1 & 1 \\
1 & 1 & 1 & 1 \\
\end{bmatrix}
\] (4.13)

IT’S JUST IDENTITY. SO EASY. The Hessian of the Dirichlet energy is then:

\[
\text{vec} \left( \frac{\partial \mathbf{P}}{\partial \mathbf{F}} \right) = 2\mathbf{I}_{9 \times 9}.
\] (4.14)

When we flatten out Hessians, we’ll often see patterns like this appear. Let’s see how far we can take it.

### 4.2.2 Neo-Hookean: Not So Easy

Let’s look at the Bonet and Wood (2008) version of Neo-Hookean again:

\[
\Psi_{\text{BW08}} = \frac{\mu}{2} \left( \|\mathbf{F}\|^2 - 3 \right) - \mu \log(J) + \frac{\lambda}{2} \left( \log(J) \right)^2 
\] (4.15)

\[
\mathbf{P} = \mu \left( \mathbf{F} - \frac{1}{J} \frac{\partial J}{\partial \mathbf{F}} \right) + \lambda \frac{\log J}{J} \frac{\partial J}{\partial \mathbf{F}} 
\] (4.16)

We’re going to have to derive several grisly expressions to get its flattened Hessian, but it will all be worth it, because these same expressions will appear in lots of different energies. In the moment, this will all feel like some sadistic exercise, but in the end, I promise that we will have forged a tool that can be used over and over.

Let’s start by trying to compute one matrix-valued entry in its 4th-order tensor:

\[
\frac{\partial \mathbf{P}}{\partial f_{ij}} = \mu \frac{\partial \mathbf{F}}{\partial f_{ij}} + \mu \frac{\partial J}{\partial f_{ij}} \frac{\partial \mathbf{J}}{\partial \mathbf{F}} - \mu \frac{\partial^2 \mathbf{J}}{\partial \mathbf{F} \partial f_{ij}} - \lambda \frac{\log J}{J} \frac{\partial^2 \mathbf{J}}{\partial \mathbf{F} \partial f_{ij}} - \lambda \frac{\partial J}{J} \frac{\partial \mathbf{J}}{\partial \mathbf{F}} + \frac{\lambda}{J^2} \frac{\partial J}{\partial f_{ij}} \frac{\partial^2 \mathbf{J}}{\partial \mathbf{F} \partial f_{ij}}.
\]

Not very clean. However, if we move things around, we can make things slightly nicer:

\[
\frac{\partial \mathbf{P}}{\partial f_{ij}} = \mu \frac{\partial \mathbf{F}}{\partial f_{ij}} + \left[ \mu + \lambda \left( 1 - \log J \right) \right] \frac{\partial J}{\partial f_{ij}} \frac{\partial \mathbf{J}}{\partial \mathbf{F}} + \left[ \frac{\lambda \log J - \mu}{J} \right] \frac{\partial^2 \mathbf{J}}{\partial \mathbf{F} \partial f_{ij}}.
\]

It’s still not that nice, so I’ll go ahead and put boxes around the non-scalar terms you should care about:

\[
\frac{\partial \mathbf{P}}{\partial f_{ij}} = \mu \frac{\partial \mathbf{F}}{\partial f_{ij}} + \left[ \mu + \lambda \left( 1 - \log J \right) \right] \frac{\partial J}{\partial f_{ij}} \frac{\partial \mathbf{J}}{\partial \mathbf{F}} + \left[ \frac{\lambda \log J - \mu}{J} \right] \frac{\partial^2 \mathbf{J}}{\partial \mathbf{F} \partial f_{ij}}.
\]
The $\mu \frac{\partial F}{\partial f_{ij}}$ term is relatively friendly-looking. If we enumerate all nine copies of this in full 4th-order tensor form, we get $\mu \frac{\partial F}{\partial f}$, and if we flatten that out, we get the same easy-looking term as the Dirichlet energy: $\text{vec} \left( \mu \frac{\partial F}{\partial f} \right) = \mu I_{9 \times 9}$.

There are two other scalar coefficients: $\left[ \mu + \lambda (1 - \log J) \right] J^2$ and $\left[ \frac{\lambda \log J - \mu}{J} \right]$. They’re not pretty, but at least they’re scalar-valued. If we can figure out what the flattened versions of the 4th-order tensors $\frac{\partial J}{\partial f_{ij}} \frac{\partial J}{\partial F}$ and $\frac{\partial^2 J}{\partial F \partial f_{ij}}$ look like, then we’ll have the complete $9 \times 9$ matrix in hand, and we’re in business.

4.2.2.1 The Gradient of $J$

First up, the $\frac{\partial J}{\partial f_{ij}} \frac{\partial J}{\partial F}$ term. Let’s call it $G_J$ in 4th-order form, since it involves two gradients of $J$:

$$G_J = \begin{bmatrix} \frac{\partial J}{\partial f_{00}} & \frac{\partial J}{\partial f_{01}} & \frac{\partial J}{\partial f_{02}} \\ \frac{\partial J}{\partial f_{10}} & \frac{\partial J}{\partial f_{11}} & \frac{\partial J}{\partial f_{12}} \\ \frac{\partial J}{\partial f_{20}} & \frac{\partial J}{\partial f_{21}} & \frac{\partial J}{\partial f_{22}} \end{bmatrix}. \quad (4.17)$$

Writing this out in its full gruesome form doesn’t add much, does it? However, when we perform the vectorization, something friendlier-looking appears:

$$\text{vec} \ G_J = \text{vec} \left( \frac{\partial J}{\partial F} \right) \text{vec} \left( \frac{\partial J}{\partial F} \right)^T. \quad (4.18)$$

It’s just the outer product of the gradient of $J$! If you don’t remember what outer products look like, don’t fret, they’re reviewed in Appendix C.4. With this in hand, we can go ahead and define

$$g_J = \text{vec} \left( \frac{\partial J}{\partial F} \right),$$

i.e. the flattened gradient of $J$, and make the whole thing look even cleaner:

$$\text{vec} \ G_J = g_J \ g_J^T. \quad (4.19)$$

The flattened version of that gnarly-looking $\left[ \frac{\mu + \lambda (1 - \log J)}{J^2} \right] \frac{\partial J}{\partial f_{ij}} \frac{\partial J}{\partial F}$ from before is just:

$$\left[ \frac{\mu + \lambda (1 - \log J)}{J^2} \right] g_J g_J^T.$$

The scalar in front is still ugly, but the rest now looks more like standard linear algebra.
4. Computing Force Gradients the Tensor Way

4.2.2.2 The Hessian of \( J \)

Next up, the \( \frac{\partial^2 J}{\partial f_{ij}^2} \) term, the Hessian of \( J \). We will refer to its full 4th-order form as \( \frac{\partial^2 J}{\partial F^2} \).

This Hessian is a little trickier, because it has a different form in 2D and 3D. Let’s look at the 2D version first, because a really easy structure appears in that case.

**The 2D version:** Remember that \( J = \det F \), so in 2D

\[
J = f_{00}f_{11} - f_{01}f_{10},
\]

and the gradient then becomes

\[
\frac{\partial J}{\partial F} = \begin{bmatrix}
  f_{11} & -f_{10} \\
 -f_{01} & f_{11}
\end{bmatrix}.
\]

The 4th-order tensor can then be written out:

\[
\frac{\partial^2 J}{\partial F^2} = \begin{bmatrix}
  0 & 0 & 0 & 1 \\
  0 & 0 & -1 & 0 \\
  0 & -1 & 0 & 0 \\
  1 & 0 & 0 & 0
\end{bmatrix}.
\] (4.20)

The flattened form is then a really simple anti-diagonal matrix:

\[
\text{vec} \left( \frac{\partial^2 J}{\partial F^2} \right) = \begin{bmatrix}
  0 & 0 & 0 & 1 \\
  0 & 0 & -1 & 0 \\
  0 & -1 & 0 & 0 \\
  1 & 0 & 0 & 0
\end{bmatrix}.
\] (4.21)

Just like we abbreviated the gradient of \( J \) to \( g_J \), let’s call this matrix \( H_J \), since it’s the Hessian of \( J \):

\[
H_J = \text{vec} \left( \frac{\partial^2 J}{\partial F^2} \right).
\] (4.22)

That’s the 2D version.

**The 3D version:** In 3D, things get messier, but it also reveals a more interesting structure. For this version, we first to have to define an operator that converts a vector into a cross-product matrix:

\[
\hat{\mathbf{x}} = \begin{bmatrix}
  0 & -x_2 & x_1 \\
  x_2 & 0 & -x_0 \\
  -x_1 & x_0 & 0
\end{bmatrix}.
\] (4.23)
function \[H\] = hessianJ(F)
zero3 = zeros(3,3);
f0 = F(:,1);
f1 = F(:,2);
f2 = F(:,3);
f0hat = crossMatrix(f0);
f1hat = crossMatrix(f1);
f2hat = crossMatrix(f2);
H = [ zero3 -f2hat f1hat;
     f2hat zero3 -f0hat;
     -f1hat f0hat zero3];
end

function [final] = crossMatrix(f)
final = [ 0 -f(3) f(2);
          f(3) 0 -f(1);
          -f(2) f(1) 0];
end

Figure 4.3.: Matlab/Octave code for the 3D version of vec \( \frac{\partial^2 J}{\partial F^2} \) = \( H_J \) from Eqn. 5.7.

The full 3D Hessian takes a long time to work out, but after lots of symbolic-machete-slaughtering, we find this form:

\[
H_J = \begin{bmatrix}
0_{3\times3} & -\hat{f}_2 & \hat{f}_1 \\
\hat{f}_2 & 0_{3\times3} & -\hat{f}_0 \\
-\hat{f}_1 & \hat{f}_0 & 0_{3\times3}
\end{bmatrix}.
\] (4.24)

The 0s above are all 3 \times 3 zero matrices, and \( f_i \) refer to the columns of \( F \). Now we see the interesting structure. The matrix \( H_J \) is full of cross-product matrices, but compare its structure to Eqn. 4.23. Its block structure follows the cross-product form! It’s a block cross-product matrix of cross-product matrices, or in other words a fractal cross-product. What does it mean? Is there some basic property of Kronecker products that gives rise to this structure? I’m not sure. Sure is interesting though.³

Coding this matrix up right can be somewhat delicate, so I’ve provided a reference Matlab implementation for you in Fig. 4.3.

4.2.2.3 The Full Hessian

Coming back to the Hessian of the Neo-Hookean energy:

\[
\frac{\partial P}{\partial f_{ij}} = \mu \frac{\partial F}{\partial f_{ij}} + \left( \mu + \lambda (1 - \log J) \right) \frac{\partial J}{\partial f_{ij}} \frac{\partial J}{\partial F} + \left( \lambda \log J - \mu \right) \frac{\partial^2 J}{\partial F \partial f_{ij}}.
\]

³To my knowledge, our paper Smith et al. (2018) was the first to observe this structure. Maybe 4D versions of \( J \) will create another fractal level? Sounds like a good homework problem.
Now we have all the pieces to write the boxed terms out in flattened form:

$$\text{vec} \left( \frac{\partial \mathbf{P}}{\partial \mathbf{F}} \right) = \mu \mathbf{I}_{9 \times 9} + \left[ \frac{\mu + \lambda (1 - \log J)}{J^2} \right] g_J g_J^T + \left[ \frac{\lambda \log J - \mu}{J} \right] \mathbf{H}_J.$$

Not too bad. We’re still stuck with the ugly scalar coefficients, but at least the matrix terms, $\mathbf{I}_{9 \times 9}$, $g_J$ and $\mathbf{H}_J$ all have compact forms. We can definitely build this, left- and right-multiply it with $\frac{\partial \mathbf{F}}{\partial \mathbf{x}}$, and get the force gradient.

You may start to worry here: am I going to have to go through this process for every stinking energy? Will there be some new and potentially more hideous $g_*$ and $H_*$ terms that I’ll need to derive every single time? Mercifully, the answer is no.

As we’ll see later, there are a very small number of $g$ and $H$ terms that can ever appear, and $g_J$ and $H_J$ are actually the worst of them. Ugliness will appear elsewhere, but it won’t be because there’s an infinite explosion of $g$- and $H$-like terms that you need to tediously arrange every single time.

### 4.2.3 St. Venant-Kirchhoff: Things Get Worse

Let’s try another energy. This time, we’ll do St. Venant-Kirchhoff, but only the stretching term, because even then, things will get ugly really fast. The energy and its PK1 are:

$$\Psi_{\text{StVK, stretch}} = \| \mathbf{E} \|_F^2$$

$$\mathbf{P} = \mathbf{F} \mathbf{E} = \mathbf{F} (\mathbf{F}^T \mathbf{F} - \mathbf{I})$$

$$= \mathbf{F}^T \mathbf{F} - \mathbf{F}.$$

We’re going to see with this energy, we need to introduce the concept of invariants to get at its flattened Hessian. This concept will become all-consumingly important in §5, so let’s initially dip our toe into the topic by seeing how it crops up in StVK.

Once again, let’s derive a matrix entry of the 4th-order tensor:

$$\frac{\partial \mathbf{P}}{\partial f_{ij}} = \left( \frac{\partial \mathbf{F}}{\partial f_{ij}} \mathbf{F}^T \mathbf{F} + \mathbf{F} \frac{\partial \mathbf{F}}{\partial f_{ij}}^T \mathbf{F} + \mathbf{F} \mathbf{F}^T \frac{\partial \mathbf{F}}{\partial f_{ij}} - \frac{\partial \mathbf{F}}{\partial f_{ij}} \right)$$

(4.26)

On the right, the $\frac{\partial \mathbf{F}}{\partial f_{ij}}$ is our friend the Dirichlet term, whose flattened Hessian will at least contain an easy $\frac{\mu}{2} \mathbf{I}_{9 \times 9}$. The rest of the terms cause more trouble.

To make further progress, we’re going to need the Cauchy-Green invariants$^4$. The invariants are a set of three scalars that often arise when dealing with hyperelastic energies. We’ve actually already seen the first invariant:

$$I_C = \| \mathbf{F} \|^2_F$$

$^4$Technically the right Cauchy-Green invariants, but we’ll drop the “right” here, because it’s not relevant.
We’ve plumbed its depths already by getting all its derivatives:
\[
\frac{\partial I_C}{\partial F} = 2F \quad \frac{\partial^2 I_C}{\partial F \partial f_{ij}} = 2 \frac{\partial F}{\partial f_{ij}} \quad \text{vec} \left( \frac{\partial^2 I_C}{\partial F^2} \right) = 2I_{9 \times 9}.
\]
It’s just the Dirichlet energy! The new trouble we’re seeing arises from the second Cauchy-Green invariant:
\[
II_C = \|F^T F\|_F^2.
\]
The derivatives we’re currently slashing through are:
\[
\frac{\partial II_C}{\partial F} = 4FF^T \quad \frac{\partial^2 II_C}{\partial F \partial f_{ij}} = 4 \left( \frac{\partial F}{\partial f_{ij}} F^T F + F \frac{\partial F^T}{\partial f_{ij}} F + FF^T \frac{\partial F}{\partial f_{ij}} \right).
\]
Hey, that’s the same hairy term from the Hessian of StVK (Eqn. 4.26)! Up to a constant factor. In total, the StVK stretching energy is really just a combination of the first and second invariants. If we can determine the flattened Hessian for the second invariant, then we have all the pieces we need to obtain the flattened Hessian for the StVK stretching term. It’s not pretty, but here it is:
\[
H_{II} = 4 \left( I_{3 \times 3} \otimes FF^T + F^T F \otimes I_{3 \times 3} + D \right).
\]
The \(\otimes\) denotes a Kronecker product, and the \(D\) term works out to:\(^5\)
\[
D = \begin{bmatrix}
  f_0 f_0^T & f_1 f_0^T & f_2 f_0^T \\
  f_0 f_1^T & f_1 f_1^T & f_2 f_1^T \\
  f_0 f_2^T & f_1 f_2^T & f_2 f_2^T
\end{bmatrix}.
\]
If you haven’t seen a Kronecker product before, or haven’t used them in a long time, they are reviewed in Appendix C.5. Like I said, it’s not pretty. But, at least we’ve baked all the ugliness into \(H_{II}\), so we can go straight from this version of the StVK Hessian,
\[
\frac{\partial P}{\partial f_{ij}} = \left( \frac{\partial F}{\partial f_{ij}} F^T F + F \frac{\partial F^T}{\partial f_{ij}} F + FF^T \frac{\partial F}{\partial f_{ij}} \right) - \frac{\partial F}{\partial f_{ij}},
\]
down to this short thing:
\[
\text{vec} \left( \frac{\partial P}{\partial F} \right) = \frac{1}{4} H_{II} - I_{9 \times 9}.
\]
At least we got something, and as with Neo-Hookean, it will turn out that the pieces we’ve just obtained can be used over and over elsewhere. Things could be worse.

\(^5\)This may look like an outer product of \(\text{vec} \, F\), but don’t be fooled. It’s slightly different. Is there another clean explanation? Not that I’ve found, so if you find one, send me an email.
4. As-Rigid-As-Possible: Things Go Terribly Wrong

With the As-Rigid-As-Possible (ARAP) energy, things finally go terribly wrong. It’s a little surprising that such a friendly-looking energy would cause such trouble:

$$\Psi_{\text{ARAP}} = \frac{\mu}{2} \| F - R \|_F^2$$

$$P = \mu (F - R).$$

If you recall from §3.6.3, ARAP was voted “Most Spring-Mass-Like” because, just like with spring-mass forces, its PK1 simply took the expression inside the norm and placed it outside of the norm.

We can write down a matrix entry in its 4th-order tensor thusly:

$$\frac{\partial P}{\partial f_{ij}} = \mu \left( \frac{\partial F}{\partial f_{ij}} - \frac{\partial R}{\partial f_{ij}} \right).$$

The full 4th-order tensor doesn’t look much different:

$$\frac{\partial P}{\partial F} = \mu \left( \frac{\partial F}{\partial F} - \frac{\partial R}{\partial F} \right).$$

The first term is easy: $\frac{\partial F}{\partial F}$ is our Dirichlet buddy yet again. The second term is the trouble-maker. What the heck is $\frac{\partial R}{\partial F}$? The $R$ comes from the polar decomposition $F = RS$, so it arrives as a result of a numerical factorization of $F$. How are we supposed to take the derivative of a numerical factorization? There are no explicit entries of $F$ that appear in $R$, so running back to your old calculus textbook will not help.

We could try applying auto-differentiation to the numerical algorithm used for the polar decomposition. I did this once (Gao et al. (2009)), and it is by far the ugliest and most complicated code that I have ever written. It was also brittle and slow. Don’t try it.

This term is so odd-looking that early works either missed or glossed over it (Müller et al. (2002); Irving et al. (2004)). Later work (Chao et al. (2010); McAdams et al. (2011); Barbic (2012)) noticed its existence, and devised a variety of techniques, of varying complexity, for dealing with it. One thing is for sure: if we want to get an expression for the ARAP energy’s Hessian, we’re going to need something more. We need a way to compute $\frac{\partial R}{\partial F}$, the rotation gradient.

This will turn out to be a can of worms that opens a trapdoor under our feet, throws us down a dusty chute, through roots and vines, and dumps us out on an entirely new forest path. It’ll need its own chapter.
Chapter 5

A Better Way For Isotropic Solids

This chapter is a loose retelling of §4.1 and §4.2 from Smith et al. (2019). If you want to just cut to the chase, the final algorithm is given in §5.5. Everything preceding it will explain where the pieces of this algorithm are coming from, and why they are necessary.

5.1 The Situation So Far

In the previous chapter, we looked at a whole bunch of different deformation energies:

\[
\begin{align*}
\Psi_{\text{Dirichlet}} &= \|F\|^2_F \\
\Psi_{\text{BW08}} &= \frac{\mu}{2} (\|F\|^2_F - 3) - \mu \log(J) + \frac{\lambda}{2} (\log(J))^2 \\
\Psi_{\text{StVK, stretch}} &= \|E\|^2_F \\
\Psi_{\text{ARAP}} &= \frac{\mu}{2} \|F - R\|^2_F
\end{align*}
\]

After jumping through lots of tensor hoops and doing a bunch of tedious manipulations, getting the force gradients for these energies turned out to be possible.\(^1\) Except for ARAP. What’s going on? Why does this friendly-looking energy cause so much trouble?

What we’re going to see is that the first three energies, \(\Psi_{\text{Dirichlet}}\), \(\Psi_{\text{BW08}}\) and \(\Psi_{\text{StVK, stretch}}\) are all Cauchy-Green energies. I mentioned the Cauchy-Green thing in §4.2.3 when we were looking at \(\Psi_{\text{StVK, stretch}}\), but now it will take center stage. Just to offer a spoiler, it will turn out that ARAP is not part of the Cauchy-Green Club, and its non-membership is the source of all the trouble. What does this even mean?

This chapter is longer than the previous ones, so again, if you want to cut to the chase, the final result is in §5.5. Otherwise, let’s get started on our journey.

\(^1\)Just so we don’t lose sight of our goal, we need these gradients if we want to use a Baraff and Witkin (1998)-style simulator.
5.2 The Cauchy-Green Invariants (a.k.a The Wrong Way)

5.2.1 The Gradients and Hessians of the Invariants

There are lots of ways to boil the matrix $F$ down to a scalar, and depending on the method you choose, it can reveal different aspects of what’s going on inside $F$. The Cauchy-Green invariants are one method, and while we will later see that they’re not the one true method, we should first see what they’re good at and what they’re bad at. Then we’ll see if we can fix the bad while keeping the good.

Here are the three Cauchy-Green invariants:

$$I_C = \| F \|_F^2$$

(5.1)

$$II_C = \| F^T F \|_F^2$$

(5.2)

$$III_C = \det \left( F^T F \right).$$

(5.3)

There are lots of identities you can apply to these, so these invariants can show up in different places in lots of sneaky disguises. For example, the first invariant has been known to use the following aliases:

$$I_C = \text{tr} \left( F^T F \right) = \text{tr}(C) = \| F \|_F^2 = F : F = \sum_i \sum_j F_{ij}^2.$$  

(5.4)

Meanwhile, the third invariant can show up in these forms:

$$III_C = \det \left( F^T F \right) = \det(C) = \det(F)^2 = J^2.$$  

(5.5)

Above, $C = F^T F$, and that’s where the $C$ subscript for the invariants comes from.\(^2\)

Why are these invariants useful? Why are we even talking about them? We got a small hint of this in the previous chapter, when figuring out the force gradient of $\Psi_{StVK,\text{stretch}}$ got a lot easier once we introduced $II_C$. Also, I mentioned in §4.2.2.3 that there is a limited rogue’s gallery of $g^*$ and $H^*$ terms that can ever appear.

The invariants bring these two advantages together. If we can get the gradients and Hessians of all three invariants, then deriving the an energy’s force gradient becomes much easier. This is because the gradients and Hessians of the invariants are the complete rogue’s gallery. Once we have these, everything becomes a mix-and-match exercise involving the same pieces over and over.\(^3\)

---

\(^2\)If you’ve taken a theory-slanted linear algebra class, or maybe a visualization class, you may recognize these as the tensor invariants that arise from the characteristic polynomial of $C$. If you don’t know what those words mean, that’s okay too.

\(^3\)Except when we try it on the ARAP energy! But, let’s first see how high we can fly with this Cauchy-Green thing before ARAP causes us to face-plant.
We saw the gradient and Hessian of $I_C$ before in §4.2.3:

$$\frac{\partial I_C}{\partial F} = 2F \quad \quad \quad \frac{\partial^2 I_C}{\partial F^2} = 2 \frac{\partial F}{\partial F}.$$  

We flatten these to get the gradient and Hessian versions:

$$g_I = \text{vec} \left( \frac{\partial I_C}{\partial F} \right) = 2 \text{vec}(F)$$  
$$H_I = \text{vec} \left( \frac{\partial^2 I_C}{\partial F^2} \right) = 2I_{9\times9}.$$  

We can do the same thing for $II_C$,

$$g_{II} = 4 \text{vec} \left( FE \right)$$  
$$H_{II} = 4 \left( I_{3\times3} \otimes FF^T + F^T F \otimes I_{3\times3} + D \right),$$  

where $D$ was defined in Eqn. 4.27. Finally, here are the gradient and Hessian of $III_C$,

$$g_{III} = 2 \det J \cdot g_J$$  
$$H_{III} = 2g_J g_J^T + 2 \det J \cdot H_J$$

where we saw $g_J$ and $H_J$ back in Eqns. 5.6 and 5.7:

$$g_J = \text{vec} \left( \frac{\partial J}{\partial F} \right) = \text{vec} \left( \begin{bmatrix} f_1 \times f_2 & f_2 \times f_0 & f_0 \times f_1 \end{bmatrix} \right) \quad (5.6)$$  
$$H_J = \begin{bmatrix} 0 & -\hat{f}_2 & \hat{f}_1 \\ -\hat{f}_2 & 0 & -\hat{f}_0 \\ -\hat{f}_1 & \hat{f}_0 & 0 \end{bmatrix}. \quad (5.7)$$  

5.2.2 Getting Any Hessian, the Cauchy-Green Way

With these in hand, we can now use the chain rule\(^4\) to obtain a generic expression for the Hessian of some arbitrary energy, $\Psi$:

$$\text{vec} \left( \frac{\partial^2 \Psi}{\partial F^2} \right) = \frac{\partial^2 \Psi}{\partial I_C^2} g_I g_I^T + \frac{\partial \Psi}{\partial I_C} H_I +$$

$$\quad \quad \quad \frac{\partial^2 \Psi}{\partial II_C^2} g_{II} g_{II}^T + \frac{\partial \Psi}{\partial II_C} H_{II} +$$

$$\quad \quad \quad \frac{\partial^2 \Psi}{\partial III_C^2} g_{III} g_{III}^T + \frac{\partial \Psi}{\partial III_C} H_{III}.$$  

\(^4\)Technically Faà di Bruno’s formula, since it’s a higher order derivative.
A compact, but slightly more obtuse, way of writing this is:

$$\text{vec} \left( \frac{\partial^2 \Psi}{\partial F^2} \right) = \sum_{x \in \{I_C, II_C, III_C\}} \frac{\partial^2 \Psi}{\partial x^2} g_x g_x^T + \frac{\partial \Psi}{\partial x} H_x.$$  (5.8)

We just got done listing all the $g_\ast$ and $H_\ast$ terms in §5.2.1, so there’s no work left to do there. Like I said, it’s the complete rogue’s gallery. Instead, the only new work to be done is deriving all the $\frac{\partial \Psi}{\partial x}$ and $\frac{\partial^2 \Psi}{\partial x^2}$ terms, such as $\frac{\partial \Psi}{\partial I_C}$ and $\frac{\partial^2 \Psi}{\partial I_C^2}$.

Those are all scalar derivatives! You can grind through those with just a high-school level grasp of calculus.\(^5\) If you’re feeling really slovenly, you can even just pawn it off on Mathematica or Wolfram Alpha.

There are now three steps to deriving the Hessian of an energy:

1. Re-write your energy $\Psi$ in terms of $I_C$, $II_C$, and $III_C$.
2. Derive the scalar derivatives $\frac{\partial \Psi}{\partial I_C}$, $\frac{\partial^2 \Psi}{\partial I_C^2}$, $\frac{\partial \Psi}{\partial II_C}$, $\frac{\partial^2 \Psi}{\partial II_C^2}$, $\frac{\partial \Psi}{\partial III_C}$ and $\frac{\partial^2 \Psi}{\partial III_C^2}$. Yes, there are six of them, but c’mon, it’s like six homework problems! And it’s not cheating to use the computer!
3. Plug the results into Eqn. 5.8. You’re all done.

### 5.2.3 St. Venant Kirchhoff Stretching, the Cauchy-Green Way

Let’s try it out on the stretching term from St. Venant Kirchhoff:

$$\Psi_{\text{StVK, stretch}} = ||E||^2_F.$$  (5.9)

**Step 1: Rewrite using invariants.** Expanding out the $E = \frac{1}{2} \left( F^T F - I \right)$ and re-writing in terms of invariants, we get

$$\Psi_{\text{StVK, stretch}} = ||E||^2_F$$  (5.10)

$$= \frac{1}{4} ||F^T F - I||^2_F$$  (Substitute E)

$$= \frac{1}{4} \left( ||F^T F||^2_F - 2 \text{tr}(F^T F) + ||I||^2_F \right)$$  (Apply B.16)

$$= \frac{1}{4} II_C - \frac{1}{2} I_C + 3$$  (Substitute invariants)

We also used $||I||^2_F = 3$ on the last line, which holds in 3D. In 2D, $||I||^2_F = 2$, but this usually isn’t that important, because it works out to a constant either way, and the constant gets burned off when we take the derivative.

\(^5\)If you don’t have a high-school level grasp of calculus, but have gotten this far, then some Srinivasa Ramanujan-type situation is going on, and probably don’t need to read any of this.
Step 2: Take the invariant derivatives. Here they are:

\[
\begin{align*}
\frac{\partial \Psi_{SVK, \text{stretch}}}{\partial I_C} &= \frac{-1}{2} \\
\frac{\partial \Psi_{SVK, \text{stretch}}}{\partial II_C} &= \frac{1}{4} \\
\frac{\partial \Psi_{SVK, \text{stretch}}}{\partial III_C} &= 0
\end{align*}
\]

\[
\frac{\partial^2 \Psi_{SVK, \text{stretch}}}{\partial I_C^2} = 0
\]

\[
\frac{\partial^2 \Psi_{SVK, \text{stretch}}}{\partial II_C^2} = 0
\]

\[
\frac{\partial^2 \Psi_{SVK, \text{stretch}}}{\partial III_C^2} = 0
\]

Wait a minute ... most of these just work out to zero. Did the professor make the homework problem super-easy by accident?! No, the derivatives are usually pretty easy. Like I said, this part turns into a calculus homework problem; one of the easy ones near the beginning, not a hard one from the end.

Step 3: Plug into Eqn. 5.8. Now let’s plug into that sum from before:

\[
\begin{align*}
\vec{\left( \frac{\partial^2 \Psi_{SVK, \text{stretch}}}{\partial \mathbf{F}^2} \right)} &= 0 \cdot \mathbf{g}_{II}^T \mathbf{g}_{II} + \frac{1}{2} \mathbf{H}_{II} + 0 \cdot \mathbf{g}_{III}^T \mathbf{g}_{III} + \frac{1}{4} \cdot \mathbf{H}_{III} + 0 \cdot \mathbf{H}_{III} \\
&= \frac{1}{4} \mathbf{H}_{II} - \frac{1}{2} \mathbf{H}_I \\
&= \frac{1}{4} \mathbf{H}_{II} - \mathbf{I}_9 \\
&= \frac{1}{4} \mathbf{H}_{II} - \mathbf{I}_9 \times 9 \\
&= \frac{1}{4} \mathbf{H}_{II} - \mathbf{I}_9 \\
\end{align*}
\]

(Plug in the definition of \( \mathbf{H}_I \))

It’s exactly the same as Eqn. 4.29, but much easier to derive! The system works.

5.2.4 Neo-Hookean, the Cauchy-Green Way

Next let’s try it out on Bonet and Wood (2008)-style Neo-Hookean:

\[
\Psi_{BW08} = \frac{\mu}{2} \left( \| \mathbf{F} \|_F^2 - 3 \right) - \mu \log(J) + \frac{\lambda}{2} \left( \log(J) \right)^2
\]

Step 1: Rewrite using invariants. We can write this as:

\[
\Psi_{BW08} = \frac{\mu}{2} (I_C - 3) - \mu \log(\sqrt{III_C}) + \frac{\lambda}{2} \left( \log(\sqrt{III_C}) \right)^2
\]

At this point, I’m going to let you in on a secret: using \( \sqrt{III_C} \) is a bad idea. Remember that \( III_C = (\det \mathbf{F})^2 \), and \( J = \det \mathbf{F} \). When we compute \( \sqrt{III_C} \), all we’re doing is taking the sign of \( J \) and throwing it away.

The sign of \( J \) is really useful! If it’s negative, it means that our element has been non-physically poked inside out, a.k.a. inverted. If we’re in this situation, we want to know about it so we can deal with it. We’ll talk about this more later on in §6.1.2. For the time being, let’s keep the \( J \)s in the energy intact:

\[
\Psi_{BW08} = \frac{\mu}{2} (I_C - 3) - \mu \log(J) + \frac{\lambda}{2} \left( \log(J) \right)^2
\]
Step 2: Take the invariant derivatives. Let’s take the derivatives, but instead of \( \frac{\partial \Psi}{\partial \text{III}_C} \) and \( \frac{\partial^2 \Psi}{\partial \text{III}_C^2} \), we’ll use \( \frac{\partial \Psi}{\partial \text{J}} \) and \( \frac{\partial^2 \Psi}{\partial \text{J}^2} \):

\[
\begin{align*}
\frac{\partial \Psi_{BW08}}{\partial \text{I}_C} &= \mu/2 \\
\frac{\partial \Psi_{BW08}}{\partial \text{II}_C} &= 0 \\
\frac{\partial \Psi_{BW08}}{\partial \text{J}} &= \lambda \log J - \mu/J \\
\frac{\partial^2 \Psi_{BW08}}{\partial \text{I}_C^2} &= 0 \\
\frac{\partial^2 \Psi_{BW08}}{\partial \text{II}_C^2} &= 0 \\
\frac{\partial^2 \Psi_{BW08}}{\partial \text{J}^2} &= \frac{\lambda (1 - \log J) + \mu}{J^2}.
\end{align*}
\]

The scalar derivatives are a little more involved than last time, but again, you can just punch it into Mathematica or Wolfram Alpha and have it do all the heavy lifting for you. It’s not cheating. It’s what I just did.

Step 3: Plug into the chain rule. For the final expression, we’re going to again replace all the \( \text{III}_C \) terms with \( \text{J} \) terms, as follows:

\[
\text{vec}\left( \frac{\partial^2 \Psi}{\partial F^2} \right) = \frac{\partial^2 \Psi_{BW08}}{\partial \text{I}_C^2} g_{IJ} g_{IJ}^T + \frac{\partial \Psi_{BW08}}{\partial \text{I}_C} H_I + \frac{\partial^2 \Psi_{BW08}}{\partial \text{II}_C^2} g_{IJ} g_{IJ}^T + \frac{\partial \Psi_{BW08}}{\partial \text{II}_C} H_J + \frac{\partial^2 \Psi_{BW08}}{\partial \text{J}^2} g_{IJ} g_{IJ}^T + \frac{\partial \Psi_{BW08}}{\partial \text{J}} H_J.
\]

Fortunately, we have expressions for \( g_J \) and \( H_J \) in Eqns. 5.6 and 5.7\(^7\), so we get:

\[
\text{vec}\left( \frac{\partial^2 \Psi_{BW08}}{\partial F^2} \right) = \frac{\mu}{2} H_I + \frac{\lambda (1 - \log J) + \mu}{J^2} g_{IJ} g_{IJ}^T + \frac{\lambda \log J - \mu}{J} H_J. \tag{5.17}
\]

\[
= \mu I_{9 \times 9} + \frac{\lambda (1 - \log J) + \mu}{J^2} g_{IJ} g_{IJ}^T + \frac{\lambda \log J - \mu}{J} H_J.
\]

(Plug in the definition of \( H_I \))

Again, it matches Eqn. 4.25 exactly! But, it was much easier to derive.

5.2.5 As-Rigid-As-Possible: Things Go Terribly Wrong (Again)

It sure seems like we now have an easy and generic way of deriving any Hessian. Let’s try it out on ARAP:

\[
\Psi_{\text{ARAP}} = \frac{\mu}{2} \| \text{F} - \text{R} \|_F^2 \tag{5.18}
\]

\(^6\)This is still kosher. You can verify it via the chain rule if it bothers you.

\(^7\)Really, the \( \text{III}_C \) versions just built on top of them. The \( \text{J} \) versions are clearly the more atomic and fundamental expressions, which we’ll see in more detail in §5.3.3.2.
Step 1: Rewrite using invariants. Things will go off the rails quickly:

\[ \Psi_{\text{ARAP}} = \frac{\mu}{2} \| F - R \|_F^2 \]  \hspace{1cm} (5.19)

\[ = \| F \|_F^2 - 2 \text{tr} \left( F^T R \right) + \| R \|_F^2 \]  \hspace{1cm} (Apply B.16)

\[ = I_C - 2 \text{tr} \left( F^T R \right) + 3. \]  \hspace{1cm} (Substitute Invariants)

What the heck is \( \text{tr} \left( F^T R \right) \)? It doesn’t correspond to any of the Cauchy-Green invariants! Like we saw in 4.2.4, the \( R \) term comes from the polar decomposition \( F = RS \), so it arises from the numerical factorization of \( F \).

But that messes everything up! The \( R \) makes it so that we can’t fold all appearances of \( F \) into Cauchy-Green invariants, and that breaks our entire Hessian derivation strategy. Nice going, ARAP.

What went wrong? The ARAP energy has now ruined things twice. In §4.2.4, we tried to do things the hard way and slash our way through a thicket of tensors and derivatives, but came up against an impenetrable rotation gradient term, \( \frac{\partial R}{\partial F} \). At that point, we were stuck. How do you take the symbolic derivative of something (\( R \)) that is purely numerical?

Then, we tried the Cauchy-Green invariant approach, but immediately ran into a \( \text{tr} \left( F^T R \right) \) term that doesn’t fit into our invariants, and again we got stuck. Why do things keep going wrong?

5.3 A Better Set of Invariants?

Our most recent problem, the \( \text{tr} \left( F^T R \right) \) term, provides a good clue. To see why, let’s look again at all the forms that the \( I_C \) invariant can take:

\[ I_C = \text{tr} \left( F^T F \right) = \text{tr}(C) = \| F \|_F^2 = F : F = \sum_i \sum_j F_{ij}^2. \]

I’ve highlighted the most relevant one in red. This sure looks similar to that \( \text{tr} \left( F^T R \right) \) term that was giving us trouble, doesn’t it? Keeping in mind again that \( R \) arose from the polar decomposition \( F = RS \), we can go ahead and write this as,

\[ \text{tr} \left( F^T R \right) = \text{tr} \left( S^T \right) = \text{tr} \left( S \right), \]  \hspace{1cm} (5.20)

where we can drop the transpose at the end because \( S \) is a symmetric matrix.

Can we just add this to our gallery of invariants? Is that even a sensible question to ask? I haven’t actually mentioned yet why these are called invariants. Invariants with respect to what? Understanding these quantities intuitively is about to become important, so let’s look at two different ways to think about them.
5.3.1 Invariants as Rotation Removers

The Cauchy-Green invariants are invariant to rotation, that irritating quantity from way back in §2.3.4 that we poured lots of effort into removing from $F$. Only then could we get a clear picture of how much stretching and squashing was going on. No matter how you rotate some deformed triangle or tetrahedron in space, if you compute $I_C$, $II_C$, or $III_C$, they should return the exact same number. In §2.3.4, we looked at two different approaches to removing the rotation.

5.3.1.1 The St. Venant-Kirchhoff Way

The first was in §2.3.4.3, where we used the StVK-style way,

$$\Psi_{\text{StVK, stretch}} = \frac{1}{2} \| F^T F - I \|_F^2,$$

which uses $F^T F$ to remove rotation. If we write this in terms of the polar decomposition, we can explicitly see the moment the rotation gets burned off:

$$F^T F = (RS)^T RS = S^T R^T RS$$

$$= S^2$$  \hspace{1cm} (Since $R^T R = I$)  \hspace{1cm} (Since $S$ is symmetric)

So really, we could add yet another alias to the list of disguises that $I_C$ can take on,

$$I_C = \text{tr} \left( F^T F \right) = \text{tr}(S^2),$$

and we could rewrite the StVK energy as:

$$\Psi_{\text{StVK, stretch}} = \frac{1}{2} \| S^2 - I \|_F^2.$$

5.3.1.2 The As-Rigid-As-Possible Way

We also looked at a second style of rotation removal in §2.3.4.4, the ARAP-style way:

$$\Psi_{\text{ARAP}} = \| F - R \|_F^2.$$

We can use the fact that the Frobenius norm does not change under rotation to burn off $R$ in this case as well:

$$\Psi_{\text{ARAP}} = \| F - R \|_F^2$$

$$= \| R^T (F - R) \|_F^2$$  \hspace{1cm} (Using rotation invariance of $\| \cdot \|_F$)

$$= \| R^T (RS - R) \|_F^2$$  \hspace{1cm} (Inserting $F = RS$)

$$= \| S - I \|_F^2$$  \hspace{1cm} (Using $R^T R = I$)

Looks pretty close to the StVK energy in this form, doesn’t it? One squares $S$ while the other one doesn’t?
5.3.1.3 Comparing the Two Ways

We can do one last manipulation to get a better understanding of what is going on. The ultimate measure of how much stretching and squashing is really going on in $F$ can be obtained using its SVD, which is $F = UΣV^T$. The diagonal matrix of singular values, $Σ$, shows exactly how much stretching is going on along each of the 3D directions. Can we write each energy in terms of that?

Using the rotation invariance of $\| \cdot \|_F^2$ again, and the definition $S = V^TΣV$, we get:

$$Ψ_{StVK, \text{stretch}} = \frac{1}{2}||S^2 - I||_F^2$$

$$= \frac{1}{2}||V \left( S^2 - I \right) V^T||_F^2$$  (Using rotation invariance of $\| \cdot \|_F$)

$$= \frac{1}{2}||V \left( V^TΣVV^TΣV - I \right) V^T||_F^2$$  (Using $S = V^TΣV$)

$$= \frac{1}{2}||ΣΣ - I||_F^2$$  (Since $Σ$ is a diagonal matrix)

$$Ψ_{ARAP} = ||S - I||_F^2$$

$$= ||V \left( S - I \right) V^T||_F^2$$

$$= ||V \left( V^TΣV - I \right) V^T||_F^2$$

$$= ||Σ - I||_F^2$$

In summary, we now have:

$$Ψ_{StVK, \text{stretch}} = \frac{1}{2}||Σ^2 - I||_F^2$$

$$Ψ_{ARAP} = ||Σ - I||_F^2$$

Ignoring the $\frac{1}{2}$ in front of $Ψ_{StVK, \text{stretch}}$, the only real difference is that StVK squares all the singular values, while ARAP doesn’t. What does that mean?

We can use the same $S = V^TΣV$ identity to obtain a new alias for $I_C$:

$$I_C = \text{tr} \left( F^TF \right) = \text{tr}(S^2) = \text{tr}(Σ^2).$$

Curiously, that irritating $\text{tr} \left( F^T R \right)$ term from ARAP works out to something similar:

$$\text{tr} \left( R^TF \right) = \text{tr}(S) = \text{tr}(Σ).$$

When you boil things down to just the singular values $Σ$, it sure looks like $\text{tr} \left( Σ^2 \right)$ measures something similar to $I_C$. That squaring of $Σ$ though, how important is it? Again, what does it even mean? To answer that, we’ll need to look at the invariants from a different perspective.

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8We’re looking at the qualitative behavior of these energies as $F$ changes, so a constant in front doesn’t matter.
5.3.2 Invariants as Geometric Measurements

In the last section, I said that to see what’s really going on in \( F \), you should look at its singular values, i.e. the \( \Sigma \) in \( F = U \Sigma V^T \). We can interpret the invariants in terms of these singular values.

Looking at the \( \text{tr}(\Sigma^2) \) version of \( I_C \), this works out to:

\[
I_C = \sigma_x^2 + \sigma_y^2 + \sigma_z^2.
\]  

(5.21)

The \( \sigma_{\{x,y,z\}} \) terms have a straightforward geometric interpretation. The \( F \) matrix characterizes rotation and scaling at a quadrature point, i.e. some infinitesimal volumetric piece of the solid. As described previously, the singular values are amount of stretching and squashing along each axis, as shown in Fig. 5.1.

The \( I_C \) invariant is the sum of the squared lengths of each side of the cube: \( I_C = \sigma_x^2 + \sigma_y^2 + \sigma_z^2 \).

I.e. it measures the overall edge lengths of the entire cube.

The \( I_C \) invariant is the sum of the squared lengths of each side of the cube (Fig. 5.2). We can work out similar terms for \( II_C \) and \( III_C \), and show that they have straightforward geometric meanings with respect to this cube.
The $II_{C}$ invariant can be rewritten thusly:

$$II_{C} = \|F^{T}F\|_{F}^{2}$$

$$= tr \left( \left( F^{T}F \right)^{T}F^{T}F \right)$$

$$= tr \left( S^{2}S^{2} \right)$$

$$= tr \left( S^{2}S^{2} \right) \quad \text{(Using } F^{T}F = S^{2}, \text{ from §5.3.1.1)}$$

$$= tr \left( V^{T}S^{4}V \right) \quad \text{(Expanding the } S^{2})$$

$$= tr \left( V^{T}S^{4}V \right) \quad \text{(Using } S = V^{T}S^{2}V)$$

$$= tr \left( V^{T}S^{4}V \right) \quad \text{(Using } VV^{T} = I)$$

$$= tr \left( V^{T}S^{4}V \right) \quad \text{(Consolidating the } \Sigma \text{s)}$$

$$= tr \left( VV^{T}S^{4} \right) \quad \text{(Using the cyclic permutation property)}$$

$$= tr \left( VV^{T}S^{4} \right) \quad \text{(Using } VV^{T} = I)$$

$$= \sigma_{x}^{4} + \sigma_{y}^{4} + \sigma_{z}^{4}$$

The cyclic permutation property of traces is: $tr(ABC) = tr(BCA) = tr(CAB)$.

Thus, the $II_{C}$ invariant is the sum of the lengths of each side of the cube, raised to the fourth power. This seems ... not that different from $I_{C}$. In fact, it seems to encapsulate the same basic information. $II_{C}$ only becomes useful when it is folded into an alternate second invariant:

$$II_{C}^{*} = \frac{1}{2}(I_{C}^{2} - II_{C}). \quad (5.22)$$

Chugging through, we get:

$$II_{C}^{*} = \frac{1}{2} \left( I_{C}^{2} - II_{C} \right)$$

$$= \frac{1}{2} \left( (\sigma_{x}^{2} + \sigma_{y}^{2} + \sigma_{z}^{2})^{2} - II_{C} \right) \quad \text{(Definition of } I_{C})$$

$$= \frac{1}{2} \left( \sigma_{x}^{4} + \sigma_{y}^{4} + \sigma_{z}^{4} + 2(\sigma_{x}\sigma_{y})^{2} + 2(\sigma_{x}\sigma_{z})^{2} + 2(\sigma_{y}\sigma_{z})^{2} - II_{C} \right) \quad \text{(Expanding the square)}$$

$$= \frac{1}{2} \left( \sigma_{x}^{4} + \sigma_{y}^{4} + \sigma_{z}^{4} + 2(\sigma_{x}\sigma_{y})^{2} + 2(\sigma_{x}\sigma_{z})^{2} + 2(\sigma_{y}\sigma_{z})^{2} - \left( \sigma_{x}^{4} + \sigma_{y}^{4} + \sigma_{z}^{4} \right) \right) \quad \text{(Definition of } II_{C})$$

$$= \frac{1}{2} \left( 2(\sigma_{x}\sigma_{y})^{2} + 2(\sigma_{x}\sigma_{z})^{2} + 2(\sigma_{y}\sigma_{z})^{2} \right) \quad \text{(Fourth order terms cancel)}$$

$$= (\sigma_{x}\sigma_{y})^{2} + (\sigma_{x}\sigma_{z})^{2} + (\sigma_{y}\sigma_{z})^{2}$$

This expression is more interesting. Whereas $I_{C}$ was the summed squared lengths of the cube, $II_{C}^{*}$ is the sum of the squared areas of the cube faces (Fig.5.3).
Figure 5.3: $II^*_C$ sums up the squared areas of each side of the cube: 

$$II^*_C = (\sigma_x \sigma_y)^2 + (\sigma_x \sigma_z)^2 + (\sigma_y \sigma_z)^2.$$ 

I.e. it measures the overall face areas of the entire cube.

Finally, let’s look at $III_C$. The determinant of a matrix is the product of its singular values, so this one is relatively easy:

$$III_C = \det(F^T F)$$

$$= \det \left( V \Sigma U^T U \Sigma V^T \right)$$

$$= \det \left( V \Sigma \Sigma V^T \right)$$

$$= (\sigma_x \sigma_y \sigma_z)^2.$$ 

Using $F = U \Sigma V^T$, and $U^T U = I$.

Thus, $III_C$ measures the square of the cube’s volume (Fig. 5.4).

Figure 5.4: $III^*_C$ is the squared volume of the cube: 

$$III^*_C = (\sigma_x \sigma_y \sigma_z)^2.$$ 

Taken together, the three invariants measure some intuitive and relevant geometric quantities:

$$I_C = \sigma_x^2 + \sigma_y^2 + \sigma_z^2 \rightarrow \text{squared edge lengths}$$

$$II^*_C = (\sigma_x \sigma_y)^2 + (\sigma_x \sigma_z)^2 + (\sigma_y \sigma_z)^2 \rightarrow \text{squared face areas}$$

$$III_C = (\sigma_x \sigma_y \sigma_z)^2 \rightarrow \text{squared volume.}$$
This looks like a fairly complete way to describe how some small cube of material can deform, so it makes sense that people have been using the Cauchy-Green invariants to describe deformation for the last 80 years (since Mooney (1940), at least).

So then what is the problem with ARAP? These three invariants look like a completely reasonable way to describe deformations, so why can’t the ARAP energy be written down in terms of them?

5.3.3 A New Set of Invariants

5.3.3.1 A New First Invariant

Back in §5.3.1.3, we saw this alias for $I_C$,

$$I_C = \text{tr}(\Sigma^2),$$

and this one for the irritatingly non-cooperative term in ARAP:

$$\text{tr} \left( R^T F \right) = \text{tr}(S) = \text{tr}(\Sigma).$$

Then in §5.3.2 we got a singular value version of $I_C$,

$$I_C = \sigma_x^2 + \sigma_y^2 + \sigma_z^2,$$

while the non-cooperative term trivially takes this form:

$$\text{tr}(S) = \sigma_x + \sigma_y + \sigma_z.$$

The non-cooperative term is an unsquared version of $I_C$. Here, it finally becomes clear why ARAP can’t be written in terms of the Cauchy-Green invariants. You can’t use a sum of squared values to express a sum of unsquared values.

This strongly suggests that $\text{tr}(S)$ should be its own invariant, because it describes something in $F$ that can’t be described with the Cauchy-Green invariants. Let’s call it:

$$I_1 = \text{tr}(S).$$ (5.24)

5.3.3.2 A New Second and Third Invariant

We have essentially just performed the following substitution:

$$I_C = \text{tr} \left( F^T F \right) \quad F^T F \rightarrow S \quad I_1 = \text{tr} (S)$$ (5.25)

Let’s go ahead and perform the same operation on the second and third invariants to round out the whole set:

$$II_C = \text{tr} \left( F^T F F^T F \right) \quad F^T F \rightarrow S \quad I_2 = \text{tr} \left( S^2 \right)$$ (5.26)

$$III_C = \text{tr} \left( F^T F F^T F \right) \quad F^T F \rightarrow S \quad I_3 = \text{det} (S)$$ (5.27)

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Is this a good set of invariants? One way to decide this is to see if we can express the original Cauchy-Green invariants using them. If we can, then we will know for sure that new invariants can describe a super-set of the phenomena that the Cauchy-Green ones could. That would be an excellent indicator that this new set is legitimate.

Well, $I_C$ is equivalent to $I_2$. We saw that $I_C = \text{tr}(S^2)$ back in §5.3.1.3, so that one is easy. Next up, $III_C = I_3^2$. We had already gotten hints about something like this back in §5.2.4, when $J = \det F$ sure looked a lot more fundamental than $III_C = \det (F^T F)$. This suspicion is now confirmed, since $I_3 = \det S = \det F = J$. This determinant of $F$ already captures all the essential information, so it should have an invariant all to itself.

This leaves $II_C$. Showing the equivalence to our new invariants is straightforward in 2D:

$$II_C = I_2^2 - 2I_3^2$$
$$= (\sigma_x^2 + \sigma_y^2)^2 - 2(\sigma_x\sigma_y)^2$$
$$= \sigma_x^4 + \sigma_y^4 + 2(\sigma_x\sigma_y)^2 - 2(\sigma_x\sigma_y)^2$$
$$= \sigma_x^4 + \sigma_y^4.$$

The 3D version takes more work, but is possible:

$$II_C = \frac{1}{2} \left( I_2^2 - I_1^2 \right) + I_1^2 I_2 + 4I_1 I_3.$$

If you already believe me that this expression is legitimate, feel free to skip forward to §5.3.4. Otherwise, here’s the proof in all its grisly glory. First some auxiliary variables:

$$\alpha = (\sigma_x\sigma_y)^2 + (\sigma_x\sigma_z)^2 + (\sigma_y\sigma_z)^2$$
$$\beta = \sigma_x^2\sigma_y\sigma_z + \sigma_x\sigma_y^2\sigma_z + \sigma_x\sigma_y\sigma_z^2$$
$$\gamma = \sigma_x^3(\sigma_y + \sigma_z) + \sigma_y^3(\sigma_x + \sigma_z) + \sigma_z^3(\sigma_x + \sigma_y)$$
$$I_2^2 = II_C + 2\alpha$$
$$I_1^4 = II_C + 6\alpha + 12\beta + 4\gamma$$
$$I_1^2 I_2 = II_C + 2\alpha + 2\beta + 2\gamma$$
$$I_1 I_3 = \beta$$

Now let’s plug them into our expression:

$$II_C = \frac{1}{2} \left( I_2^2 - I_1^2 \right) + I_1^2 I_2 + 4I_1 I_3$$
$$II_C = \frac{1}{2} (II_C + 2\alpha - II_C - 6\alpha - 12\beta - 4\gamma) + I_1^2 I_2 + 4I_1 I_3 \quad \text{(Expand $I_2^2$ and $I_1^4$)}$$
$$II_C = -4 (\alpha + 3\beta + \gamma) + I_1^2 I_2 + 4I_1 I_3 \quad \text{(Simplify the expansion)}$$
$$II_C = -4 (\alpha + 3\beta + \gamma) + II_C + 2\alpha + 2\beta + 2\gamma + 4\beta \quad \text{(Expand $I_1^2 I_2$ and $I_1 I_3$)}$$
$$II_C = -4 (\alpha + 3\beta + \gamma) + II_C + 4(\alpha + 3\beta + \gamma) \quad \text{(Simplify the expansion)}$$
$$II_C = II_C$$

Our new invariants look legitimate. They are a super-set of Cauchy-Green.
5.3.4 Does ARAP Work Now?

Now let’s revisit §5.2.5 and try to write the ARAP energy in terms of our new invariants:

\[ I_1 = \text{tr} \mathbf{S} \quad I_2 = \text{tr} \left( \mathbf{S}^2 \right) \quad I_3 = \text{det} \mathbf{S} \quad (5.28) \]

Can we do it?

\[
\Psi_{\text{ARAP}} = \| \mathbf{F} - \mathbf{R} \|_F^2 \\
= \| \mathbf{F} \|_F^2 - 2 \text{tr} \left( \mathbf{F}^T \mathbf{R} \right) + \| \mathbf{R} \|_F^2 \\
= I_2 - 2I_1 + 3 \quad \text{(Apply B.16)}
\]

(Since \( \| \mathbf{F} \|_F^2 = I_C = I_2 \) and \( \text{tr} \mathbf{F}^T \mathbf{R} = \text{tr} \mathbf{S} = I_1 \))

Hooray! Now let’s see if we can get the PK1, since we will need it to compute a force:

\[
\frac{\partial \Psi_{\text{ARAP}}}{\partial \mathbf{F}} = \frac{\partial}{\partial \mathbf{F}} (I_2 - 2I_1 + 3) \\
= \frac{\partial I_2}{\partial \mathbf{F}} - 2 \frac{\partial I_1}{\partial \mathbf{F}}. \quad (5.30)
\]

Do we know these derivatives? Since \( I_2 = I_C \), we already have that one from §4.2.3:

\[
\frac{\partial I_2}{\partial \mathbf{F}} = \frac{\partial I_C}{\partial \mathbf{F}} = 2 \mathbf{F}.
\]

What about \( \frac{\partial I_1}{\partial \mathbf{F}} \)? Since \( I_1 \) is our brand-new invariant, we’re going to need to establish a new expression. Fortunately, it turns out to be quite simple:

\[
\frac{\partial I_1}{\partial \mathbf{F}} = \frac{\partial}{\partial \mathbf{F}} \text{tr} \mathbf{S} = \frac{\partial}{\partial \mathbf{F}} \left( \text{tr} \mathbf{R} \mathbf{F}^T \right) = \mathbf{R}. \quad (5.32)
\]

It’s just the rotation component of \( \mathbf{F} \)!

Now we can write the full PK1:

\[
\frac{\partial \Psi_{\text{ARAP}}}{\partial \mathbf{F}} = \frac{\partial I_2}{\partial \mathbf{F}} - 2 \frac{\partial I_1}{\partial \mathbf{F}} \\
= 2 \mathbf{F} - 2 \mathbf{R} \quad (5.34)
\]

\[
= 2(\mathbf{F} - \mathbf{R}) \quad (5.35)
\]

This matches what we got in §4.2.4, so everything checks out\(^9\). Now if we can just get an expression for the Hessian of ARAP, we’re finally all done:

\[
\frac{\partial^2 \Psi_{\text{ARAP}}}{\partial \mathbf{F}^2} = 2 \frac{\partial}{\partial \mathbf{F}} (\mathbf{F} - \mathbf{R}) \\
= 2 \left( \frac{\partial \mathbf{F}}{\partial \mathbf{F}} - \frac{\partial \mathbf{R}}{\partial \mathbf{F}} \right). \quad (5.36)
\]

Oh ... right. There was that \( \frac{\partial \mathbf{R}}{\partial \mathbf{F}} \) rotation gradient term from §4.2.4 that we didn’t know how to deal with. Does our brand new \( I_1 \) invariant help us out with \( \frac{\partial \mathbf{R}}{\partial \mathbf{F}} \)? Not in any way that I can see. ARAP is making trouble again. ARAP, why can’t you just behave?

\(^9\)I dropped the \( \frac{\mu^2}{2} \) term, just to make things a little cleaner.
5.4 The Eigenmatrices of the Rotation Gradient

We have a new set of promising invariants, but one key piece is missing. If we can’t figure out a good way to write down $\frac{\partial^2 I_1}{\partial F^2} = \frac{\partial R}{\partial F}$ then everything was for naught. Or rather, we can use some ugly numerical method to obtain $\frac{\partial R}{\partial F}$ (like the one in Papadopoulo and Lourakis (2000)), and our quest to find a clean and slick method for finding Hessians will come to a messy and vaguely unsatisfying end.

The core issue is that we don’t know how to obtain a tidy symbolic derivative for the numerical quantity $R$. However, as we will see, this derivative does indeed have a simple and clean structure, if you look at it from the right perspective.

To get to that perspective, we need to look along two counter-intuitive directions:

- The eigendecomposition of $\frac{\partial R}{\partial F}$ will have a simple structure, i.e. we will be able to form it directly $Q$ and $\Lambda$ from $\frac{\partial R}{\partial F} = Q\Lambda Q^T$. This is the opposite of how things usually go.

  Usually there’s a simple way to write down some matrix $A$, and then you compute the eigendecomposition, $A = Q\Lambda Q^T$, using some byzantine numerical method. The entries of $Q$ and $\Lambda$ then form an impenetrable sea of numbers with no discernible structure. Only in a few special cases do you ever get a simple, closed-form representation of $Q$.

- We won’t be able to see anything if we flatten $\frac{\partial R}{\partial F}$ out to a matrix and look at its eigenvectors. Instead, we have to recognize $\frac{\partial R}{\partial F}$ for what it really is: a 4th-order tensor (§4.1). We need to gaze at its eigenstructure in all its 4th-order glory.

  In the 4th-order universe, an eigenvector becomes an eigenmatrix. The eigenmatrix view will reveal the fundamental structure of the rotation gradient.

Let’s take the first point for granted: the eigendecomposition of the rotation gradient contains a simple structure that is waiting to be discovered. From there, you’re probably wondering what the eigendecomposition of a 4th-order tensor looks like, and what the heck an eigenmatrix is.

5.4.1 What’s an Eigenmatrix?

Let’s recall the basic eigenvalue problem:

$$Aq_0 = \lambda_0 q_0.$$

The eigenvalue $\lambda_0$ and the eigenvector $q_0$ form an eigenpair of $A$. The vector $q_0$ is special because even after you push it through a multiply with $A$, it remains exactly the same. Except, it was scaled by $\lambda_0$.

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10Like Fourier series popping out of the heat equation. That’s a pretty important one.
For some 4th-order tensor

$$A = \begin{bmatrix}
a_0 & a_2 & a_8 & a_{10} 
a_1 & a_3 & a_9 & a_{11}
a_4 & a_6 & a_{12} & a_{14}
a_5 & a_7 & a_{13} & a_{15}
\end{bmatrix}$$  \hspace{1cm} (5.38)

we can define a similar eigenpair, \((\lambda_0, Q_0)\). The equivalent eigenvalue problem is,

$$\hat{A} : Q_0 = \lambda_0 Q_0,$$

where instead of the \(A q_0\) matrix-vector multiply, we have a double-contraction. We can write down everything down in its grisly, fully-expanded verbosity like this:

$$\hat{A} : Q_0 = \begin{bmatrix}
a_0 & a_2 & a_8 & a_{10} 
a_1 & a_3 & a_9 & a_{11}
a_4 & a_6 & a_{12} & a_{14}
a_5 & a_7 & a_{13} & a_{15}
\end{bmatrix} \begin{bmatrix}q_0 
q_2 
q_1 
q_3\end{bmatrix} = \begin{bmatrix}a_0 q_0 + a_1 q_1 + a_2 q_2 + a_3 q_3 
a_4 q_0 + a_5 q_1 + a_6 q_2 + a_7 q_3 
a_8 q_0 + a_9 q_1 + a_{10} q_2 + a_{11} q_3 
a_{12} q_0 + a_{13} q_1 + a_{14} q_2 + a_{15} q_3\end{bmatrix} = \lambda_0 \begin{bmatrix}q_0 
q_2 
q_1 
q_3\end{bmatrix} = \lambda_0 Q_0.$$

Again, \(Q_0\) is a special matrix that, even after going through that baroque double-contraction operation, emerges essentially unscathed. Except, it was scaled by \(\lambda_0\).

Now the thing is, the \(A q_0\) matrix-vector and \(\hat{A} : Q_0 = \lambda_0 Q_0\) versions are exactly equivalent. If you find some eigenmatrix \(Q_0\) of the tensor \(\hat{A}\), then it is also an eigenvector of the flattened out matrix \(A = \text{vec} \hat{A}\). Again, spelling everything out verbosely:

$$(\text{vec} \hat{A})^T (\text{vec} Q_0) = A^T q_0 = \begin{bmatrix}
a_0 & a_4 & a_8 & a_{12} 
a_1 & a_5 & a_9 & a_{13}
a_2 & a_6 & a_{10} & a_{14}
a_3 & a_7 & a_{11} & a_{15}
\end{bmatrix}^T \begin{bmatrix}q_0 
q_1 
q_2 
q_3\end{bmatrix} = \begin{bmatrix}a_0 q_0 + a_1 q_1 + a_2 q_2 + a_3 q_3 
a_4 q_0 + a_5 q_1 + a_6 q_2 + a_7 q_3 
a_8 q_0 + a_9 q_1 + a_{10} q_2 + a_{11} q_3 
a_{12} q_0 + a_{13} q_1 + a_{14} q_2 + a_{15} q_3\end{bmatrix} = \lambda_0 \begin{bmatrix}q_0 
q_1 
q_2 
q_3\end{bmatrix} = \lambda_0 q_0 = \lambda_0 \text{vec} Q_0.$$
This is good news! We don’t have to come up with a whole new suite of numerical methods to find the eigenmatrices of some tensor \( A \). Instead, we can just flatten things out to \( A = \text{vec} \, \hat{A} \), and run whatever eigenvalue routine you want from Matlab, Eigen, or LAPACK. The \((\lambda_i, q_i)\) pairs that come back can then just be re-folded back up into the eigenmatrix-based pair, \((\lambda_i, Q_i)\).

But if this is the case, why do we even care about eigenmatrices? If they contain the exact same entries as eigenvectors, just repackaged into matrix form, why not use the more familiar vector form?

5.4.2 Structures Lurk in the Decomposition of an Eigenmatrix

The reason we care about eigenmatrices is that once you have a matrix, you can apply a variety of matrix-based tools that are not available for vectors. For example, you could take the eigendecomposition of an eigenmatrix. It would be weirdly meta-mathematical and fractal, but you could do it. Equivalently, you could take the QR decomposition or SVD of an eigenmatrix.

Or, we can apply the results of the SVD from related matrices, and see if those reveals any new structures. The most important matrix we’ve been dealing with throughout these chapters, and ever since I highlighted it in bright red back in §2.3.3, is the deformation gradient, \( F \). Let’s say that we had the SVD of that deformation gradient, \( F = U \Sigma V^T \).

Let’s also say that we went ahead and did the horribly ugly thing by computing the rotation gradient, \( \frac{\partial R}{\partial F} \), using the numerical approach of Papadopoulo and Lourakis (2000). I’ll even give you the actual Matlab code I used to do this in Figs. 5.5 and 5.6.

To keep things simple, let’s look at the 2D version of \( \frac{\partial R}{\partial F} \). In that case, if we poke at the matrices coming out of \texttt{DRDF\_Horrible} in Fig. 5.5, we will quickly discover that it’s a rank-one matrix composed of a single eigenpair, \((\lambda_0, Q_0)\). Hoping to discover some simple structure, we can stare at the entries of \( Q_0 \), or even \( q_0 = \text{vec} \, Q_0 \). As usual, it will just be an impenetrable wall of numbers. Not helpful.

We can poke at it more in Matlab and take its SVD, QR, and eigendecomposition to see if something interesting shows up. Alas, I can tell you from experience that this will not yield any insight either. Finally, while running a few errands over the weekend, and chewing this problem over in the backs of our minds, we get the nagging feeling that everything we’re dealing with really just boils down to \( F \). Why not pound on \( Q_0 \) using the decomposition of \( F \) and see what happens?

We can try rotating \( Q_0 \) into the same space as \( F \) using \( UQ_0V^T \). Nothing much to see here, just the usual sea of numbers. How about we rotate \( Q_0 \) out of the space of \( F \):

\[
U^TQ_0V = \begin{bmatrix} 0 & -0.70711 \\ 0.70711 & 0 \end{bmatrix}
\]  \hspace{1cm} (5.39)

62
function [final] = DRDF_Horrible(R,S)
    H = zeros(4,4);
    for index = 1:4
        i = mod(index - 1, 2);
        j = floor((index - 1) / 2);

        % promote it by a dimension
        i = i + 1;
        j = j + 1;
        index3 = i + (j - 1) * 3;

        R3 = eye(3,3);
        S3 = eye(3,3);
        R3(1:2, 1:2) = R;
        S3(1:2, 1:2) = S;

        [DR3, DS3] = DRDF_Column(R3,S3,index3);

        DR = DR3(1:2,1:2);
        DS = DS3(1:2,1:2);
        column = DR;

        H(:,index) = reshape(column , 4, 1);
    end
    final = H;
end

Figure 5.5.: Matlab code to compute the rotation gradient using a horrible numerical method. We only use this temporarily on our way to finding a clean expression for $\frac{\partial R}{\partial F}$. 

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function [DR, DS] = DRDF_Column(R,S,index)
    i = mod(index - 1, 3);
    j = floor((index - 1) / 3);
    % Matlab indexing
    i = i + 1;
    j = j + 1;
    eiej = zeros(3,3);
    eiej(i,j) = 1;
    G = (eye(3,3) * trace(S) - S) * R';
    Rij = R' * eiej;
    % extract the skew vector
    Rijsym = (Rij - Rij') * 0.5;
    skew = zeros(3,1);
    skew(1) = -Rijsym(2,3);
    skew(2) = Rijsym(1,3);
    skew(3) = -Rijsym(1,2);
    skew = 2 * skew;
    omega = (G^-1) * skew;
    cross = [
              0 -omega(3) omega(2);
              omega(3) 0 -omega(1);
              -omega(2) omega(1) 0];
    DR = cross * R;
    DS = R' * (eiej - DR * S);
end

Figure 5.6.: Matlab code to compute one column of the 3D rotation gradient. Like in Fig. 5.5, we only use this as a waypoint on our journey to find a clean expression for $\frac{\partial R}{\partial F}$ in DRDF_Horrible, I pinned one dimension of this 3D version so that I could mess around with it in 2D.
Hang on a minute. A bunch of zeros appeared, which is usually a good sign. Also, that 0.70711 sure looks familiar from trigonometry. Wasn’t it $\sqrt{\frac{2}{2}} = \frac{1}{\sqrt{2}}$, or something like that? If we pull that constant out, we see that the structure of $Q_0$ is actually:

$$Q_0 = \frac{1}{\sqrt{2}} U \begin{bmatrix} 0 & -1 \\ 1 & 0 \end{bmatrix} V^T. \quad (5.40)$$

That’s ... pretty simple! It’s similar to $F$, but instead of the singular value matrix $\Sigma$ in the middle, it’s just the constant matrix $\begin{bmatrix} 0 & -1 \\ 1 & 0 \end{bmatrix}$. This matrix corresponds to an infinitesimal rotation in the 2D plane, so we call the whole thing a *twist* matrix. The $\frac{1}{\sqrt{2}}$ makes sense too. Eigenvectors have unit magnitude, and since $U$ and $V$ are already unitary matrices, the only things left to normalize are the 1 and $-1$ in the center matrix. Since $\sqrt{1^2 + (-1)^2} = \sqrt{2}$, the normalization factor is $\frac{1}{\sqrt{2}}$.

If this entire process sounds improbably informal, and worryingly outside the hypothesis-experiment-conclusion loop of scientific discovery you were taught in high school, just know that real-world research doesn’t always conform to that model. I know I’m not the only researcher who stumbles across new things by just messing around in Matlab.

### 5.4.3 What About the Eigenvalue?

Let’s look again at our horribly ugly, numerically-based rotation gradient code, `DRDF_Horrible`. Just to see how the $\lambda_0$ eigenvalue evolves, we can start plugging easy-looking integers into the singular values, like $\sigma_x = 2$ and $\sigma_y = 3$, $\sigma_x = 2$ and $\sigma_y = 5$, and so on. You can literally try this out yourself – the code to run these experiments is in Fig. 5.7.

From there, we can see $\lambda_0$ take on values that look suspiciously like some tidy fractions. For example, $(\sigma_0, \sigma_1) = (1, 3)$ yielded 0.4, and $(\sigma_0, \sigma_1) = (2, 5)$ yields 0.28571. Just to confirm our suspicion that these are fractions, we can push them through Matlab’s `rats` function, and see that $\text{rats}(0.4) \rightarrow \frac{2}{5}$ and $\text{rats}(0.28571) \rightarrow \frac{2}{7}$ (last line, Fig. 5.7).

Running a variety of experiments with different $U$, $\Sigma$ and $V$, it will quickly become clear that the eigenvalue takes the general form:

$$\lambda_0 = \frac{2}{\sigma_x + \sigma_y}. \quad (5.41)$$

If the informality and empiricism of this process bothers you, think of it like running experiments to build up some numerical intuition. Once you see a pattern, you can posit a testable theory.

### 5.4.4 Building the Rotation Gradient (Finally)

We now have the single eigenpair that composes the 2D rotation gradient:

$$\lambda_0 = \frac{2}{\sigma_x + \sigma_y} \, \quad Q_0 = \frac{1}{\sqrt{2}} U \begin{bmatrix} 0 & -1 \\ 1 & 0 \end{bmatrix} V^T. \quad (5.42)$$
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```matlab
% Pick your sigmas
Sigma = [2 0,
        0 5];

% Pick some rotations
angleU = 0;
U = [cos(angleU) -sin(angleU);
     sin(angleU)  cos(angleU)];

angleV = 0;
V = [cos(angleV) -sin(angleV);
     sin(angleV)  cos(angleV)];

% compose your F
F = U * Sigma * V';

% compose your polar decomposition
R = U * V';
S = V * Sigma * V';

% get the rotation gradient
H = DRDF_Horrible(R,S)

% what's the eigendecomposition look like?
[Q Lambda] = eig(H)

% is the eigenvalue a rational number?
rats(Lambda(4,4))

Figure 5.7.: Matlab code to experiment with the single eigenvalue of the rotation gradient. Plug different integers into the diagonal of Sigma and see suspiciously rational-looking quantities pop out at the end.
```
function [final] = DRDF_Clean_2D(U, Sigma, V)
Q0 = (1 / sqrt(2)) * U * [0, -1; 1, 0] * V';
q0 = vec(Q0);
lambda0 = 2 / (Sigma(1,1) + Sigma(2,2));
final = lambda0 * (q0 * q0');
endfunction

Figure 5.8.: Fast and compact Matlab code for building the 2D rotation gradient. This implements Eqn. 5.43. Looks nicer than Figs. 5.5 and 5.6, don’t you think?

We can then use an outer product to arrive at a very simple way to compute the flattened rotation gradient:

$$\text{vec}\left( \frac{\partial R}{\partial F} \right) = \frac{2}{\sigma_x + \sigma_y} \text{vec}(Q_0) \text{vec}(Q_0)^T.$$  \hspace{1cm} (5.43)

This also maps onto some very simple Matlab code, given in Fig. 5.8. We can chug through the entire experimental process again to find a similar expression for the 3D rotation gradient. I will spare you all the intermediate Matlab muddling, and instead tell you that the 3D rotation gradient is rank-3, and the three eigenpairs follow a pattern that is similar to 2D:

$$\lambda_0 = \frac{2}{\sigma_x + \sigma_y} \quad Q_0 = \frac{1}{\sqrt{2}} U \begin{bmatrix} 0 & -1 & 0 \\ 1 & 0 & 0 \\ 0 & 0 & 0 \end{bmatrix} V^T \hspace{1cm} (5.44)$$

$$\lambda_1 = \frac{2}{\sigma_y + \sigma_z} \quad Q_1 = \frac{1}{\sqrt{2}} U \begin{bmatrix} 0 & 0 & 0 \\ 0 & 0 & 1 \\ 0 & -1 & 0 \end{bmatrix} V^T \hspace{1cm} (5.45)$$

$$\lambda_2 = \frac{2}{\sigma_x + \sigma_z} \quad Q_2 = \frac{1}{\sqrt{2}} U \begin{bmatrix} 0 & 0 & 1 \\ 0 & 0 & 0 \\ -1 & 0 & 0 \end{bmatrix} V^T. \hspace{1cm} (5.46)$$

We can then use the same outer product form as the 2D case to compute the final $\text{vec}\left( \frac{\partial R}{\partial F} \right)$ matrix,

$$\text{vec}\left( \frac{\partial R}{\partial F} \right) = \sum_{i=0}^{2} \lambda_i \text{vec}(Q_i) \text{vec}(Q_i)^T,$$  \hspace{1cm} (5.47)

where $\lambda_i$ and $Q_i$ correspond to Eqns. 5.44 to 5.46. The corresponding Matlab code is given in Fig. 5.9.

5.5 Building a Generic Hessian (Finally)

We now have everything in hand needed to compute the Hessian for an isotropic energy. Remember what happened back in §5.2.2: we wrote things in terms of the Cauchy-Green
function [H] = DRDF_Clean_3D(U, Sigma, V)
% get the twist modes
T0 = [0 -1 0;
      1 0 0;
      0 0 0];
T0 = (1 / sqrt(2)) * U * T0 * V';
T1 = [0 0 0;
      0 0 1;
      0 -1 0];
T1 = (1 / sqrt(2)) * U * T1 * V';
T2 = [0 0 1;
      0 0 0;
      -1 0 0];
T2 = (1 / sqrt(2)) * U * T2 * V';
%
% get the flattened versions
t0 = vec(T0);
t1 = vec(T1);
t2 = vec(T2);
%
% get the singular values
sx = Sigma(1,1);
sy = Sigma(2,2);
sz = Sigma(3,3);
H = (2 / (sx + sy)) * (t0 * t0');
H = H + (2 / (sy + sz)) * (t1 * t1');
H = H + (2 / (sx + sz)) * (t2 * t2');
end

Figure 5.9.: Fast and compact Matlab code for building the 3D rotation gradient. This is
the outer product of the eigenmatrices from Eqns. 5.44 - 5.46.
invariants, and it looked like we were on easy street for building the final expression. Then we tried to approach on ARAP, which didn’t fit into the Cauchy-Green formalism, and blew it all to smithereens.

Now we do have a set of invariants that encompass ARAP, so let’s use them to get a similarly generic approach. Starting from our new Smith et al. (2019) invariants,

\[
I_1 = \text{tr} (S) \quad I_2 = \text{tr} \left( F^T F \right) \quad I_3 = \det F, \quad (5.48)
\]

we have the gradient (\(g_1\)) and Hessian (\(H_1\)) of each:

\[
g_1 = \text{vec} (R) \quad H_1 = \sum_{i=0}^{2} \lambda_i \text{vec} (Q_i) \text{vec} (Q_i)^T \quad (\text{Eqn. 5.47})
\]

\[
g_2 = \text{vec} (2F) \quad H_2 = 2I_{9 \times 9}
\]

\[
g_J = \text{vec} \left( \begin{bmatrix} f_1 \times f_2 & f_2 \times f_0 & f_0 \times f_1 \end{bmatrix} \right) \quad H_J = \begin{bmatrix} 0 & -\hat{f}_2 & \hat{f}_1 \\ -\hat{f}_2 & 0 & -\hat{f}_0 \\ -\hat{f}_1 & \hat{f}_0 & 0 \end{bmatrix}
\]

Now we can write an alternative expression for the energy Hessian:

\[
\text{vec} \left( \frac{\partial^2 \Psi}{\partial F^2} \right) = \frac{\partial^2 \Psi}{\partial I_1^2} g_1 \otimes g_1^T + \frac{\partial \Psi}{\partial I_1} H_1 + \frac{\partial^2 \Psi}{\partial I_2^2} g_2 \otimes g_2^T + \frac{\partial \Psi}{\partial I_2} H_2 + \frac{\partial^2 \Psi}{\partial I_3^2} g_3 \otimes g_3^T + \frac{\partial \Psi}{\partial I_3} H_3 \quad (5.49)
\]

\[
= \sum_{i=1}^{3} \frac{\partial^2 \Psi}{\partial I_i^2} g_i \otimes g_i^T + \frac{\partial \Psi}{\partial I_i} H_i. \quad (5.50)
\]

Finally, we follow the same three-step process as before, but using the Smith et al. (2019) invariants:

1. Re-write your energy \( \Psi \) using \( I_1, I_2 \) and \( I_3 \).
2. Derive the scalar derivatives \( \frac{\partial \Psi}{\partial I_1}, \frac{\partial^2 \Psi}{\partial I_1^2}, \frac{\partial \Psi}{\partial I_2}, \frac{\partial^2 \Psi}{\partial I_2^2}, \frac{\partial \Psi}{\partial I_3}, \frac{\partial^2 \Psi}{\partial I_3^2} \) and \( \frac{\partial^2 \Psi}{\partial I_i^2} \).
3. Plug the results into Eqn. 5.50.

Let’s put this algorithm through its paces. First, let’s confirm it does the same thing as Cauchy-Green by confirming that we still get the same results for Neo-Hookean.

### 5.5.1 Neo-Hookean, the Smith et al. (2019) Way

Once again, here’s Bonet and Wood (2008)-style Neo-Hookean:

\[
\Psi_{BW08} = \frac{\mu}{2} (\|F\|_F^2 - 3) - \mu \log(J) + \frac{\lambda}{2} (\log(J))^2
\]
Step 1: Rewrite using invariants. This takes the form:

\[ \Psi_{BW08} = \frac{\mu}{2} (I_2 - 3) - \mu \log(I_3) + \frac{\lambda}{2} (\log(I_3))^2 \]

We’re now using \( I_3 \) instead of \( J \), or the even more ridiculous-looking \( \sqrt{\text{III}} \). So, things are already looking cleaner.

Step 2: Take the invariant derivatives. These are similar to before, which is good sign:

\[
\begin{align*}
\frac{\partial \Psi_{BW08}}{\partial I_1} &= 0 \\
\frac{\partial \Psi_{BW08}}{\partial I_2} &= \frac{\mu}{2} \\
\frac{\partial \Psi_{BW08}}{\partial I_3} &= \lambda \log I_3 - \mu \\
\frac{\partial^2 \Psi_{BW08}}{\partial I_1^2} &= 0 \\
\frac{\partial^2 \Psi_{BW08}}{\partial I_2^2} &= 0 \\
\frac{\partial^2 \Psi_{BW08}}{\partial I_3^2} &= \frac{\lambda(1 - \log I_3) + \mu}{I_3}.
\end{align*}
\]

Step 3: Plug into the chain rule. Plugging into Eqn. 5.50, but dropping everything with a zero coefficient, we get:

\[
\text{vec} \left( \frac{\partial^2 \Psi_{BW08}}{\partial F^2} \right) = \mu I_{9 \times 9} + \frac{\lambda(1 - \log I_3) + \mu}{I_3} g_3 g_3^T + \frac{\lambda \log I_3 - \mu}{I_3} H_3.
\]

It matches both Eqn. 4.25 and Eqn. 5.17: mission accomplished! The Smith et al. (2019) Way is looking legit.

5.5.2 ARAP, the Smith et al. (2019) Way

Now let’s look again at our troublemaking friend, ARAP:

\[ \Psi_{ARAP} = ||F - R||_F^2. \]

Step 1: Rewrite using invariants. Unlike the last time we tried this in §5.2.5, we now have the \( I_1 \) invariant at our disposal. We actually already did this rewrite in §5.3.4, and it worked out to:

\[ \Psi_{ARAP} = I_2 - 2I_1 + 3. \]

Step 2: Take the invariant derivatives. Here we go:

\[
\begin{align*}
\frac{\partial \Psi_{ARAP}}{\partial I_1} &= -2 \\
\frac{\partial \Psi_{ARAP}}{\partial I_2} &= 1 \\
\frac{\partial \Psi_{ARAP}}{\partial I_3} &= 0 \\
\frac{\partial^2 \Psi_{ARAP}}{\partial I_1^2} &= 0 \\
\frac{\partial^2 \Psi_{ARAP}}{\partial I_2^2} &= 0 \\
\frac{\partial^2 \Psi_{ARAP}}{\partial I_3^2} &= 0.
\end{align*}
\]
function [H] = ARAP_Hessian(F)

[U Sigma V] = svd_rv(F);

% get the twist modes
T0 = [0 -1 0; 1 0 0; 0 0 0];
T0 = (1 / sqrt(2)) * U * T0 * V';

T1 = [0 0 0; 0 0 1; 0 -1 0];
T1 = (1 / sqrt(2)) * U * T1 * V';

T2 = [0 0 1; 0 0 0; -1 0 0];
T2 = (1 / sqrt(2)) * U * T2 * V';

% get the flattened versions
t0 = vec(T0);
t1 = vec(T1);
t2 = vec(T2);

% get the singular values in an order that is consistent
% with the numbering in [Smith et al. 2019]
s0 = Sigma(1,1);
s1 = Sigma(2,2);
s2 = Sigma(3,3);

H = 2 * eye(9,9);
H = H - (4 / (s0 + s1)) * (t0 * t0');
H = H - (4 / (s1 + s2)) * (t1 * t1');
H = H - (4 / (s0 + s2)) * (t2 * t2');
end

Figure 5.10.: Matlab code to compute the exact Hessian of the ARAP energy in 3D. The code for svd_rv is given in Fig. F.1.

This time it was easy. I didn’t even have to go running to Mathematica for this one.

Step 3: Plug into the chain rule. Again but dropping everything with a zero coefficient, we get:

\[ \text{vec} \left( \frac{\partial^2 \Psi_{\text{ARAP}}}{\partial F^2} \right) = 2I_{9 \times 9} - 2H_1. \]

FINALLY, A SIMPLE EXPRESSION. I ALWAYS KNEW YOU HAD IT IN YOU, ARAP. The Matlab code to compute this is shown in Fig. 5.10.

5.5.3 Symmetric Dirichlet, the Smith et al. (2019) Way

Just to build some confidence that this approach is indeed generic and we’re not fooling ourselves, let’s try it out on an energy we haven’t seen before, the Symmetric Dirichlet energy from Smith and Schaefer (2015a):

\[ \Psi_{\text{Symmetric}} = \|F\|^2_F + \|F^{-1}\|^2_F. \] (5.57)
Step 1: Rewrite using invariants. We’re not quite on easy street with this energy. Is it possible to write $\|F^{-1}\|^2$ in terms of the invariants? Indeed it is, and just to keep things moving along, I’m going to pull it in as an identity from Eqn. B.34. Then we get:

$$\Psi_{\text{Symmetric}} = I_2 + \frac{1}{4} \left( \frac{I_1^2 - I_2}{I_3} \right) - 2 \frac{I_1}{I_3}.$$  (5.58)

Step 2: Take the invariant derivatives. This part is tedious but straightforward. As always, you can have Mathematica or Matlab do this part for you:

$$\frac{\partial \Psi_{\text{Symmetric}}}{\partial I_1} = \frac{1}{I_3} \left( \frac{I_1}{2} - 2 \right) \quad \frac{\partial^2 \Psi_{\text{Symmetric}}}{\partial I_1^2} = \frac{1}{2 I_3}$$

$$\frac{\partial \Psi_{\text{Symmetric}}}{\partial I_2} = 1 - \frac{1}{4 I_3} \quad \frac{\partial^2 \Psi_{\text{Symmetric}}}{\partial I_2^2} = 0$$

$$\frac{\partial \Psi_{\text{Symmetric}}}{\partial I_3} = \frac{1}{(I_3)^2} \left( 2 I_1 - \frac{I_1^2 - I_2}{4} \right) \quad \frac{\partial^2 \Psi_{\text{Symmetric}}}{\partial I_3^2} = \frac{1}{(I_3)^3} \left( -4 I_1 + \frac{I_1^2 - I_2}{2} \right).$$  (5.59)

Once again, I used Mathematica for this, just a minute ago. It’s not cheating.

Step 3: Plug into the chain rule. It’s not quite as clean and pretty as the other energies:

$$\text{vec} \left( \frac{\partial^2 \Psi_{\text{Symmetric}}}{\partial F^2} \right) = \frac{1}{2 I_3} g_1 g_1^T + \frac{1}{I_3} \left( \frac{I_1}{2} - 2 \right) H_1 + \left( 1 - \frac{1}{4 I_3} \right) H_2 + \frac{1}{(I_3)^2} \left( -4 I_1 + \frac{I_1^2 - I_2}{2} \right) g_3 g_3^T + \frac{1}{(I_3)^2} \left( 2 I_1 - \frac{I_1^2 - I_2}{4} \right) H_3.$$  (5.61)

But, it worked, once we got past the $\|F^{-1}\|^2$ hurdle. The system looks legit! You now have a mechanical method for computing the Hessian of any isotropic energy. You need to take some high-school-level derivatives, and maybe dig out a new identity during Step 1. Otherwise, the process is close to automatic.
Chapter 6

A Friendlier Neo-Hookean Energy

This chapter is loose retelling of §3 from Smith et al. (2018). If you want to cut to the chase and see the final energy, it’s in §6.3.3. It’s not the final energy listed in Smith et al. (2018), but it is the one that is used internally at Pixar. More details appear in a footnote 8.

6.1 Cauchy-Green vs. Smith et al. (2019)

In the previous chapter, we found that the Cauchy-Green invariants, which have been used extensively for the last 80 years, were not up to the task of generating energy Hessians. ARAP’s membership application to the Cauchy-Green Club was roundly rejected, causing all sorts of problems. Instead, we built a new, more inclusive club for ARAP: the invariants from Smith et al. (2019). Once ARAP stepped inside this new clubhouse, the simple and elegant structure that had been living inside it all along was finally revealed.

This raises a question though: where have these more inclusive invariants been for the last 80 years? Why did Mooney (1940) choose to use the unnecessarily restrictive Cauchy-Green invariants? Unfortunately, we can’t ask Melvin Mooney directly because he passed away over 52 years ago. George Green has been gone for 179 years, and Augustin-Louis Cauchy for 163, so they can’t help either. Thus, we are left to speculate. Having mulled this over for a while, I can think of two reasons.

6.1.1 Maybe Mooney Didn’t Know About the Polar Decomposition

The Smith et al. (2019) invariants are based on the polar decomposition, which has been known in math circles for more than a century. For example, Higham (1986) cites Autonne (1902). However, it may not have been widely known in Mooney’s immediate physics and rheology communities in the 1940s.

Just 15 years earlier in 1925, Werner Heisenberg had not known about basic matrix operations. While investigating quantum mechanics, he unknowingly re-derived basic
linear algebraic identities, and did not make the connection until Max Born pointed it out to him (Gribbin (2011), Chapter 6). At the time, Heisenberg was in Göttingen, Germany, the epicenter of all math and science research, not the jerkwater nowheresville of Passaic, New Jersey where Mooney was sitting.¹

Matrix notation had not even been standardized by 1940, and numerical methods were still an extremely niche discipline. The hardware to run them did not exist yet, though this would change a few years later when the Allies built massive machines to break Nazi codes. Even if these machines had existed, standard methods for computing the polar decomposition, such as those involving the SVD, would need techniques like Golub-Kahan bidiagonalization (Golub and Kahan (1965)). These were still 25 years in the future.

Thus, it would make sense that Mooney (1940) did not have the the polar decomposition at his fingertips. His paper does not list a single matrix, and is firmly a work of analysis.

6.1.2 Maybe Mooney Didn’t Care About Inversion

Alternatively, it is possible that Mooney did know about the polar decomposition, or some analytic equivalent, and decided to ignore it. To see why, we should ask which physical phenomena are captured by Smith et al. (2019) that are missing from Cauchy-Green, and whether Mooney would have cared about them.

We saw a piece of this in §5.2.4 when I let you in on the secret that nobody actually uses that weird-looking $\sqrt{\text{III}_C}$ term. Instead, everybody uses $J$. Later on in §5.3.3.2, we saw $J$ getting the recognition it deserves with a promotion to first-class invariant, $I_3 = J = \det F$.

As previously mentioned, this $I_3$ version preserves the sign of the determinant:

\[
I_3 = \sigma_x \sigma_y \sigma_z \tag{6.1}
\]

\[
\sqrt{\text{III}_C} = \sqrt{\sigma_x^2 \sigma_y^2 \sigma_z^2} = |\sigma_x \sigma_y \sigma_z|. \tag{6.2}
\]

But then why does the Cauchy-Green version wrap the $|\cdot|$ around the whole thing? Why is it so eager to throw away the sign?

A negative determinant signals that an element has inverted². As shown in Fig. 6.1, this happens if you squish a tetrahedron so hard that it pokes inside out. This sort of configuration is not possible in nature, but due to the fact that we can numerically represent phenomena that don’t occur in nature³, it can happen in the simulation.

---

¹Down the road in Murray Hill, NJ, Bell Labs would become the epicenter soon enough.

²As far as I can tell, this use of the term inversion comes from Irving et al. (2004), though they acknowledge that the concept was observed earlier (Espinosa et al. (1998)). This creates an unfortunate nomenclature collision with matrix inversion, which is a related but distinct phenomenon. Thinking of it instead as element reversal or a backwards element, or the anatomically graphic prolapsed element may help here.

³Simulation could be viewed as coercing the computer into obeying the physics of our current universe, instead of drifting into one of the myriad other ones that the math permits, but our physical laws do not.
In animation, inversions happen \textit{all the time}. We frequently squash things down to Wile-E-Coyote-style pancakes, and if we then keep squashing, we get an inversion. Thus, the energies we use need to be able to support this phenomenon.

Apparently this often happens in geometry processing as well, because one of the main strengths of the ARAP model is that it is \textit{also} able to see inversion. If an element has been squished inside out with sufficient violence, the $I_1 = \text{tr } S$ term that is so important in ARAP also becomes negative (Fig. 6.2). In contrast, the original $I_C = \text{tr } (F^T F)$ invariant squares all of its singular values, so it can never become negative.

From Mooney’s perspective, it may have made sense to use the Cauchy-Green invariants because the idea of an inversion was absurdly non-physical, so the math should preclude...
its very existence. Square everything! Nature abhors a negative invariant! He wasn’t performing any numerical computations or animating any movies, so Cauchy-Green was the way to go.\footnote{If we want to go further down this rabbit hole, Mooney never actually invokes the Cauchy-Green invariants, the follow-on analysis of Rivlin (1948) does 8 years later in Eqn. 3.4, which further muddies the provenance. I think we’ve travelled down this hole far enough for the moment though.}

\section*{6.2 \makebox[1em]{ARAP (And Others)} Don’t Do Great}

Having poured all this effort into understanding ARAP, how do the final results look when we push them into a solid mechanics simulation? To answer that, we should first establish what we want out of a solid mechanics simulation. Our target application is computer animation, and expressively realistic \textit{squashing and stretching} are some of the most important visual features, according to the Disney animators who wrote Johnston and Thomas (1981) and Blair (2003).

Let’s look at what we expect under squashing in stretching. In Fig. 6.3, we have a square of bubble gum, held between two metal plates. Under squashing, we expect that stuff will pooch out of the top and bottom, and under stretching, we expect it to form a graceful curve that flattens out to a line along the middle.

\begin{figure}[h]
\centering
\includegraphics[width=3in]{fig6.3.png}
\caption{What we want from a 2D square of rubber (left). Under squashing, stuff squeezes out the top and bottom (center). Under stretching, it swoops down to a nice clean line along the middle (right).}
\end{figure}

What happens when we try to subject ARAP to these conditions? The results shown in Fig. 6.4 aren’t great. This partially explains why ARAP is popular in \textit{geometry processing}, but less so in simulation. While it’s relatively robust\footnote{Provided you compute the exact rotation gradient!}, it’s not very realistic, and never pretended to be. If we look at the energy closely, \( \Psi_{ARAP} = I_2 - 2I_1 - 3 \), we see that it...
only involves the first and second invariants. However, we saw back in §5.3.2 and Fig. 5.4, that the third invariant is the one that deals with volume.

The ARAP energy does not explicitly perform any volume preservation, because it doesn’t even bother to compute the volume of each element. Instead, it only cares about preserving relative lengths. That’s why Fig. 6.4 looks like a length of chickenwire being stretched and squashed.

6.2.1 A Brief Aside: The Lamé Parameters

The ARAP model only preserves lengths, not volumes. Thus, we didn’t need parameters describing how much relative length and volume preservation we wanted. But, we’re going to need such parameters in a moment, so let’s describe them now.

These are usually specified using the Lamé parameters, $\mu$ and $\lambda$. The $\mu$ parameter controls length preservation, and is also sometimes called the shear modulus or Lamé’s second parameter. The $\lambda$ parameter controls volume preservation, and is sometimes called Lamé’s first parameter.\(^6\) If you want lots of volume preservation, you set $\lambda$ to be much larger than $\mu$. Conversely, if you want more length preservation, you push up the value of $\mu$.

Just to make your life more difficult, many simulators, including Fizt, don’t take in values for $\mu$ and $\lambda$ directly. Instead, they accept an equivalent set of parameters: the Young’s modulus $E$, and Poisson’s ratio, $\nu$. The conversion from $(E, \nu)$ and $(\mu, \lambda)$ is:

$$\mu = \frac{E}{2(1+\nu)} \quad (6.3)$$

$$\lambda = \frac{E\nu}{(1+\nu)(1-2\nu)}. \quad (6.4)$$

Hold on. When $\nu = \frac{1}{2}$, the value of $\lambda$ creates a divide-by-zero that blasts off to infinity? Is this a mistake?

---

\(^6\)I know. The $\lambda$ sets up a naming collision with the eigenvalues $\lambda_i$. I don’t make the rules here, but it should be clear from context which one I mean.
It’s correct: specifying $\nu = \frac{1}{2}$ means that volume must be preserved at all costs, no matter how much it ruins the stability of the simulation. Biological tissues like muscle and fat are known to have $\nu \approx \frac{1}{2}$ (Greaves et al. (2011)), but they’re not exactly $\frac{1}{2}$, so we can still chug ahead.

### 6.2.2 St. Venant Kirchhoff Doesn’t Do Better

With the Lamé preliminaries out of the way, let’s look at St. Venant Kirchhoff to see how it compares to ARAP. We’re moving past the stretching-only version and finally adding the “volume preservation” term to the right to arrive at the full model:

$$\Psi_{StVK} = \mu \lVert E \rVert_F^2 + \frac{\lambda}{2} \text{tr}^2(E).$$ (6.5)

Since we care how fleshy our cartoon characters look, let’s try this out with $\nu = 0.49$ in Fig. 6.5. First the good news: the stretching looks better than ARAP. At least it bows a little in the middle, even though it doesn’t give us the clean line we want along the middle. But what is going on under squashing?! It poofs up slightly along the top and bottom, but then some bizarre overlap and collapse happens along the sides!

Figure 6.5.: StVK, under squashing (middle) and stretching (right). What is going on with squashing?!

---

7If you’re a stickler, the Lamé conversion formulas should be slightly different for StVK because it’s a quartic model. We’re going to abandon StVK as a whole in a minute though, so I’m not going to spend time on that.
What we’re seeing is that StVK is a card-carrying member of the Cauchy-Green Club:

\[ \Psi_{\text{StVK}} = \mu \|E\|^2_F + \frac{\lambda}{2} \text{tr}^2(E) \]

\[ = \frac{\mu}{4} \|E\|^2_F + \frac{\lambda}{8} \text{tr}^2(F^T F - I) \quad \text{(Definition of E, and pull out the } \frac{1}{2} \text{)} \]

\[ = \frac{\mu}{4} \|E\|^2_F + \frac{\lambda}{8} \text{tr}(F^T F - I) \text{tr}(F^T F - I) \quad \text{(Make the squaring explicit)} \]

\[ = \frac{\mu}{4} \|E\|^2_F + \frac{\lambda}{8} \left( \text{tr}(F^T F)^2 - 6 \text{tr}(F^T F) + 9 \right) \quad \text{(Eqn. B.13 and tr I = 3)} \]

\[ = \frac{\mu}{4} \|E\|^2_F + \frac{\lambda}{8} \left( I_C^2 - 6I_C + 9 \right) \quad \text{(Definition of } I_C \text{)} \]

\[ = \frac{\mu}{4} \left( I_C^2 - 2I_C + 3 \right) + \frac{\lambda}{8} \left( I_C^2 - 6I_C + 9 \right). \quad \text{(From Eqn. 5.10)} \]

As we just saw in §6.1.2, the Cauchy-Green invariants don’t know about inversion. In fact, they deliberately put on blinders by squaring it away. Thus, the bizarrely collapsed and overlapping elements in Fig. 6.5 are elements that have inverted. StVK thinks they’re not inverted, due to its Cauchy-Green myopia, so it does not exert any forces that try to correct things.

Thus, StVK makes some progress under stretching, but this behavior under squashing simply won’t do.

6.2.3 Co-Rotational Doesn’t Do Better

Next up, let’s try that co-rotational model that everybody (Müller et al. (2002); Etzmuss et al. (2003); Irving et al. (2004)) simultaneously re-discovered back in the mid-2000s:

\[ \Psi_{\text{CoRot}} = \mu \|F - R\|^2_F + \frac{\lambda}{2} \text{tr}^2(S - I). \quad (6.6) \]

As you can see in Fig. 6.6, things get really weird. The freakiness around squashing we saw with StVK is gone, but now it’s replaced with bigger freakishness under stretching! Why did the slab turn into a prolapsed trampoline!?

Let’s unpack that so-called “volumepreserving” term. Following the same steps as StVK, we get:

\[ \Psi_{\text{CoRot}} = \mu \|F - R\|^2_F + \frac{\lambda}{2} \left( I_C^2 - 6I_C + 9 \right). \quad (6.7) \]

What a minute ... where’s \( I_3 \)? What kind of a shady volume term doesn’t even compute the volume? A \textit{linearized} volume term, that’s what kind. That won’t do here.
Figure 6.6.: Co-rotational, with increasing $\nu$. As $\nu \to \frac{1}{2}$, stretching gets really strange.

### 6.2.4 Neo-Hookean Is Okay Unless It Burns The House Down

Finally, let’s try the Neo-Hookean material, which we can re-write in terms of the Smith et al. (2019) invariants:

$$
\Psi_{BW08} = \frac{\mu}{2}(\|F\|_F^2 - 3) - \mu \log(J) + \frac{\lambda}{2} (\log(J))^2
$$

$$
= \frac{\mu}{2}(I_2 - 3) - \mu \log(I_3) + \frac{\lambda}{2} (\log(I_3))^2.
$$
The $I_3$ invariant shows up for sure, so at least we’re not getting suckered into using some sham volume preservation term that doesn’t even know the definition of “volume”.

The results of the simulation are in Fig. 6.7. First the good news: under stretching, Neo-Hookean finally gives us that nice, stretched-out bubble gum look that we wanted. Then then bad: under squashing, Neo-Hookean produces a NaN that burns down the rest of the simulation.

Figure 6.7.: Neo-Hookean material looks great under stretching (right), but produces a NaN and explodes under squashing.

The culprit is that $\log(I_3)$ term, since as $I_3 \to 0$, we get the singularity $\log(I_3) \to -\infty$. It’s actually slightly worse, since there’s a squared $(\log(J))^2$ term that blasts off even more rapidly towards infinity.

Now our speculative motivations from §6.1.2 about Mooney using the squared invariants rears its ugly head. Under this interpretation, nature abhors a negative invariant, so a $\log(I_3)$ term has been inserted as a barrier function to ensure that we never cross the $I_3 = 0$ threshold. In this world view, we abhor a negative invariant so much that we give the energy permission to burn down the entire simulation if it looks like a negative is about to appear.

Computer animation does not ascribe to this view. Again, Wile-E-Coyote-style pancakes happen all the time. They don’t persist very long; after the initial impact, the Road Runner’s anvil usually falls away, and the coyote blows into his thumb to re-inflate himself. A good energy in this scenario should allow pancakes and inversions to occur, but once the anvil forces are removed, sproing the coyote back into his original shape.

6.3 A Better Neo-Hookean Energy?

6.3.1 So Many Neo-Hookeans

We looked at one Neo-Hookean energy in §6.2.4, but there are a whole lot of energies that call themselves “Neo-Hookean”. We even saw one way back in Eqn. 2.6 which was just the Dirichlet energy in disguise. If Fig. 6.7, we saw that we can get good-looking
stretching shapes out of Neo-Hookean, but it explodes under squashing. Maybe one of these other energies that calls itself “Neo-Hookean” can do better?

Here’s a few I found in the literature:

\[
\Psi_A = \frac{\mu}{2}(I_2 - 3) - \mu \log I_3 + \frac{\lambda}{2}(I_3 - 1)^2 \quad (\text{Ogden (1997)})
\]

\[
\Psi_B = \frac{\mu}{2} \left( \frac{I_2}{I_3^2} - 3 \right) + \frac{\lambda}{2}(I_3 - 1)^2 \quad (\text{Bower (2009)})
\]

\[
\Psi_C = \frac{\mu}{2} \left( \frac{I_2}{I_3^2} - 3 \right) + \frac{\lambda}{2}(I_3 - 1). \quad (\text{Wang and Yang (2016)})
\]

We saw before that any \(\log(I_3)\) will create an infinity under pancake states, so \(\Psi_A\) is already out of the running. The \(\Psi_B\) and \(\Psi_C\) energies don’t contain a \(\log(I_3)\), but they instead have \(\frac{I_2}{I_3^2}\) terms. These terms have the same problem: as \(I_3 \to 0\), the \(\frac{I_2}{I_3^2}\) is standing there with a match and a can of gasoline, ready to burn everything down with an \(\infty\). That’s not helpful either.

### 6.3.2 Let’s Mix-And-Match Our Own

Still, if we look at \(\Psi_A\), \(\Psi_B\), and \(\Psi_C\), we can see some glimmers of hope. That \(\frac{\mu}{2}(I_2 - 3)\) in front of \(\Psi_A\), which also appears in front of \(\Psi_{BW08}\), is a well-behaved Dirichlet-like term. It doesn’t give us the behavior we want, but at least it will never generate a simulation-destroying singularity.

All three of the energies also incorporate a \(\frac{\lambda}{2}(I_3 - 1)^2\) term. This is well-behaved as well! It can never create a divide-by-zero error, and unlike the co-rotational model, it uses the 100% certified, honest-to-goodness volume-measuring, \(I_3\) invariant. Maybe we can just mix-and-match these two components to get our own Neo-Hookean flavor:

\[
\Psi_D = \frac{\mu}{2}(I_2 - 3) + \frac{\lambda}{2}(I_3 - 1)^2.
\]

This looks quite reasonable, doesn’t it? Until you try to simulate it. In Fig. 6.8, we see what happens at the first step of the simulation, even when no forces have been applied. The mesh shrinks! The artifact becomes less pronounced as \(\nu \to \frac{1}{2}\), but is still clearly visible. Other biomechanics works have observed similar phenomena (e.g. Blemker et al. (2005)), though not in the context of authoring an inversion-friendly Neo-Hookean energy. The solution is usually to set \(\lambda \approx 1000\mu\), which corresponds to ignoring the artifact by always keeping \(\nu\) really close to \(\frac{1}{2}\). Let’s try to do better.

### 6.3.3 A Stable Neo-Hookean Energy

Why does the mesh shrink to begin with? First, it has a Dirichlet-like term, \(\frac{\mu}{2}(I_2 - 3)\), which we first saw in §2.3.4 when it tried to collapse our entire mesh down to a point.
Figure 6.8: $\Psi_D$, our first attempt at a mix-and-match Neo-Hookean energy, freakishly causes the mesh to shrink, even if no forces are being applied. As $\nu$ decreases, the problem gets worse, but it visibly persists even as $\nu \approx \frac{1}{2}$.

Second, it has a volume preservation term, $\frac{\lambda}{2}(I_3 - 1)^2$, which tries to keep $I_3$ as close to the original volume (which corresponds to 1) as possible. Even in the absence of any external forces, these two terms are pulling in opposition to each other, and the true “rest-state” that a mesh will have is the middle ground where these two forces cancel out.

This cancellation point will not be when $I_3 = 1$. It will instead be at some compromise point with the dueling Dirichlet force, which wants to pull everything down into the zero-volume $I_2 = 0$ state. In this duel, if we increase $\lambda$, then $\frac{\lambda}{2}(I_3 - 1)^2$ gains the upper hand and the artifact become smaller.

Here’s an idea: why don’t we give the $\frac{\lambda}{2}(I_3 - 1)^2$ upper hand by making the element fatter? Instead of trying to get back to the $I_3 = 1$ state, why don’t we tell the energy to return to some $I_3 = \alpha$ state, where $\alpha > 1$? Then when it has to compromise with the Dirichlet term, the slightly shrunken version that they balance out to will be the $I_3 = 1$ state we originally wanted?

We can write this out in more detail by taking the PK1 of $\Psi_D$:

$$\frac{\partial\Psi_D}{\partial\mathbf{F}} = \mu \mathbf{F} - \lambda (I_3 - 1) \frac{\partial I_3}{\partial \mathbf{F}}$$

$$= \mu \mathbf{F} - \lambda (I_3 - 1) \begin{bmatrix} f_{11} & -f_{10} \\ -f_{01} & f_{00} \end{bmatrix}.$$  \hspace{1cm} \text{(In 2D)}$$

If everything was working as it should the PK1 would be all zeros under zero deformation,
\[ F = I, \text{ but we see that it doesn't work out that way:} \]
\[
\frac{\partial \Psi_D(I)}{\partial F} = \mu I - \lambda (1 - 1) I \\
= \mu I.
\]

That’s not great, but we knew it was going to happen. Now let’s introduce our \( \alpha \) factor, and see how much chubbier we need to make each element:

\[
\frac{\partial \Psi_\alpha}{\partial F} = \mu F - \lambda (I_3 - \alpha) \left[ \begin{array}{cc} f_{11} & -f_{10} \\ -f_{01} & f_{00} \end{array} \right] \\
\left[ \begin{array}{cc} 0 & 0 \\ 0 & 0 \end{array} \right] = \mu I - \lambda (1 - \alpha) I.
\]

(Plugging in \( F = I, \frac{\partial \Psi_\alpha}{\partial F} = 0 \))

The last line encodes the constraint \( 0 = \mu - \lambda (1 - \alpha) \). Solving for \( \alpha \) yields:

\( \alpha = 1 - \frac{\mu}{\lambda} \). (6.8)

We have not seen the \( \frac{\partial \Psi}{\partial F} = 0 \) explicitly named anywhere, so in Smith et al. (2019) we decided to call it the “rest-stability” criterion. Our rest-stable Neo-Hookean energy is then:

\[
\Psi_{SNH} = \frac{\mu}{2} (I_2 - 3) + \frac{\lambda}{2} (I_3 - \alpha)^2 \\
= \frac{\mu}{2} (I_2 - 3) + \frac{\lambda}{2} \left( I_3 - 1 + \frac{\mu}{\lambda} \right)^2.
\]

Expanding the square, we get:

\[
\Psi_{SNH} = \frac{\mu}{2} (I_2 - 3) - \mu (I_3 - 1) + \frac{\lambda}{2} (I_3 - 1)^2 + \left( \frac{\mu}{\lambda} \right)^2.
\]

Constant factors only translate the overall energy and get burned off under differentiation, so this is equivalent to:

\[
\Psi_{SNH} = \frac{\mu}{2} (I_2 - 3) - \mu (I_3 - 1) + \frac{\lambda}{2} (I_3 - 1)^2. \quad (6.9)
\]

This is our final Stable Neo-Hookean (SNH) energy.\(^8\)

\(^8\)The Smith et al. (2019) paper also adds an origin barrier term that further complicates things. That version is *not the one* that is used internally at Pixar. The \( \Psi_{SNH} \) energy is. The origin barrier was added because the paper’s referees insisted that a spurious but extremely-difficult-to-reach root at \( F = 0 \) would destabilize the energy. Extensive production experience has shown that it does not.
6.4 A Bunch of Other Stable Energies

We can apply this approach more generally. Just to drive you crazy, a bunch of other energies are usually written down without any volume preservation term. The assumption is that you will somehow add your own. For example, the two-term Mooney Rivlin model can be written:

$$\Psi_{MR} = \mu_0(I_C - 3) + \frac{\mu_1}{2}(I_C^2 - II_C - 3).$$

If you try to simulate this, you get the collapsed mesh on the left of Fig. 6.9. Clearly, a volume term is needed. We could be stubborn, insist that we learned nothing from the previous section, and just patch on a $(I_3 - 1)^2$ to obtain:

$$\Psi_{MR^+} = \mu_0(I_C - 3) + \frac{\mu_1}{2}(I_C^2 - II_C - 3) + \frac{\lambda}{2}(I_3 - 1)^2.$$

Predictably, we then encounter the same mesh contraction problems, as seen on the right side of Fig. 6.9. We can do better.

Figure 6.9.: Mooney-Rivlin, without a volume term (left), and with a volume term, but without rest stability (right). Same contraction problems as before: the mesh shrinks, even though no forces are being applied.

6.4.1 Stable Mooney-Rivlin

I’m being slightly abusive here and mixing the Cauchy-Green and Smith et al. (2019) invariants. But, as we saw in §5.3.3.2, the $II_C$ term gets a lot bigger when written in terms of the new invariants. Here’s the Stable Mooney-Rivlin energy,

$$\Psi_{SMR} = \mu_0(I_C - 3) + \frac{\mu_1}{2}(I_C^2 - II_C - 3) + \frac{\lambda}{2}(I_3 - \alpha)^2,$$

and the task is now to solve for $\alpha$ in the PK1 when $F = I$. The PK1 is:

$$\frac{\partial \Psi_{SMR}}{\partial F} = 2\mu_0 F + \mu_1 I_C F - 2\mu_1 F F^T F + \lambda(I_3 - \alpha) \begin{bmatrix} f_{11} & -f_{10} \\ -f_{01} & f_{00} \end{bmatrix}.$$
6. A Friendlier Neo-Hookean Energy

Pushing through, the alpha works out to:

\[ 0 = 2\mu_0 I + 3\mu_1 I - 2\mu_1 I + \lambda (1 - \alpha) I \]
\[ 2\mu_0 I + \mu_1 I + \lambda (1 - \alpha) I \]
\[ \alpha = \frac{2\mu_0 + \mu_1}{\lambda} + 1 \]

The final Stable Mooney-Rivlin energy takes on a form similar to Stable Neo-Hookean:

\[
\Psi_{\text{SMR}} = \mu_0 (I_C - 3) + \frac{\mu_1}{2} (I_C^2 - 3) + \frac{\lambda}{2} \left( I_3 - \frac{2\mu_0 + \mu_1}{\lambda} - 1 \right)^2 \tag{6.10}
\]

\[
= \mu_0 (I_C - 3) + \frac{\mu_1}{2} (I_C^2 - 3) + \frac{\lambda}{2} (I_3 - 1)^2 - (2\mu_0 + \mu_1)(I_3 - 1). \tag{6.11}
\]

The new terms are shown in red.

6.4.2 Stable Arruda-Boyce

Let’s do it again for the energy of Arruda and Boyce (1993). A three-term version of this is:

\[
\Psi_{\text{AB}} = \mu (I_2 - 3) + \frac{\mu_1}{4} (I_2^2 - 9) + \frac{\mu_2}{6} (I_2^3 - 27).
\]

Our stable version is,

\[
\Psi_{\text{SAB}} = \mu (I_2 - 3) + \frac{\mu_1}{4} (I_2^2 - 9) + \frac{\mu_2}{6} (I_2^3 - 27) + \frac{\lambda}{2} (I_3 - \alpha)^2,
\]

and after obtaining \( \alpha = 1 + \frac{\mu}{\lambda}(1 + 3\beta_1 + 9\beta_2) \), the final version is

\[
\Psi_{\text{SAB}} = \mu (I_2 - 3) + \frac{\mu_1}{4} (I_2^2 - 9) + \frac{\mu_2}{6} (I_2^3 - 27) + \frac{\lambda}{2} \left( I_3 - 1 - \frac{\mu}{\lambda}(1 + 3\beta_1 + 9\beta_2) \right)^2 \tag{6.12}
\]

\[
= \mu (I_2 - 3) + \frac{\mu_1}{4} (I_2^2 - 9) + \frac{\mu_2}{6} (I_2^3 - 27) + \frac{\lambda}{2} (I_3 - 1)^2 - \mu(1 + 3\beta_1 + 9\beta_2)(I_3 - 1). \tag{6.13}
\]

Again, the new terms are in red.

6.4.3 Stable Fung Hardening

Finally, let’s look at the hardening model of Fung (2013). This energy usually doesn’t appear all by itself, but is added so that a material becomes dramatically stiffer (hardens) under large stretching. Here we append it to our Stable Neo-Hookean model,

\[
\Psi_{\text{SNH+Fung}} = \frac{\mu_0}{2} (I_2 - 3) + \frac{\lambda}{2} (I_3 - \alpha)^2 + \frac{\gamma}{2} \left( e^{\frac{\mu_0}{2} (I_2 - 3)} - 1 \right),
\]
and after solving for \( \alpha = 1 + \frac{\mu_0 + \gamma \mu_1}{\lambda} \), we obtain

\[
\Psi_{SNH+Fung} = \frac{\mu_0}{2} (I_2 - 3) + \frac{\lambda}{2} \left( I_3 - 1 - \frac{\mu_0 + \gamma \mu_1}{\lambda} \right)^2 + \frac{\gamma}{2} \left( e^{\frac{\mu_1}{2} (I_2 - 3)} - 1 \right) \tag{6.14}
\]

\[
= \frac{\mu_0}{2} (I_2 - 3) + \frac{\lambda}{2} (I_3 - 1)^2 + \frac{\gamma}{2} \left( e^{\frac{\mu_1}{2} (I_2 - 3)} - 1 \right) - (\mu_0 + \gamma \mu_1)(I_3 - 1). \tag{6.15}
\]

**In general:** When obtaining the rest stabilization term \( \alpha \), the expressions tend to take the form \( \alpha = 1 + \beta \). When inserted into \((I_3 - \alpha)^2\), these then separate (up to constant) into the original \( \frac{\lambda}{2} (I_3 - 1)^2 \) volume term, and an energy-specific stabilization term \(-\beta(I_3 - 1)\).

Go forth and design your own stable energy!
Chapter 7

The Analytic Eigensystems of Isotropic Energies

This chapter is a loose retelling of §4 from Smith et al. (2018) and §4 and §5 in Smith et al. (2019). If you want to cut to the chase, Matlab code that implements the final algorithm is given in Fig. 7.7. Plug your favorite isotropic energy into line 11, and it will spit out the analytic eigenvalues.

7.1 Keeping Everything Semi-Positive-Definite

The core of Fitz and Baraff and Witkin (1998)-style integration is the use of a preconditioned conjugate gradients (PCG) solver. PCG requires that the underlying matrix \( A \) is semi-positive-definite, i.e. that all of its eigenvalues are greater than or equal to zero. This is usually written somewhat obtusely in linear algebra textbooks as:

\[
\mathbf{x}^T \mathbf{A} \mathbf{x} \geq 0 \quad \forall \mathbf{x}.
\]

Another way to look at this is through the product \( \mathbf{A} \mathbf{x} = \mathbf{y} \). Positive-definiteness implies that the sign of each entry in \( \mathbf{y} \) must always match the corresponding entry in \( \mathbf{x} \). The matrix \( \mathbf{A} \) is not allowed to flip the sign of any entry in \( \mathbf{x} \), no matter what the entries in \( \mathbf{x} \) are. If it does, when we compute the dot product \( \mathbf{x}^T \mathbf{y} \), the negatives of \( \mathbf{x} \) and \( \mathbf{y} \) will not mutually annihilate, and you can end up with a result less than zero.

We care about this property because if the system isn’t semi-positive-definite, it implies the existence of multiple, physically valid, energetically equivalent solutions. In that case, the optimization will not know which solution to pick, and can oscillate indecisively between them forever, or explode spectacularly to infinity. This ambiguity can and does happen in the physical systems we are examining, but let’s put that fact aside for the moment. In the moment, we want to be able to hand to our existing Fitz solver a semi-positive-definite system. How can we know for sure that all the Hessians that came out of all of our \( \frac{\partial \Psi}{\partial \mathbf{F}} \) and \( \frac{\partial \Psi}{\partial \mathbf{x}} \) manipulations produced all-positive eigenvalues?
One way would be to compute the eigendecomposition of the global system matrix you assembled using the $\frac{\partial^2 \Psi}{\partial F^2}$ from every element in your simulation (Nocedal and Wright (2006)). If any of those eigenvalues are less than zero, you can perform a projection to snap those back to zero, reassemble a modified matrix, and then perform PCG using this modified matrix. We would prefer not to do this, because PCG runs in roughly $O(N^2)$ time, but an eigendecomposition of the global matrix takes $O(N^3)$. The cure is worse than the disease. We can instead try a per-element projection in the style of Teran et al. (2005), where we compute a eigendecomposition for each element’s $\frac{\partial^2 \Psi}{\partial F^2}$, which only involves taking the eigendecomposition of a bunch of $9 \times 9$ matrices, not the global $N \times N$ matrix. The sum of semi-positive-definite matrices is known to also be semi-positive definite, so the strategy is sound.

7.2 Can ARAP Go Indefinite?

Maybe we’re worrying about nothing, and the energies we’re interested in can’t even go indefinite. To see if this is true, let’s look at the ARAP energy.

Fortunately, the slick representation we found for the rotation gradient $\frac{\partial R}{\partial F}$ already gives analytic expressions for its own eigendecomposition. From there, it’s only a short skip and jump to obtain the analytic eigenvalues of the entire ARAP energy. From §5.5.2, the flattened Hessian for ARAP is:

$$\text{vec} \left( \frac{\partial^2 \Psi_{\text{ARAP}}}{\partial F^2} \right) = 2I_{9 \times 9} - 2H_1.$$ 

We know what the eigendecomposition of $2I_{9 \times 9}$ is. All the eigenvalues are $\lambda_{0...8} = 2$, and since the eigenvalues are all the same, the eigenvectors are not unique. Any basis than spans a rank-9 subspace will do.

We also know what the eigensystem of $H_1$ from §5.4.4. Here they are again:

$$\lambda_0 = \frac{2}{\sigma_x + \sigma_y} \quad Q_0 = \frac{1}{\sqrt{2}} U \begin{bmatrix} 0 & -1 & 0 \\ 1 & 0 & 0 \\ 0 & 0 & 0 \end{bmatrix} V^T$$

$$\lambda_1 = \frac{2}{\sigma_y + \sigma_z} \quad Q_1 = \frac{1}{\sqrt{2}} U \begin{bmatrix} 0 & 0 & 0 \\ 0 & 0 & 1 \\ 0 & -1 & 0 \end{bmatrix} V^T$$

$$\lambda_2 = \frac{2}{\sigma_x + \sigma_z} \quad Q_2 = \frac{1}{\sqrt{2}} U \begin{bmatrix} 0 & 0 & 1 \\ 0 & 0 & 0 \\ -1 & 0 & 0 \end{bmatrix} V^T$$

$$\lambda_{3...8} = 0 \quad Q_{3...8} = \text{subspace orthogonal to } Q_{0,1,2}.$$
Multiplying a matrix by a constant only scales its eigenvalues, so $-2\mathbf{H}_1$ has the eigenvalues

$$
\begin{align*}
\lambda_0 &= \frac{-4}{\sigma_x + \sigma_y} \\
\lambda_1 &= \frac{-4}{\sigma_y + \sigma_z} \\
\lambda_2 &= \frac{-4}{\sigma_x + \sigma_z} \\
\lambda_{3\ldots8} &= 0,
\end{align*}
$$

and the eigenvectors remain the same.

So, what’s the eigensystem of their sum, $\mathbf{I}_{9\times9} - 2\mathbf{H}_1$? Unfortunately, there is no magic theorem that takes the eigendecomposition of two matrices and tells you the eigendecomposition of their sum.

Unless if one of the matrices has a special structure, like $\mathbf{I}_{9\times9}$! In that case, you can just add the eigenvalues from the diagonal matrix to the eigenvalues of your second matrix. The analytic eigendecomposition of the ARAP energy is then:

$$
\begin{align*}
\lambda_0 &= 2 - \frac{4}{\sigma_x + \sigma_y} & Q_0 &= \frac{1}{\sqrt{2}} \begin{bmatrix} 0 & -1 & 0 \\ 1 & 0 & 0 \\ 0 & 0 & 0 \end{bmatrix} \mathbf{V}^T \\
\lambda_1 &= 2 - \frac{4}{\sigma_y + \sigma_z} & Q_1 &= \frac{1}{\sqrt{2}} \begin{bmatrix} 0 & 0 & 0 \\ 0 & 0 & 1 \\ 0 & -1 & 0 \end{bmatrix} \mathbf{V}^T \\
\lambda_2 &= 2 - \frac{4}{\sigma_x + \sigma_z} & Q_2 &= \frac{1}{\sqrt{2}} \begin{bmatrix} 0 & 0 & 1 \\ 0 & 0 & 0 \\ -1 & 0 & 0 \end{bmatrix} \mathbf{V}^T \\
\lambda_{3\ldots8} &= 2 \\
\mathbf{Q}_{4\ldots8} &= \text{subspace orthogonal to } Q_{0,1,2}.
\end{align*}
$$

First of all, who knew that ARAP’s eigendecomposition had such crisp expressions? I’m used to seeing the Hessian as a chaotic churn of numbers, and taking its eigendecomposition just plunges you further into the primordial soup. But, this crystalline eigenstructure was sitting there at the bottom this whole time! A pleasant surprise.

Second, and returning to the task at hand, can the energy go indefinite? Certainly $\lambda_{3\ldots8}$ are all safely positive for all time, but when do $\lambda_{0,1,2}$ go negative? Let’s look at $\lambda_0$ as an example:

$$
\begin{align*}
\lambda_0 &= 2 - \frac{4}{\sigma_x + \sigma_y} \\
2 &\leq \frac{4}{\sigma_x + \sigma_y} \\
\sigma_x + \sigma_y &\leq 2.
\end{align*}
$$

Similar expressions appear for the other two:

$$
\begin{align*}
\lambda_1 &\leq 0 \iff \sigma_y + \sigma_z \leq 2 \\
\lambda_2 &\leq 0 \iff \sigma_x + \sigma_z \leq 2.
\end{align*}
$$
Figure 7.1: From left to right: An undeformed element is already semi-definite. Squashing along \( y \) makes it indefinite. Stretching enough in \( x \) makes it semi-definite again. Squashing \( y \) even further pushes us back into indefiniteness.

Uh oh. Don’t these conditions get tripped all the time? Let’s put a single element through its paces in Fig. 7.1. If an element is undergoing zero deformation, \( \sigma_x = \sigma_y = 1 \), it is already flirting with disaster, as \( \lambda_0 = 0 \). If any of these directions gets squashed, e.g. \( \lambda_0 = \frac{1}{2} \) while \( \lambda_1 = 1 \), then this eigenvalue will be negative. The other direction could pick up the slack, i.e. \( \lambda_0 = \frac{1}{2} \) while \( \lambda_1 = \frac{3}{2} \), but if any direction is more squashed than the other direction is stretched, then the eigenvalue will become negative. The only situation where \( \lambda_0 \) is safely positive is if both directions are being stretched.

The appearance of negative eigenvalues under squashing is well-known in mechanical engineering, as it corresponds to the onset of buckling, as shown in Fig. 7.2. If you squash a bar of material, will it buckle up or down? From a deformation energy standpoint, both configurations will give the exact same score for \( \Psi_{\text{ARAP}} \). Therefore, we are currently stuck between multiple, equally enticing solutions, which corresponds to a saddle point in the energy. Saddle points possess both positive and negative curvature, a.k.a. indefinite Hessians, and so the negative eigenvalues make sense.

Figure 7.2: From left to right: An undeformed bar. We squash it. Which direction should it buckle in? Up or down? As far as the deformation energy goes, both have exactly the same score!
So yes: ARAP can go indefinite. However, we now also know its eigensystem in closed form, so we can devise a projected Hessian quite easily. If we detect that any of the eigenvalues are less than zero, we can efficiently snap it back to zero. It works out to a few extra lines on top of ARAP_Hessian from Fig. 5.10, which we’ll call ARAP_Hessian_Filtered and give the code for in Fig. 7.3.

We didn’t have to numerically compute a $9 \times 9$ eigendecomposition anywhere, which is both faster and cleaner. Additionally, we got better intuition of when and why elements go indefinite, and some fast, compact code for projected Hessians. Wouldn’t it be nice if we could do this for every energy?

7.3 The Eigendecompositions of Arbitrary Energies

An analytic decomposition is not only available for the ARAP energy, it can be obtained for any isotropic energy. It will take a little bit of work to find a generic expression, but once we do, I’ll bake everything into a Matlab script for you (Fig. 7.5)

7.3.1 The General Eigensystem of $I_3$

We’re actually $2/3$ of the way to a generic approach already. We have the analytic eigensystem for $\frac{\partial^2 I_3}{\partial F^2}$, which was given in §5.4.4 and §7.2. We also have the eigensystem of $\frac{\partial^2 I_2}{\partial F^2}$, which was easy to get, because it’s a diagonal matrix.

If we can get the analytic eigensystem for $\frac{\partial^2 I_3}{\partial F^2}$, then maybe, just maybe, we can combine the eigendecompositions for all three in some way to arrive at a generic approach. If you’re an analysis rockstar, you can probably derive the eigensystem by taking the variational derivative of $I_3$ or using some related principle.

I am not an analysis rockstar, so this is not what I did. Just like in §5.4.2, I messed around in Matlab with some easy cases where $U = V = I$ and $\Sigma$ was loaded up with easy integers, until I found something that looked like a pattern, and then verified their correctness symbolically using Mathematica. Not the most straight-line route to finding analytic expressions, but there’s more than one way to skin a cat.
function [H] = ARAP_Hessian_Filtered(F)

[U Sigma V] = svd_rv(F);

% get the twist modes
T0 = [0 -1 0; 1 0 0; 0 0 0];
T0 = (1 / sqrt(2)) * U * T0 * V';

T1 = [0 0 0; 0 0 1; 0 -1 0];
T1 = (1 / sqrt(2)) * U * T1 * V';

T2 = [0 0 1; 0 0 0; -1 0 0];
T2 = (1 / sqrt(2)) * U * T2 * V';

% get the flattened versions
t0 = vec(T0);
t1 = vec(T1);
t2 = vec(T2);

% get the singular values in an order that is consistent
% with the numbering in the paper
s0 = Sigma(1,1);
s1 = Sigma(2,2);
s2 = Sigma(3,3);

% build the filter the non-trivial eigenvalues
lambda0 = 2 / (s0 + s1);
lambda1 = 2 / (s1 + s2);
lambda2 = 2 / (s0 + s2);
if (s0 + s1 < 2)
    lambda0 = 1;
end
if (s1 + s2 < 2)
    lambda1 = 1;
end
if (s0 + s2 < 2)
    lambda2 = 1;
end

H = eye(9,9);
H = H - lambda0 * (t0 * t0');
H = H - lambda1 * (t1 * t1');
H = H - lambda2 * (t2 * t2');
H = 2 * H;
end

Figure 7.3.: Matlab code to compute the projected Hessian of the ARAP energy in 3D. Any eigenvalue that is less than zero is snapped back to zero. The lambda terms are not set to zero directly, because they still need to cancel off the entries of eye(9,9) when they are added to H.
Weirdly enough, three of the eigenvectors of $\frac{\partial^2 I_3}{\partial F^2}$ are exactly the same as $\frac{\partial^2 I_1}{\partial F^2}$,

$$\lambda_3 = \sigma_z \quad Q_3 = \frac{1}{\sqrt{2}} U \begin{bmatrix} 0 & -1 & 0 \\ 1 & 0 & 0 \\ 0 & 0 & 0 \end{bmatrix} V^T,$$

$$\lambda_4 = \sigma_x \quad Q_4 = \frac{1}{\sqrt{2}} U \begin{bmatrix} 0 & 0 & 0 \\ 0 & 0 & 1 \\ 0 & -1 & 0 \end{bmatrix} V^T,$$

$$\lambda_5 = \sigma_y \quad Q_5 = \frac{1}{\sqrt{2}} U \begin{bmatrix} 0 & 0 & 1 \\ 0 & 0 & 0 \\ -1 & 0 & 0 \end{bmatrix} V^T,$$

and the eigenvalues are even simpler: just the singular values $\sigma_{\{x,y,z\}}$. Just like we saw with Eqn. 5.40, these correspond to infinitesimal rotations, so we call these the \textit{twist} matrices. The next three eigenpairs are so similar to the first three that you’re going to think I typed them in wrong:

$$\lambda_6 = -\sigma_z \quad Q_6 = \frac{1}{\sqrt{2}} U \begin{bmatrix} 0 & 1 & 0 \\ 1 & 0 & 0 \\ 0 & 0 & 0 \end{bmatrix} V^T,$$

$$\lambda_7 = -\sigma_x \quad Q_7 = \frac{1}{\sqrt{2}} U \begin{bmatrix} 0 & 0 & 0 \\ 0 & 0 & 1 \\ 0 & 1 & 0 \end{bmatrix} V^T,$$

$$\lambda_8 = -\sigma_y \quad Q_8 = \frac{1}{\sqrt{2}} U \begin{bmatrix} 0 & 0 & 1 \\ 0 & 0 & 0 \\ 1 & 0 & 0 \end{bmatrix} V^T.$$
eigenvalues are the roots of the depressed cubic equation,
\[ \lambda_i = 2\sqrt{\frac{I_2}{3}} \cos \left[ \frac{1}{3} \left( \arccos \left( \frac{3I_3}{I_2} \sqrt{\frac{3}{I_2}} \right) + 2\pi(i - 1) \right) \right], \quad (7.9) \]
where \( i \in \{0, 1, 2\} \). The eigenvectors are then a linear combination of the scaling modes,
\[ Q_i = \sum_{j=0}^{2} z_j D_j \quad \text{where} \quad \begin{cases} z_0 = \sigma_x \sigma_z + \sigma_y \lambda_i \\ z_1 = \sigma_y \sigma_z + \sigma_x \lambda_i \\ z_2 = \lambda_i^2 - \sigma_z^2 \end{cases} , \]
and the exact weights in the combination varies based on the current deformation.\(^1\)

### 7.3.2 All Isotropic Energies Have the Exact Same Eigenvectors

The nine eigenmatrices that we have just seen arise from \( I_3 \) are in fact the eigenvectors of all isotropic energies. No matter what isotropic energy you write down, these will always be the eigenvectors. That primordial soup we saw when taking the numerical eigendecomposition of \( \Re^{9 \times 9} \) Hessians? It wasn’t a soup after all, it was a stack of the same nine Lego blocks all along.

Given their importance, let’s list them again here, once and for all:

\[ Q_i \in \{0, 1, 2\} = \sum_{j=0}^{2} z_j D_j \quad \text{where} \quad \begin{cases} z_0 = \sigma_x \sigma_z + \sigma_y \lambda_i \\ z_1 = \sigma_y \sigma_z + \sigma_x \lambda_i \\ z_2 = \lambda_i^2 - \sigma_z^2 \end{cases} , \quad (7.10) \]

\[ Q_3 = \frac{1}{\sqrt{2}} U \begin{bmatrix} 0 & -1 & 0 \\ 1 & 0 & 0 \\ 0 & 0 & 0 \end{bmatrix} V^T \quad Q_4 = \frac{1}{\sqrt{2}} U \begin{bmatrix} 0 & 0 & 0 \\ 0 & 0 & 1 \\ 0 & -1 & 0 \end{bmatrix} V^T \quad Q_5 = \frac{1}{\sqrt{2}} U \begin{bmatrix} 0 & 0 & 1 \\ 0 & 0 & 0 \\ -1 & 0 & 0 \end{bmatrix} V^T \quad (7.11) \]

\[ Q_6 = \frac{1}{\sqrt{2}} U \begin{bmatrix} 0 & 1 & 0 \\ 1 & 0 & 0 \\ 0 & 0 & 0 \end{bmatrix} V^T \quad Q_7 = \frac{1}{\sqrt{2}} U \begin{bmatrix} 0 & 0 & 0 \\ 0 & 0 & 1 \\ 0 & 1 & 0 \end{bmatrix} V^T \quad Q_8 = \frac{1}{\sqrt{2}} U \begin{bmatrix} 0 & 0 & 1 \\ 0 & 0 & 0 \\ 1 & 0 & 0 \end{bmatrix} V^T . \quad (7.12) \]

How do we know that these eigenvectors are so universal? We had that generic expression for the Hessians from Eqn. 5.50 back in §5.5:
\[ \text{vec} \left( \frac{\partial^2 \Psi}{\partial F^2} \right) = \sum_{i=1}^{3} \frac{\partial^2 \Psi}{\partial I_i} g_t g_i^T + \frac{\partial \Psi}{\partial I_i} H_i . \quad (7.13) \]

\(^1\)Ever since we found these equations, it has bugged me that these three eigenpairs are much less parsimonious than the other six. I suspect that there’s a slicker way to write these down, but it clearly involves a trick I haven’t thought of yet.
There are only two places that eigenvectors can come from: the $H_i$ matrices and $g_i g_i^T$ outer products. We already have complete knowledge of the eigenvectors that arise in the $H_i$ terms:

- The Hessian of $I_1$ has the twist eigenvectors (Eqn. 7.11). Past those, it doesn’t care – the rest spans an arbitrary subspace.
- The Hessian of $I_2$ is diagonal, so it is one big arbitrary, “don’t care” subspace.
- The Hessian of $I_3$ has the twists eigenvectors, just like $I_1$, so those will already align. Then it has the flip and the scaling eigenvectors, but since $I_1$ and $I_2$ both had arbitrary subspaces along those spans, they will just get pinned to flip and scaling.\(^2\)

That leaves $g_i g_i^T$ as the only term that could ruin our claim that we’ve found the nine eigenvectors that can exist. Since it is an outer-product, each $g_i g_i^T$ term is its own eigenvector, i.e. the eigenvector is the normalized version of $g_i$.

All three gradients, $g_1 = \text{vec } R$, $g_2 = \text{vec } 2F$, $g_3 = \text{Eqn. B.21}$ are orthogonal to the twist and flip eigenvectors.\(^3\) Therefore, these get chunked in with the $Q_i \in \{0,1,2\}$ eigenvectors, so nothing new can appear from them either.

If you have an isotropic energy, these are the eigenvectors. It doesn’t matter how gnarly the energy looks, these are the nine.\(^4\)

### 7.3.3 Cranking Out Analytic Eigenvalues

If the eigenvectors are exactly the same for all isotropic energies, the only place where energies stand apart must be the eigenvalues. We saw a generic (albeit ugly) way to compute the analytic eigenvalues of $I_3$, and we can generalize this by performing a deflation (see e.g. Bunch et al. (1978)). We will explicitly project off the twist and flip eigenmodes, so that what’s left over is a $\mathbb{R}^{3\times3}$ system of scaling modes.

\(^2\)A bunch of linear algebra identities are getting thrown around here that I haven’t mentioned before. Don’t feel bad if you can’t follow it entirely.

\(^3\)This appears as Lemma B.1 in Smith et al. (2019). Take $g_2 = \text{vec } 2F$ and $Q_3$ as an example. First, observe that the dot product $\text{vec } (2F)^T \text{vec } (Q_3) = 0$ can be written as the trace $\text{vec } (2F)^T \text{vec } (Q_3) = 2F : Q_3 = \text{tr } (2F^T Q_3)$. Then $\text{tr } (2F^T Q_3)$ can be expanded using $F = U \Sigma V^T$ and the definition of $Q_3$ to get

$$\text{tr } \begin{pmatrix} 2 \Sigma U^T U & 0 & 0 \\ 0 & 1 & 0 \\ 0 & 0 & 1 \end{pmatrix} V^T$$

and simplified to $\text{tr } \begin{pmatrix} 2 & 0 & 0 \\ 0 & \sigma_z & 0 \\ 0 & 0 & \sigma_z \end{pmatrix} \begin{pmatrix} 0 & -1 & 0 \\ -1 & 0 & 0 \\ 0 & 0 & 0 \end{pmatrix}$. The product of the two matrices produces all zeros along the diagonal, so the trace is zero. Moreover, as long as the middle matrix in $Q_3$ has all zeros along the diagonal, i.e. is a hollow matrix, this will always hold. Since the middle matrix in all of $Q_{1,3}$ are all hollow, they are all orthogonal.

\(^4\)The form of Eqn. 5.50 needs to be slightly more general to encompass all isotropic energies; some mixed $g_i g_j^T$ where $i \neq j$ terms can appear too. This doesn’t make a difference; no new eigenvectors appear.
The entries of this scaling-only, projected matrix $\mathbf{A}$ are:

$$a_{ij} = \frac{\partial \Psi}{\partial I_3} + 4 \frac{I_3}{\sigma_k} \frac{\partial^2 \Psi}{\partial I_2^2} + \sigma_k I_3 \frac{\partial^2 \Psi}{\partial I_3^2} + 2 \sigma_k \left( I_2 - \sigma_k^2 \right) \frac{\partial^2 \Psi}{\partial I_2 \partial I_3} + (I_1 - \sigma_k) \left( \frac{\sigma_k}{\partial I_3} \frac{\partial^2 \Psi}{\partial I_1 \partial I_2} + 2 \frac{\partial^2 \Psi}{\partial I_1 \partial I_2} \right)$$  \hspace{1cm} (7.14)

$$a_{ii} = 2 \frac{\partial \Psi}{\partial I_2} + \frac{\partial^2 \Psi}{\partial I_2^2} + 4 \sigma_i^2 \frac{\partial^2 \Psi}{\partial I_2^2} + I_3 \frac{\partial^2 \Psi}{\partial I_2 \partial I_3} + 4 \sigma_i \frac{\partial^2 \Psi}{\partial I_2 \partial I_3} + 4 I_3 \frac{\partial^2 \Psi}{\partial I_2 \partial I_3} + 2 \frac{I_3}{\sigma_i} \frac{\partial^2 \Psi}{\partial I_1 \partial I_2}. \hspace{1cm} (7.15)$$

In these equations, I have used a slightly irritating $\{i, j, k\}$ notation. The symbol $a_{ii}$ denotes a diagonal entry, where if $i$ lines up with the $x$ direction, the $\sigma$ on the right-hand-side is replaced with $\sigma_x$. The symbol $a_{ij}$ denotes an off-diagonal entry, and if this corresponds to the $(x, y)$ off-diagonal entry, then $\sigma_k$ on the right-hand-side becomes $\sigma_z$, i.e. whichever coordinate does not appear as $i$ or $j$. If you’re still confused, the Matlab code for this is given in Fig. 7.4.

The first three eigenvalues of the larger system are now the eigenvalues of the matrix $\mathbf{A}$. These can be solved for any number of ways, both numerical and analytic, using the cubic equation, the method of Jenkins and Traub (1970), the method of Smith (1961), a cornucopia of methods from Golub and Van Loan (2013), or plain old Eigen (Guennebaud et al. (2010)).

The other six eigenvalues, on the other hand, have a much more compact general form:

$$\lambda_3 = \frac{2}{\sigma_z + \sigma_y} \frac{\partial \Psi}{\partial I_1} + 2 \frac{\partial \Psi}{\partial I_2} + \frac{\partial \Psi}{\partial I_3}$$  \hspace{1cm} (7.16)

$$\lambda_4 = \frac{2}{\sigma_y + \sigma_z} \frac{\partial \Psi}{\partial I_1} + 2 \frac{\partial \Psi}{\partial I_2} + \frac{\partial \Psi}{\partial I_3}$$  \hspace{1cm} (7.17)

$$\lambda_5 = \frac{2}{\sigma_x + \sigma_z} \frac{\partial \Psi}{\partial I_1} + 2 \frac{\partial \Psi}{\partial I_2} + \frac{\partial \Psi}{\partial I_3}$$  \hspace{1cm} (7.18)

$$\lambda_6 = 2 \frac{\partial \Psi}{\partial I_2} - \sigma_z \frac{\partial \Psi}{\partial I_3}$$  \hspace{1cm} (7.19)

$$\lambda_7 = 2 \frac{\partial \Psi}{\partial I_3} - \sigma_z \frac{\partial \Psi}{\partial I_3}$$  \hspace{1cm} (7.20)

$$\lambda_8 = 2 \frac{\partial \Psi}{\partial I_2} - \sigma_y \frac{\partial \Psi}{\partial I_3}. \hspace{1cm} (7.21)$$

7.3.3.1 St. Venant-Kirchhoff

Let’s push through one instance of these eigenvalues, using our old friend the StVK energy:

$$\Psi_{\text{StVK}} = \mu \|\mathbf{E}\|_F^2 + \frac{\lambda}{2} \text{tr}^2(\mathbf{E}). \hspace{1cm} (7.22)$$

function [A] = Get_Stretching_System(Psi)
syms I1 I2 I3 sigma0 sigma1 sigma2 sigmai sigmak real;
A = sym(zeros(3,3));

% first derivatives
firstI1 = diff(Psi, I1);
firstI2 = diff(Psi, I2);
firstI3 = diff(Psi, I3);

% second derivatives
secondI1 = diff(firstI1, I1);
secondI2 = diff(firstI2, I2);
secondI3 = diff(firstI3, I3);

% mixed derivatives
secondI1I2 = diff(firstI1, I2);
secondI1I3 = diff(firstI1, I3);
secondI2I3 = diff(firstI2, I3);

% get the diagonal entry
aii = 2 * firstI2 + secondI1 + 4 * sigmai^2 * secondI2 + ...
(I3 / sigmai)^2 * secondI3 + 4 * sigmai * secondII2 + ...
4 * I3 * secondII1I3 + 2 * (I3 / sigmai) * secondII1I3;
A(1,1) = subs(aii, sigmai, sigma0);
A(2,2) = subs(aii, sigmai, sigma1);
A(3,3) = subs(aii, sigmai, sigma2);

% get the off-diagonal entry
aij = sigmak * firstI3 + secondI1 + 4 * (I3 / sigmak) * secondI2 + ...
sigmatk * I3 * secondI3 + ...
2 * sigmak * (I2 - sigmak^2) * secondI2I3 + ...
(I1 - sigmak) * (sigmak * secondII1I3 + 2 * secondII1I2);
A(1,2) = subs(aij, sigmak, sigma2);
A(1,3) = subs(aij, sigmak, sigma1);
A(2,3) = subs(aij, sigmak, sigma0);

% symmetrize and simplify
A(2,1) = A(1,2);
A(3,1) = A(1,3);
A(3,2) = A(2,3);
A = Simplify_Invariants(A);
end

Figure 7.4.: Matlab/Octave code to get an analytic expressions for stretching subspace of any isotropic energy $\Psi$.  

7. The Analytic Eigensystems of Isotropic Energies
The entries of the $A$ matrix evaluate to something not so bad-looking:

$$a_{ii} = -\mu + \frac{\lambda}{2} (I_2 - 3) + (\lambda + 3\mu) \sigma_i^2$$
$$a_{ij} = \lambda \sigma_i \sigma_j.$$

The last six eigenvalues are available directly:

$$\lambda_3 = -\mu + \frac{\lambda}{2} (I_2 - 3) + \mu \left(\sigma_y^2 + \sigma_z^2 - \sigma_y \sigma_z\right)$$ (7.23)
$$\lambda_4 = -\mu + \frac{\lambda}{2} (I_2 - 3) + \mu \left(\sigma_x^2 + \sigma_z^2 - \sigma_x \sigma_z\right)$$ (7.24)
$$\lambda_5 = -\mu + \frac{\lambda}{2} (I_2 - 3) + \mu \left(\sigma_x^2 + \sigma_y^2 - \sigma_x \sigma_y\right)$$ (7.25)
$$\lambda_6 = -\mu + \frac{\lambda}{2} (I_2 - 3) + \mu \left(\sigma_x^2 + \sigma_y^2 + \sigma_x \sigma_y\right)$$ (7.26)
$$\lambda_7 = -\mu + \frac{\lambda}{2} (I_2 - 3) + \mu \left(\sigma_x^2 + \sigma_y^2 + \sigma_x \sigma_y\right)$$ (7.27)
$$\lambda_8 = -\mu + \frac{\lambda}{2} (I_2 - 3) + \mu \left(\sigma_x^2 + \sigma_y^2 + \sigma_x \sigma_y\right)$$ (7.28)

A little verbose, but not too bad.

### 7.3.4 If You’re Lucky, Things Get Simpler

If we grind $\Psi_{ARAP}$ through Eqs. 7.14 - 7.21, we will get exactly the same eigenvalue expressions that we obtained in Eqs. 7.2 - 7.5. But hang on a minute ... where did all the ugly stuff from Eqs. 7.14 and 7.15 go?

In some cases, if you’re lucky, the first three eigenvectors work out to the scaling modes in Eqn. 7.8. In these cases, the eigenvalues also take on the following closed forms:

$$\lambda_0 = 2 \frac{\partial \Psi}{\partial I_2} + \frac{\partial^2 \Psi}{\partial I_1^2} + 4\sigma_x \frac{\partial^2 \Psi}{\partial I_2 \partial I_1} + 4\sigma_y \frac{\partial^2 \Psi}{\partial I_3 \partial I_1} + 4I_3 \frac{\partial^2 \Psi}{\partial I_3 \partial I_1} + 2\sigma_y \sigma_z \frac{\partial^2 \Psi}{\partial I_3 \partial I_1}$$ (7.29)
$$\lambda_1 = 2 \frac{\partial \Psi_q}{\partial I_2} + \frac{\partial^2 \Psi_q}{\partial I_1^2} + 4\sigma_y \frac{\partial^2 \Psi_q}{\partial I_2 \partial I_1} + 4I_3 \frac{\partial^2 \Psi_q}{\partial I_3 \partial I_1} + 2\sigma_y \sigma_z \frac{\partial^2 \Psi_q}{\partial I_3 \partial I_1}$$ (7.30)
$$\lambda_2 = 2 \frac{\partial \Psi_q}{\partial I_2} + \frac{\partial^2 \Psi_q}{\partial I_1^2} + 4\sigma_z \frac{\partial^2 \Psi_q}{\partial I_2 \partial I_1} + 4I_3 \frac{\partial^2 \Psi_q}{\partial I_3 \partial I_1} + 2\sigma_z \sigma_x \frac{\partial^2 \Psi_q}{\partial I_3 \partial I_1}$$ (7.31)

They’re still not that pretty, but the expression you get out at the end might be,\(^5\) and the need to call any additional solver has been removed entirely.

---

\(^5\)Also, remember I promised I’ll bake all this out into a Matlab script for you.

99
How do you know if you’ve hit the **Eigenvalue Jackpot**? If all the off-diagonal entries \( a_{ij} \) work out to zero, then the scaling modes become decoupled, and you’ve hit the jackpot. You can use Eqns. 7.29 - 7.31 instead of the building out an unwieldy matrix.

A bunch of energies hit the jackpot.

### 7.3.4.1 Symmetric Dirichlet

The Symmetric Dirichlet energy from Smith and Schaefer (2015b),

\[
\Psi_{SD} = \frac{\left(\|F\|_F^2 + \|F^{-1}\|_F^2\right)}{2},
\]  

(7.32)

has the following eigenvalues:

\[
\begin{align*}
\lambda_0 &= 1 + \frac{3}{\sigma_x^2} \\
\lambda_1 &= 1 + \frac{3}{\sigma_y^2} \\
\lambda_2 &= 1 + \frac{3}{\sigma_z^2} \\
\lambda_3 &= 1 + \frac{\sigma_x^2}{I_x^3} + \frac{(I_x^2 - \sigma_x^2)\sigma_x^3}{I_x^3} \\
\lambda_4 &= 1 + \frac{\sigma_y^2}{I_y^3} + \frac{(I_y^2 - \sigma_y^2)\sigma_y^3}{I_y^3} \\
\lambda_5 &= 1 + \frac{\sigma_z^2}{I_z^3} + \frac{(I_z^2 - \sigma_z^2)\sigma_z^3}{I_z^3} \\
\lambda_6 &= 1 + \frac{\sigma_x^2}{I_x^3} - \frac{\sigma_y^2}{I_y^3} - \frac{\sigma_z^2}{I_z^3} \\
\lambda_7 &= 1 + \frac{\sigma_y^2}{I_y^3} - \frac{\sigma_z^2}{I_z^3} - \frac{\sigma_x^2}{I_x^3} \\
\lambda_8 &= 1 + \frac{\sigma_z^2}{I_z^3} - \frac{\sigma_x^2}{I_x^3} - \frac{\sigma_y^2}{I_y^3}.
\end{align*}
\]

### 7.3.4.2 Symmetric ARAP

The Symmetric ARAP energy from Shtengel et al. (2017),

\[
\Psi_{SARAP} = \frac{\mu}{2} \left(\|F - R\|^2 + \|F^{-1} - R^{-1}\|^2\right),
\]  

(7.33)

also hits the jackpot:

\[
\begin{align*}
\lambda_0 &= \mu \left(1 - \frac{2}{\sigma_x^2} + \frac{3}{\sigma_z^2}\right) \\
\lambda_1 &= \mu \left(1 - \frac{2}{\sigma_y^2} + \frac{3}{\sigma_z^2}\right) \\
\lambda_2 &= \mu \left(1 - \frac{2}{\sigma_z^2} + \frac{3}{\sigma_x^2}\right) \\
\lambda_3 &= \mu + \frac{\mu}{\sigma_x + \sigma_z} \left[\frac{1}{\sigma_y^2} + \frac{1}{\sigma_z^2} - \frac{1}{\sigma_y^2} - \frac{1}{\sigma_z^2} - 2\right] \\
\lambda_4 &= \mu + \frac{\mu}{\sigma_x + \sigma_z} \left[\frac{1}{\sigma_x^2} + \frac{1}{\sigma_z^2} - \frac{1}{\sigma_x^2} - \frac{1}{\sigma_z^2} - 2\right] \\
\lambda_5 &= \mu + \frac{\mu}{\sigma_x + \sigma_y} \left[\frac{1}{\sigma_x^2} + \frac{1}{\sigma_y^2} - \frac{1}{\sigma_x^2} - \frac{1}{\sigma_y^2} - 2\right]
\end{align*}
\]
7. The Analytic Eigensystems of Isotropic Energies

\[
\begin{align*}
\lambda_6 &= \mu \left[ 1 + \frac{1}{(\sigma_y \sigma_z)^2} + \frac{\sigma_y^2 + \sigma_z^2}{(\sigma_y \sigma_z)^3} - \frac{1}{\sigma_y^2 \sigma_z} - \frac{1}{\sigma_y \sigma_z^2} \right] \\
\lambda_7 &= \mu \left[ 1 + \frac{1}{(\sigma_x \sigma_z)^2} + \frac{\sigma_x^2 + \sigma_z^2}{(\sigma_x \sigma_z)^3} - \frac{1}{\sigma_x^2 \sigma_z} - \frac{1}{\sigma_x \sigma_z^2} \right] \\
\lambda_8 &= \mu \left[ 1 + \frac{1}{(\sigma_x \sigma_y)^2} + \frac{\sigma_x^2 + \sigma_y^2}{(\sigma_x \sigma_y)^3} - \frac{1}{\sigma_x^2 \sigma_y} - \frac{1}{\sigma_x \sigma_y^2} \right]
\end{align*}
\]

7.3.4.3 Co-rotational (sort of)

The Co-rotational energy hits two out of three cherries on the slot-machine,

\[\Psi_{CR} = \mu \| \mathbf{F} - \mathbf{R} \|^2 + \frac{\lambda}{2} \text{tr}^2 (\mathbf{S} - \mathbf{I})\]

and has the scaling modes weakly couple in a single rotation mode:

\[
\begin{align*}
\lambda_0 &= 2\mu + 3\lambda \\
\lambda_{1,2} &= 2\mu \\
Q_0 &= \frac{1}{\sqrt{3}} \mathbf{R} \\
Q_{1,2} &= 2\text{D subspace orthogonal to } \mathbf{R}.
\end{align*}
\]

\[
\begin{align*}
\lambda_3 &= 2\mu + 2\lambda \frac{(I_1 - 3 - 2\mu)}{(\sigma_y + \sigma_z)} \\
\lambda_4 &= 2\mu + 2\lambda \frac{(I_1 - 3 - 2\mu)}{(\sigma_x + \sigma_z)} \\
\lambda_5 &= 2\mu + 2\lambda \frac{(I_1 - 3 - 2\mu)}{(\sigma_x + \sigma_y)} \\
\lambda_6 &= 2\mu \\
\lambda_7 &= 2\mu \\
\lambda_8 &= 2\mu.
\end{align*}
\]

For all of these energies, projecting to semi-positive-definiteness is now straightforward: look at the eigenvalues, and if they’re below zero, snap them to zero. As we did with ARAP, we can also derive the exact conditions under which they become indefinite in order to better understand their behavior.

None of these energies use the \( I_3 \) invariant, and that’s not a coincidence. The invariant \( I_3 = \sigma_x \sigma_y \sigma_z \) couples the scaling modes; the exact thing that all these jackpot winners don’t do. If your energy has \( I_3 \) lurking in it somewhere, then it’s inevitable that the first three eigenvalues will become non-linearly coupled.

7.3.4.4 Finally, Some Code, To Save You From Error-Riddled Typing

As promised, the code to produce these analytic eigensystems is provided in Fig. 7.5. These implement the Simple Eigenvalue Jackpot case. If \( I_3 \) appears in your energy, you’ll still need to build out your \( \mathbf{A} \) matrix using Eqns. 7.14 and 7.15. We will see an example of that next.
function [lambdas] = Get_Analytic_Eigenvalues(Psi)
    syms I1 I2 I3 real;
    syms sigma0 sigma1 sigma2 sigmai sigmaj sigmak real;
    lambdas = sym(zeros(9,1));

    % x-twist, y-twist, and z-twist
    lambdas(4) = (2 / (sigma1 + sigma2)) * diff(Psi, I1) + ... 
                  2 * diff(Psi, I2) + sigma0 * diff(Psi, I3);
    lambdas(6) = (2 / (sigma0 + sigma2)) * diff(Psi, I1) + ... 
                  2 * diff(Psi, I2) + sigma1 * diff(Psi, I3);
    lambdas(5) = (2 / (sigma0 + sigma1)) * diff(Psi, I1) + ... 
                  2 * diff(Psi, I2) + sigma2 * diff(Psi, I3);

    % x-flip, y-flip, and z-flip
    lambdas(7) = 2 * diff(Psi, I2) - sigma2 * diff(Psi, I3);
    lambdas(8) = 2 * diff(Psi, I2) - sigma0 * diff(Psi, I3);
    lambdas(9) = 2 * diff(Psi, I2) - sigma1 * diff(Psi, I3);

    % x-scale, y-scale and z-scale
    lambdaScale = 2 * diff(Psi, I2) + diff(Psi, I1, 2) + ... 
                   4 * sigmai^2 * diff(Psi, I2, 2) + ... 
                   sigmaj^2 * sigmak^2 * diff(Psi, I3, 2) + ... 
                   4 * sigmai * diff(diff(Psi, I1), I2) + ... 
                   4 * I3 * diff(diff(Psi, I2), I3) + ... 
                   2 * sigmaj * sigmak * diff(diff(Psi, I3), I1);
    lambdas(1) = subs(lambdaScale, [sigmai, sigmaj, sigmak], ... 
                      [sigma0, sigma1, sigma2]);
    lambdas(2) = subs(lambdaScale, [sigmai, sigmaj, sigmak], ... 
                      [sigma1, sigma0, sigma2]);
    lambdas(3) = subs(lambdaScale, [sigmai, sigmaj, sigmak], ... 
                      [sigma2, sigma0, sigma1]);

    % try to get the simplest expression
    lambdas = Simplify_Invariants(lambdas);
end

Figure 7.5.: Matlab/Octave code to get analytic expressions for the eigenvalues of any isotropic energy Psi. The routine for Simplify_Invariants is defined in Fig. 7.6.
function [out] = Simplify_Invariants(in)
syms I1 I2 I3 sigma0 sigma1 sigma2 real;
out = subs(in, I1, sigma0 + sigma1 + sigma2);
out = subs(out, I2, sigma0^2 + sigma1^2 + sigma2^2);
out = subs(out, I3, sigma0 * sigma1 * sigma2);
out = simplify(out);
out = subs(out, sigma0 + sigma1 + sigma2, I1);
out = subs(out, sigma0^2 + sigma1^2 + sigma2^2, I2);
out = subs(out, sigma0 * sigma1 * sigma2, I3);
out = simplify(out);
end

Figure 7.6.: Matlab/Octave code for simplifying expressions using the definitions of the invariants. This is called from Get_Analytic_Eigenvalues in Fig. 7.5.

7.4 The Stable Neo-Hookean Eigensystem

Let’s take a closer look at the Stable Neo-Hookean eigensystem, because it exhibits some qualitatively different behaviors from all the “jackpot” energies we just saw.

We can obtain the complete eigenmodes by punching the $\Psi_{SNH}$ symbolically into Matlab/Octave code and running the code from Figs. 7.4 and 7.5. An example of the calls is given in Fig. 7.7.

The resulting scaling system is:

$$a_{ii} = \mu + \frac{I_3^2}{\sigma_i^2}$$

$$a_{ij} = \sigma_k \left( \lambda (2I_3 - 1) - \mu \right).$$

The twist and flip eigenvalues are:

$$\lambda_3 = \mu + \sigma_z \left( \lambda (I_3 - 1) - \mu \right)$$  \hspace{1cm} (7.34)

$$\lambda_4 = \mu + \sigma_x \left( \lambda (I_3 - 1) - \mu \right)$$  \hspace{1cm} (7.35)

$$\lambda_5 = \mu + \sigma_y \left( \lambda (I_3 - 1) - \mu \right)$$  \hspace{1cm} (7.36)

$$\lambda_6 = \mu - \sigma_z \left( \lambda (I_3 - 1) - \mu \right)$$  \hspace{1cm} (7.37)

$$\lambda_7 = \mu - \sigma_x \left( \lambda (I_3 - 1) - \mu \right)$$  \hspace{1cm} (7.38)

$$\lambda_8 = \mu - \sigma_y \left( \lambda (I_3 - 1) - \mu \right).$$  \hspace{1cm} (7.39)

7.4.1 When Does It Go Indefinite?

When do these eigenvalues go indefinite? These are slightly more difficult to analyze, because the relations are still non-linear. When we were obtaining the eigensystem of $I_3$, the flip modes corresponding to $\lambda_6, \ldots, \lambda_8$ were always negative, so let’s take a look at what happens after they’ve been plugged into the Stable Neo-Hookean energy.
% Boilerplate to load up the symbolic package in Octave
isOctave = exist('OCTAVE_VERSION', 'builtin') ~= 0;
if (isOctave)
    pkg load symbolic;
end

% Declare the invariants and constants
syms I1 I2 I3 mu lambda real;

% Build the SNH energy
Psi = (mu / 2) * (I2 - 3) - mu * (I3 - 1) + (lambda / 2) * (I3 - 1)^2;

% Build the ARAP energy
% Psi = I2 - 2 * I1 + 3

% Get the analytic twist and flip eigenvalues, and
% maybe the stretching values too, if you hti the jackpot
[lambdas] = Get_Analytic_Eigenvalues(Psi);

% Get the stretching matrix
A = Get_Stretching_System(Psi);

% Did we hit the Simple Eigenvalue Jackpot?
jackpot = A(1,2) + A(1,3) + A(2,3);
if (jackpot == sym(0))
    printf('You HIT the Jackpot!
');
    printf('No stretching matrix is needed.
');
    printf('Here are your eigenvalues:
');
    lambdas
else
    printf('You MISSED the Jackpot!
');
    printf('The stretching eigenvalues are the three
');
    printf('eigenvalues from this matrix:
');
    A
    printf('The last six are these:
');
    lambdas(4:9)
end

Figure 7.7.: Matlab/Octave code for obtaining the analytic eigenvalues and stretching system for the Stable Neo-Hookean energy, $\Psi_{SNH}$. If you want to see what it feels like to win the Simple Eigenvalue Jackpot, uncomment line 13 and run everything on $\Psi_{ARAP}$.
In particular, we can focus on $\lambda_6$, and examine the conditions under which it becomes zero, i.e. when it is on the cusp of becoming indefinite. Solving for the $\sigma_x$ roots of the quadratic

$$
\mu - \sigma_z \left( \lambda (\sigma_x \sigma_y \sigma_z - 1) - \mu \right) = 0
$$

(7.40)
yields the two roots:

$$
\sigma_x = \frac{\lambda + \mu \pm \sqrt{4\sigma_y \sigma_z \lambda \mu + (\lambda + \mu)^2}}{2\lambda \sigma_y \sigma_z}.
$$

(7.41)

Let’s examine a specific instance of $\lambda$ and $\mu$. As $\nu \rightarrow \frac{1}{2}$, which corresponds to biological tissue like muscle and skin, $\lambda \rightarrow \infty$ and overwhelms the $\mu$ term.

Another way to phrase this domination of $\lambda$ over $\mu$ is to set $\lambda = 1$ and $\mu = 0$:

$$
\sigma_x = 1 \pm \frac{1}{2 \sigma_y \sigma_z} = \left\{ 0, \frac{1}{\sigma_y \sigma_z} \right\}.
$$

(7.42)

The $\sigma_x = 0$ corresponds to the total pancake case where the elements has been squashed to zero volume. Indefiniteness make sense there; there are myriad ways to un-pancake yourself.

The $\sigma_x = \frac{1}{\sigma_y \sigma_z}$ condition is interesting if we view $\sigma_y \sigma_z$ as the cross-sectional area orthogonal to the $\sigma_x$ direction:

$$
\sigma_x = \frac{1}{\text{area}_\perp}.
$$

If the $x$ direction has been squashed so much that its length is less than the inverse of $\text{area}_\perp$, then the material will buckle. These cases are illustrated in Fig. 7.8. If the middle of your element does not bulge out far enough to stabilize your current amount of squashing, the material has no choice but to buckle.

The twist eigenvalues produce the exact same roots under $\lambda = 1$, $\mu = 0$, so those modes are not safe from indefiniteness either. These conditions only apply to $\nu = \frac{1}{2}$, but nonetheless, our analytic eigenanalysis gives a glimpse into the qualitative behavior of the Stable Neo-Hookean energy that was not possible before.

I wonder what running this eigenanalysis on your favorite energy will reveal?
Figure 7.8.: From left to right: An undeformed volume is safely positive definite. As it is squashed by $\frac{1}{2}$, the cross-sectional area has to be at least 2 to prevent it from buckling. If this condition isn’t met, the volume does not have base wide enough to hold up, and must buckle.
Chapter 8

A Better Way For Anisotropic Solids

This chapter is a loose retelling of §4 and §5 from Kim et al. (2019). If you want to cut to the chase, the Matlab code that spits out the analytic eigenvalues of an anisotropic energy is given in §8.2.5.

8.1 What’s Anisotropy?

So far, we have looked at isotropic solids. If you squish them down, they bulge out the exactly the same amount in all directions. If you stretch them out, they resist the same amount, no matter which direction you’re pulling in. Most real-world solids don’t do this exactly; they can be stiffer in some directions and softer in others.

Let’s compare a length of rubber band and a length of hemp rope sagging under gravity (Fig. 8.1). The rubber band sags a whole bunch. While the rope sags too, it doesn’t dip as far down, because it is composed of bundles of stiff fibers that all run in a preferred direction.

Figure 8.1.: Left pair: A length of rubber band sags a lot under gravity. Right pair: A length of hemp rope sags less, because it contains stiff fibers that all run in a preferred direction.

as far down, because it is composed of bundles of stiff fibers that all run in a single, preferred direction. In the extreme, if we ran a steel reinforcement rod down the length of the rope, it wouldn’t sag at all.
The material is anisotropic. It’s stiffer in some directions than it is in others.

The same thing can happen in a volumetric solid. When we squash down an isotropic material (Fig. 8.2), it bulges out equally in all directions. If the solid contains fibers that all run in a specific direction, then it will be reluctant to pooch out in that direction. One everyday material that does this is biological muscle. All the muscles in your body contain fibers, and when you flex a muscle, your brain is telling those aligned fibers to contract, i.e. become shorter (Fig. 8.3). When these fibers become shorter, they pull on the bones on either ends of the muscle, and that’s why your body moves. End of anatomy lesson.

8.2 The (Wrong) Cauchy-Green Invariants, Again

8.2.1 The $IV_C$ and $V_C$ Invariants

How do we tell our energies that there’s a preferred anisotropy direction that we want to be stiffer? Even if we have some anisotropy direction $a \in \mathbb{R}^3$ in mind, there’s nowhere to plug it into our existing invariants.
In biomechanics (see e.g. the survey of Chagnon et al. (2015)), this is solved by introducing a new set of invariants that explicitly take into account some vector \( a \). These new invariants gets stacked onto the end of our existing sequence of Cauchy-Green invariants: \(^1\)

\[
IV_C = a^T Ca \quad \quad V_C = a^T C^T Ca,
\]

where, as before, \( C = F^T F \). The anisotropy direction only needs to be defined at the rest shape. It gets automatically carried along as the object deforms.

\[\text{shape (Fig. 8.4). To see why, we can unpack } IV_C \text{ into:}\]

\[
IV_C = a^T F^T Fa = (Fa)^T Fa = \left( D_s D_m^{-1} a \right)^T D_s D_m^{-1} a.
\]

On the right, we can see that the invariant is just two applications of \( D_s D_m^{-1} a \). The first multiply, \( D_m^{-1} a \), rotates the fiber direction out of the original rest shape. Then it gets rotated into the current deformation by the second multiply, \( D_s \). Thus, the invariant will automatically transform the original \( a \) into the appropriate orientation for the current deformation.

**8.2.2 Gradients and Hessians, Again**

Let’s look at how to take the gradient and Hessian of an anisotropic energy.

**8.2.2.1 Anisotropic St. Venant Kirchhoff**

There’s a bunch of different anisotropic energies appearing in the literature that we can pick from, but for some reason people seem reluctant to name them. This one appears in anonymous form in both Liu et al. (2017) and Holzapfel (2005):

\[
\Psi_{ASVK} = \frac{\mu}{2} (IV_C - 1)^2.
\]

\(^1\)If you’re a real invariant nerd, it may bother you that these invariants don’t actually arise from the characteristic polynomial of \( F \). Where’s their generating equation? I don’t know. They’re definitely translation and rotation invariant though, so they at least fit the definition of “invariant” in that sense.
Since it follows the StVK approach of squaring a C-like term, let’s call it Anisotropic StVK. The gradient and Hessian are,

\[
\frac{\partial \Psi_{\text{ASVK}}}{\partial F} = \mu (IV_C - 1) \frac{\partial IV_C}{\partial F} \tag{8.4}
\]

\[
\text{vec} \left( \frac{\partial^2 \Psi_{\text{ASVK}}}{\partial F^2} \right) = \mu \left( (IV_C - 1) H_{IV} + g_{IV} g_{IV}^T \right), \tag{8.5}
\]

where

\[
\frac{\partial IV_C}{\partial F} = 2Faa^T \tag{8.6}
\]

\[
g_{IV} = \text{vec} \left( \frac{\partial IV_C}{\partial F} \right) = \text{vec} \left( 2Faa^T \right) \tag{8.7}
\]

\[
H_{IV} = \text{vec} \left( \frac{\partial^2 IV_C}{\partial F^2} \right) = 2aa^T \otimes I. \tag{8.8}
\]

Rather than slugging through these derivatives every time, just like in the isotropic case (Eqns. 5.8 and 5.50), there is a generic, oh-thank-my-lucky-stars-I-only-need-scalar-derivatives way of deriving the Hessian:

\[
\text{vec} \left( \frac{\partial^2 \Psi}{\partial F^2} \right) = \frac{\partial \Psi}{\partial IV_C} H_{IV} + \frac{\partial^2 \Psi}{\partial IV_C^2} g_{IV} g_{IV}^T. \tag{8.9}
\]

Once you have the relatively easy \( \frac{\partial \Psi}{\partial IV_C} \) and \( \frac{\partial^2 \Psi}{\partial IV_C^2} \) terms worked out\(^2\), you’re all done.

8.2.2.2 Anisotropic Square Root

Let’s try the generic approach on another (usually unnamed) energy that appears in biomechanics (Alastrué et al. (2008)):

\[
\Psi_{\text{ASqrt}} = \frac{\mu}{2} \left( \sqrt{IV_C} - 1 \right)^2. \tag{8.10}
\]

It’s almost exactly the same as the Anisotropic StVK energy, except for the square root. For that reason, let’s call it the Anisotropic Square Root model. The gradient is:

\[
\frac{\partial \Psi_{\text{ASqrt}}}{\partial F} = \mu \left( 1 - \frac{1}{\sqrt{IV_C}} \right) Faa^T. \tag{8.11}
\]

We can get the Hessian by taking the scalar derivatives,

\[
\frac{\partial \Psi_{\text{ASqrt}}}{\partial IV_C} = \frac{\mu}{2} \left( 1 - \frac{1}{\sqrt{IV_C}} \right) \quad \frac{\partial^2 \Psi_{\text{ASqrt}}}{\partial IV_C^2} = \frac{\mu}{4IV_C^3}, \tag{8.12}
\]

\(^2\)Again, don’t be ashamed to use Mathematica or Matlab. Really.
and plugging them in to get our final Hessian:\footnote{This expression looks slightly different from the one I gave in Kim et al. (2019) because I pushed some factors of 2 around in a different way. However, I assure you that if you work through all the terms, they are indeed equivalent.}:

\[
\text{vec} \left( \frac{\partial^2 \Psi_{\text{ASqt}}}{\partial F^2} \right) = \frac{\mu}{2} \left( 1 - \frac{1}{\sqrt{IVC}} \right) H_{IV} + \frac{\mu}{4IVC^2} g_{IV} g_{IV}^T. \tag{8.12}
\]

All done. Except that the Hessian might be indefinite.

### 8.2.3 The Eigensystem of $IV_C$

With Eqn. 8.9 in hand, we can apply the same approach as §7.3. If we can get an analytic expression for the eigendecomposition of $H_{IV}$, then we can get the analytic eigenstructure of any energy based on $IV_C$.

Fortunately, Matlab hack-and-slashing quickly reveals that the eigensystem of $H_{IV}$ has a rank-six null-space, so there are only three non-trivial eigenpairs:

\[
\lambda_{0,1,2} = 2, \quad Q_0 = \begin{bmatrix} a^T \\ 0 \\ 0 \\ 0 \end{bmatrix}, \quad Q_1 = \begin{bmatrix} 0 & 0 & 0 \\ a^T \\ 0 & 0 & 0 \end{bmatrix}, \quad Q_2 = \begin{bmatrix} 0 & 0 & 0 \\ 0 & 0 & 0 \\ a^T \end{bmatrix}. \tag{8.13}
\]

First off, it is an arbitrary subspace, so the forms given for $Q_{0,1,2}$ are not unique. They’re convenient to write in this form though. Second, the most general form of the eigenvalues is $\lambda_{0,1,2} = 2\|a\|_2^2$, but usually $a$ is specified as a unit vector. The purpose of $a$ is to convey that you want things stiffer in this direction, but the amount of stiffness you want would be communicated by some sort of $\mu$ constant. Baking it into the magnitude of $a$ would just cause confusion.

### 8.2.4 The Eigensystems of Arbitrary $IV_C$ Energies

With the eigensystem of $H_{IV}$ in hand, we can now build the eigensystems of arbitrary $IV_C$-based energies. Looking again at the generic form of these Hessians,

\[
\text{vec} \left( \frac{\partial^2 \Psi}{\partial F^2} \right) = \frac{\partial \Psi}{\partial IV_C} H_{IV} + \frac{\partial^2 \Psi}{\partial IV_C^2} g_{IV} g_{IV}^T,
\]

we already know the eigensystem of the first term on the right, $\frac{\partial \Psi}{\partial IV_C} H_{IV}$. The eigenvectors are still those from Eqn. 8.14, and the eigenvalues all get scaled by $\frac{\partial \Psi}{\partial IV_C}$.

Adding the $\frac{\partial^2 \Psi}{\partial IV_C^2} g_{IV} g_{IV}^T$ term complicates things, because (again) there’s no general theorem that helps you get the eigensystem of a sum of matrices, even if you already
know the eigensystems of the summands. But, while we’re not adding to a diagonal matrix like we did in §7.2 and §7.3, we’re doing something similar. We’re adding a rank-one update, $g_{IV}$, to a matrix with an arbitrary subspace, $H_{IV}$.

In our case, what happens is that one of the directions in the arbitrary subspace gets pinned to $g_{IV}$, i.e. the direction of the rank-one update. The other two directions are still arbitrary, but they then get pinned to the subspace that is orthogonal to $g_{IV}$.

More concretely, the first eigenpair is:

$$
\lambda_0 = 2 \left( \frac{\partial \Psi}{\partial IV} + 2IV \frac{\partial^2 \Psi}{\partial IV^2} \right) \quad \text{and} \quad Q_0 = \frac{1}{\sqrt{IV}} \text{Faa}^T.
$$

Keeping in mind that $g_{IV} = 2 \text{vec} \left( \text{Faa}^T \right)$, the eigenmatrix $Q_0$ has indeed been pinned to a normalized version of $g_{IV}$.

The other two eigenvalues are:

$$
\lambda_{1,2} = 2 \frac{\partial \Psi}{\partial IV}.
$$

The arbitrary subspace takes a little effort to construct, and in practice you never actually compute these explicitly. However, to show that it can be done, we first define the twist matrices,

$$
T_x = \begin{bmatrix} 0 & 0 & 0 \\ 0 & 0 & 1 \\ 0 & -1 & 0 \end{bmatrix}, \quad T_y = \begin{bmatrix} 0 & 0 & -1 \\ 0 & 0 & 0 \\ 1 & 0 & 0 \end{bmatrix}, \quad T_z = \begin{bmatrix} 0 & 1 & 0 \\ -1 & 0 & 0 \\ 0 & 0 & 0 \end{bmatrix}.
$$

Since the last two eigenvectors are arbitrary, you have to pick one of these matrices to start with. Just for the sake of argument, let’s pick $T_x$. Then the second eigenmatrix becomes:

$$
Q_1 = UT_x \Sigma V^T A.
$$

This effectively pins third eigenmatrix, which can be written in terms of $T_y$ and $T_z$,

$$
Q_2 = (\sigma_y \hat{a}_y) UT_y \Sigma V^T A - (\sigma_z \hat{a}_z) UT_z \Sigma V^T A,
$$

where $\hat{a}_{y,z}$ are the corresponding entries from $\hat{a} = V^T a$.

### 8.2.5 Matlab/Octave Implementation

Code that implements these expressions is given in Fig. 8.5, and examples running them on $\Psi_{AS}$ and $\Psi_{Sqrt}$ are in Fig. 8.6. Using these scripts, we obtain the following analytic eigenvalues for Anisotropic StVK,

$$
\lambda_0 = 2\mu(I_5 - 1) + 4\mu I_5 \\
\lambda_{1,2} = 2\mu(I_5 - 1).
$$
function [lambdas] = Get_Analytic_Eigenvalues(Psi)
    syms mu I5 real;
    lambdas = sym(zeros(3,1));
    firstI5 = diff(Psi, I5);
    secondI5 = diff(firstI5, I5);
    lambdas(1) = 2 * (firstI5 + 2 * I5 * secondI5);
    lambdas(2) = 2 * firstI5;
    lambdas(3) = 2 * firstI5;
    lambdas = simplify(lambdas);
end

Figure 8.5.: Matlab/Octave code for obtaining the analytic eigenvalues of an IVC-based anisotropic energy. For reasons that will become clear in the next section, IVC is instead called I5 in this code.

% Boilerplate to load up the symbolic package in Octave
isOctave = exist('OCTAVE_VERSION', 'builtin') ~= 0;
if (isOctave)
    pkg load symbolic;
end

% Declare the invariants and constants
syms I5 mu real;

% Build the Anisotropic StVK energy
Psi_AS = (mu / 2) * (I5 - 1)^2;
[lambdas_AS] = Get_Analytic_Anisotropic_Eigenvalues(Psi_AS)

% Build the Anisotropic Sqrt energy
Psi_Sqrt = (mu / 2) * (sqrt(I5) - 1)^2;
[lambdas_Sqrt] = Get_Analytic_Anisotropic_Eigenvalues(Psi_Sqrt)

Figure 8.6.: Matlab/Octave example calls to Get_Analytic_Anisotropic_Eigenvalues for $\Psi_{\text{AS}}$ and $\Psi_{\text{Sqrt}}$. 
and for Anisotropic Square Root:

\[ \lambda_0 = \mu \]  
\[ \lambda_{1,2} = \mu \left( 1 - \frac{1}{\sqrt{I_5}} \right) . \]  

(8.21)  
(8.22)

8.3 Better Invariants, Again

8.3.1 An Inversion-Aware Invariant

The energies we’ve looked all suffer from a version of the problem we saw in §6.1.2. They can’t tell if an element has been poked inside-out, i.e. inverted / reversed / prolapsed. Like the other Cauchy-Green-style invariants that came before it, \( IV_C \) has gone out of its way to throw away any reflection information, so it’s not a simple matter of re-jiggering its usage in the energy. I haven’t said much about the \( V_C \) invariant. Looking at it again, \( V_C = a^T C^T C a \), it just doubles down on the squaring strategy that stomps the information we care about, so it won’t help us either. Once again, we need to dream up a new invariant.

Way back in §5.3.3.2, we got a new set of invariants by performing a \( F^T F \rightarrow S \) substitution. Then the pesky \( F^T F \) doesn’t get the chance to stomp any reflections. We’re going to do the same thing here. Our new anisotropic invariants are:

\[ I_4 = a^T S a \]  
\[ I_5 = a^T S^T S a. \]  

(8.23)  
(8.24)

Similar to §5.3.3.2, we’ve introduced a new lower-order invariant, so some renumbering is needed. The new \( I_5 \) invariant is equivalent to \( IV_C^4 \), but since we need to make room for our new \( I_4 \), we’re renumbering it to 5. It’s not ideal, but the Arabic numeral subscript versus the non-subscripted Roman numeral at least has a distinct visual contrast.

8.3.2 The Gradient of \( I_4 \)

If we want to use the new \( I_4 \) in our energies, we will first need its gradient:

\[ \frac{\partial I_4}{\partial \mathbf{F}} = \frac{\partial \mathbf{R}}{\partial \mathbf{F}} : \mathbf{Faa}^T + \mathbf{Raa}^T. \]  

(8.25)

Uh-oh, that \( \frac{\partial \mathbf{R}}{\partial \mathbf{F}} : \mathbf{Faa}^T \) term doesn’t look too friendly. Fortunately, we poured tons of effort into understanding the structure of \( \frac{\partial \mathbf{R}}{\partial \mathbf{F}} \) back in §5.4. In 2D, this works out to:

\[ \frac{\partial \mathbf{R}}{\partial \mathbf{F}} : \mathbf{Faa}^T = \left( \begin{bmatrix} 1 & 1 \end{bmatrix} \mathbf{V}^T \mathbf{a} \right) \left( \begin{bmatrix} \sigma_x - \sigma_y \\ \sigma_x + \sigma_y \end{bmatrix} \right) \mathbf{U} \begin{bmatrix} 0 & -1 \\ 1 & 0 \end{bmatrix} \mathbf{V}^T. \]  

(8.26)

\[ ^4 \text{Since } IV_C = a^T F^T F a = a^T \Sigma \Sigma^T \Sigma \Sigma^T U a = a^T \Sigma \Sigma \Sigma \Sigma \Sigma^T U a = a^T S^T S a = I_5 \]
There’s a lot going on here, but the term to focus on is $\frac{\sigma_x - \sigma_y}{\sigma_x + \sigma_y}$, because as $\sigma_x + \sigma_y \to 0$, the gradient (and forces) will explode to infinity.

Is this some weird corner case that will never actually happen? Or do simulations hit this state all the time, and using this gradient will doom all of our simulations to unexpectedly and spontaneously explode? In order to investigate, I went ahead and coded up this gradient the first time I derived it, and the infinity case was hit on the VERY FIRST SIMULATION I RAN. I ran a few other ones, hoping that maybe I had just gotten unlucky. The explosions occurred with alarming frequency. Clearly, this invariant is too volatile to be used in its raw form.$^5$

### 8.3.3 The Sign of $I_4$

The information that is discarded by $IV_C$ is whether the element has inverted in the anisotropy direction. This distinction is slightly different from the isotropic case. As shown

![Figure 8.7.](image)

Figure 8.7.: An element inverts, but not in the fiber direction, shown in red. Therefore, the value of $I_4$ remains the same the entire time.

in Fig. 8.7, an element can be inverted, but unless it’s inverted in the fiber direction, the invariant doesn’t, and shouldn’t, care about it. It is only when the fiber itself has been pushed inside out (Fig. 8.8) that a negative appears in $I_4$. What happens if we just try to extract and use only the sign information from $I_4$? Do the gradient and Hessian become less volatile?

Let’s write a sign-extracting signum function $S(x)$:

$$S(x) = \begin{cases} 
-1 & \text{if } x < 0 \\
0 & \text{if } x = 0 \\
1 & \text{if } x > 0 
\end{cases}$$

(8.27)

$^5$As for the Hessian of $I_4$, let’s just say I had to figure out how to print things in landscape mode on legal-sized paper just to look at it. It’s not pretty, so let’s not even discuss it.
The derivative of $S(x)$ is a fancy-looking-and-sounding Dirac delta function:

$$
\delta(x) = \begin{cases} 
0 & \text{if } x \neq 0 \\
\infty & \text{if } x = 0 
\end{cases}
$$

There’s still an infinity in there, so things aren’t perfect, but this is still a pretty good result. Just to be clear, we are looking at some supremely uncomplicated functions here (Fig. 8.9). The $S(x)$ is just a step function, and $\delta(x)$ would be the most boring function of all, a line of zeros, if it wasn’t for its one spike to infinity. When we wrap $I_4$ inside $S(I_4)$, the gradient becomes:

$$
\frac{\partial S(I_4)}{\partial F} = \delta(I_4) \frac{\partial I_4}{\partial F}.
$$

The volatile $\frac{\partial I_4}{\partial F}$, rife with infinities, still appears. However, any explosions to infinity are contained by the $\delta(I_4)$ term, which stomps (almost) everything to zero.\footnote{Ooooh Paul Dirac, quantum mechanics, etc.} With this better-behaved result in hand, we can now look at building a better anisotropic energy.

\footnote{We could argue about whether the zero should overwhelm the infinity in the limit, but I don’t see a practical reason to.}
8.4 An Inversion-Aware Anisotropic Energy

8.4.1 Previous Models, and Their Issues

The square-root model from before had a few attractive properties:
\[
\Psi_{\text{ASqrt}} = \frac{\mu}{2} \left( \sqrt{I_5} - 1 \right)^2.
\]

We’ve now replaced \(IV_C\) with \(I_5\) to follow our new numbering scheme, but they’re equivalent.

The ASTVK model was too non-linear; it raised everything to an unnecessarily high 4\textsuperscript{th} power, so the forces shoot off the top and bottom of the graph if you try to plot them out (see Fig. 8.10). In addition to introducing really big forces, this will also ruin the conditioning of the global Hessian, and slow down solver convergence. In contrast, the ASqrt model shows a nearly-linear response in Fig. 8.11. The forces won’t grow too large, even if you stretch your objects out really far. The conditioning of the global Hessian will get worse under large stretches, but again, not too quickly. Unfortunately, there is

the distressing issue that the energy plot has a kink (Fig. 8.11 (a)) which then makes the force response discontinuous (Fig. 8.11 (b)).
Things are actually worse than that. The PK1 plot for ASqrt contains two roots, at 1 and -1, which means that both are stable states for the energy. If the element is perfectly inverted, i.e. the rest state experiences a perfect reflection in the fiber direction, then the energy will think it is at the rest state, and exert no force.

This is the danger of using $I_5$ instead of $I_4$: it can’t tell the difference between a deformation of 1 (zero deformation) and -1 (perfect reflection) in the fiber direction. So the question is: can we build an anisotropic energy that has all the nice linear-looking properties of ASqrt, but doesn’t have the discontinuity issues and spurious stable states?

### 8.4.2 An Anisotropic ARAP Model

The following energy accomplishes both of these goals:

$$\Psi_{AA} = \frac{\mu}{2} \left( \sqrt{I_5} - S(I_4) \right)^2. \quad (8.31)$$

The plots for this new energy is given in Fig. 8.12. It gives a nice, kink-free parabola for the energy, and the PK1 resolves to just a line. In essence, we patched everything to the left of $I_4 = 0$ to give us the result we wanted. The only remaining issue is that at $I_4 = 0$, the Dirac delta will produce an an infinity.

Fortunately, by just looking at the plots, we can see that this isn’t your usual catastrophic singularity that starts exploding towards infinity if you’re anywhere near its neighborhood. It’s a point singularity that generates a bad value in one specific case, but as long as you don’t step directly on it, you’re just fine.

A straightforward patch suggests itself almost immediately. For $\Psi_{ARAP}$, patching in the value of $\Psi_{ARAP}(2)$ whenever we hit the case of $\Psi_{ARAP}(0)$ should work just fine. For the PK1, if you ever get unlucky and hit $\frac{\partial \Psi_{ARAP}}{\partial F}(0)$ dead-on, then replace the infinity with $-\frac{\partial \Psi_{ARAP}}{\partial F}(2)$. The smooth, linear response will then be maintained.
The PK1 of this energy is:

\[
\frac{\partial \Psi_{\text{AARAP}}}{\partial F} = \mu \left( \sqrt{I_5} - S(I_4) \right) \left[ \frac{\partial \sqrt{I_5}}{\partial F} - \frac{\partial S(I_4)}{\partial F} \right],
\]

but if use the fact that \( \delta(I_4) \frac{\partial I_4}{\partial F} = 0 \) almost everywhere, this simplifies to

\[
\frac{\partial \Psi_{\text{AARAP}}}{\partial F} = \mu \left( \sqrt{I_5} - S(I_4) \right) \frac{1}{2\sqrt{I_5}} Faa^T - \frac{\partial S(I_4)}{\partial F}.
\]

Taking this expression forward, the Hessian is then,

\[
\frac{\partial^2 \Psi_{\text{AA}}}{\partial F^2} = \mu \left[ \left( 1 - \frac{S(I_4)}{\sqrt{I_5}} \right) H_5 + 2 \frac{S(I_4)}{I_5^{3/2}} g_5 g_5^T \right],
\]

where, identical to the \( IV_C \) case,

\[
g_5 = \text{vec} \left( \frac{\partial I_5}{\partial F} \right) = \text{vec} \left( 2Faa^T \right),
\]

\[
H_5 = \text{vec} \left( \frac{\partial^2 I_5}{\partial F^2} \right) = 2aa^T \otimes I.
\]

If we treat \( S(I_4) \) as a constant, we can push everything through the code in Figs. 8.5 and 8.6 to obtain the eigenvalues

\[
\lambda_0 = \mu \quad (8.37)
\]

\[
\lambda_{1,2} = \mu \left( 1 - \frac{S(I_4)}{\sqrt{I_5}} \right). \quad (8.38)
\]

Building a semi-positive definite Hessian is then a matter of checking if \( \lambda_{1,2} < 0 \). If they are, we only need to build a rank-one matrix using \( \lambda_0 \) and the eigenmatrix \( Q_0 = \frac{1}{\sqrt{I_5}} Faa^T \).

### 8.4.3 What Other Models Are Out There?

We’ve only looking at one possible anisotropic model that uses the new \( I_4 \) invariant. We found that \( I_4 \) possesses some difficult properties, so we tried to keep the good while discarding the bad. If you can think of a better one, that’s great. Publish it as a paper, post the code publicly, or use it as the secret sauce in your proprietary simulator\(^8\).

\(^8\)Bear in mind that whenever I hear “mine’s better, but it’s a secret” I usually don’t start wondering what the secret is. I start wondering whether the speaker is a dummy, a liar, or both.
Chapter 9

Tips on Force-Jacobians

Computing the second derivative of our deformation gradient is necessary if we want to implement methods like Backward-Euler in the next chapter. Hand rolling the individual terms can be a daunting task. Over the years, I’ve picked up a few tricks that I thought I’d share. In the presentation I said this wasn’t going to be covered in the notes, but I now have the time. I also want to illustrate that I can’t be trusted and that you should forever remain a healthy skeptic of the things I say.

Let’s imagine we have one of our constitutive models on an element defined over four particles. We’ve taken the derivative of the deformation gradient $F$ to compute the forces that act on the particles. Taking the derivatives of these forces with respect to their positions $x_i$ yields the $12 \times 12$ Jacobian shown in figure 9.1.

\[
\begin{pmatrix}
\frac{\partial f_0}{\partial x_0} & \frac{\partial f_0}{\partial x_1} & \frac{\partial f_0}{\partial x_2} & \frac{\partial f_0}{\partial x_3} \\
\frac{\partial f_1}{\partial x_0} & \frac{\partial f_1}{\partial x_1} & \frac{\partial f_1}{\partial x_2} & \frac{\partial f_1}{\partial x_3} \\
\frac{\partial f_2}{\partial x_0} & \frac{\partial f_2}{\partial x_1} & \frac{\partial f_2}{\partial x_2} & \frac{\partial f_2}{\partial x_3} \\
\frac{\partial f_3}{\partial x_0} & \frac{\partial f_3}{\partial x_1} & \frac{\partial f_3}{\partial x_2} & \frac{\partial f_3}{\partial x_3}
\end{pmatrix}
\]

Figure 9.1.: An example $12 \times 12$ force Jacobian matrix composed of $3 \times 3$ blocks

If you don’t take the "Tensor-Way" to derive the force Jacobian matrix and instead decide to crank these terms by hand, there are some tricks one can play to reduce the effort. The Jacobian is symmetric, so we only need to compute the upper-triangular blocks and only the upper-triangular portion of the diagonal blocks are needed. To save us even more effort we should remember the property that our forces must sum to zero.

$$\sum_{i=0}^{3} f_i = 0$$
This means that the column blocks of our Jacobian should sum to a $3 \times 3$ matrix of zeros. Again because of our symmetry property, this means that the row blocks should also sum to a zero matrix. One can use this property to check your work for errors. There will likely be some numerical round-off accumulation, so things won’t exactly be zero, but if you make a mistake it will likely fail this property in an obvious way.

What else can we do with this knowledge? Well if we have the off-diagonal blocks of our first row, we can compute $\frac{\partial f_0}{\partial x_0}$ them them as follows:

$$\frac{\partial f_0}{\partial x_0} = -\left[ \frac{\partial f_0}{\partial x_1} + \frac{\partial f_0}{\partial x_2} + \frac{\partial f_0}{\partial x_3} \right]$$

Then let’s imagine that $\frac{\partial f_1}{\partial x_2}$ is has particularly nasty derivatives to perform, but that we have $\frac{\partial f_1}{\partial x_1}$ and $\frac{\partial f_1}{\partial x_3}$ along with the top row of blocks. We can compute $\frac{\partial f_1}{\partial x_2}$ in a similar fashion:

$$\frac{\partial f_1}{\partial x_2} = -\left[ \frac{\partial f_0}{\partial x_1} + \frac{\partial f_1}{\partial x_1} + \frac{\partial f_1}{\partial x_3} \right]$$

We can play similar games with other terms as we work our way down the matrix rows to reduce the amount of work needed by any hand derivations or code implementation. Even if you don’t crank the entries of these block terms by hand and go the “Tensor-Way” of Chapter 4, you might want to take advantage of our Jacobian properties in another way. Numerical round-off accumulation may result in forces that don’t exactly sum to zero and a force Jacobian that is not numerically symmetric. Restoring the property to the forces is straightforward.

$$f_0 = -\left[ f_1 + f_2 + f_3 \right]$$

For the force Jacobian blocks we can take a different approach. One can average the off-diagonal blocks and assign the average back to the strictly upper-diagonal block, $i < j$.

$$\frac{\partial f_i}{\partial x_j} = \frac{1}{2} \left[ \frac{\partial f_i}{\partial x_1} + \frac{\partial f_j}{\partial x_1} \right]$$

One then assigns the transpose of this average to the strictly lower-diagonal block, $j > i$

$$\frac{\partial f_j}{\partial x_i} = \frac{\partial f_i}{\partial x_j}^T$$

The diagonal blocks are then defined by the negative sum of the off-diagonal terms in the row.

$$\frac{\partial f_i}{\partial x_i} = -\sum_{j=1}^{i-1} \frac{\partial f_i}{\partial x_j}$$

---

1Thanks to Tim Haynes for showing me this property many years ago at Disney’s Secret Lab.
This will make the Jacobian symmetric. The rows and columns won’t sum exactly to zero, but the error is usually reduced. This process is not required for things to work properly in a simulator. However, it can be handy in debugging for the force Jacobian to better reflect the properties that we know it should possess.

9.1 A Warning To Non-Members

Let’s imagine you’ve derived all the necessary terms for the force Jacobian, verified them with a symbolic math program and then verified you have made absolutely no errors transforming the expressions into code. That is quite an achievement, but unfortunately you have fallen into a trap that has been known by a secret cabal of physics programmers for quite some time.\(^2\) The right math is wrong! What does that mean? It simply means that the force Jacobian can become indefinite which will cause your simulator a myriad of problems. In some cases the force Jacobian will be indefinite, but other terms will mask it in the global matrix and you’ll be blissfully unaware. Other times it will lead to simulation artifacts or your solver failing to converge.

So what can one do? Well, if you go through the rigor of computing the analytic eigensystem as presented in Chapter 7, you don’t need to worry. That work pays off and allows one to precisely avoid the force Jacobian from becoming indefinite. If you cranked the individual terms by hand though, you have probably taken a brute-force approach that does not allow for this analysis. In that case you will need to perform some experiments and make some ad-hoc adjustments to the math.\(^3\) I recommend that anyone writing a simulator, that uses the force Jacobian, have a mode that performs an eigendecomposition of it at the element level. Have this capability for all the instances of an implicit force in your simulator! This will help detect these issues and to help to analyze where and when they occur. For the cloth forces in Baraff and Witkin (1998), they occur only in the second term of the equation below from their paper, below:\(^4\)

\[
\frac{\partial f_i}{\partial x_j} = k \left[ \frac{\partial C}{\partial x_i} \frac{\partial C^T}{\partial x_j} + \frac{\partial C^2}{\partial x_i \partial x_j} \right]
\]

Sometimes the the second term can be completely omitted. Other times it is necessary to identify the deformation conditions lead to indefiniteness and only discard it then. This was illustrated for the case of a simple linear spring in Choi and Ko (2002). Even when such ad-hoc approaches are employed, the properties of the force Jacobian from before must still hold with respect to the row and column block sums. Besides the effort of deriving the force Jacobian, I suspect the indefiniteness pitfall has lead to many abandon

\(^2\)The first rule of the cabal is to always give credit where credit is due. This is something I had to painfully discover on my own, but over time found that many others had solved before me. Unfortunately that means I cannot properly uphold the first rule. I hope that I am not banished or worse.

\(^3\)This is the approach taken with the cloth forces in Pixar’s simulator Fizt.

\(^4\)The paper Baraff and Witkin (1998) has a negative on this equation, but when implementing implicit methods, such as Backward-Euler we negate the term again. One should be perform the eigendecomposition with respect to the final sign of the matrix that will need to be inverted.
their implementation of Baraff and Witkin (1998) and pivot to either Position Based Dynamics, (PBD), or Extended PBD which do not require the force Jacobian. Since we have lifted the mask on this problem, we will move on to discuss our motivation for computing the force Jacobian in the first place, implicit integration with Backward-Euler.
Chapter 10

Constrained Backward-Euler

Large Steps in Cloth Simulation, Baraff and Witkin (1998), ushered in a new era of deformable simulation by clearly demonstrating the benefits of implicit integration\(^1\). Despite the time that has passed, it remains the foundation of Pixar’s cloth and volume simulator, called Fizt. Using Backward-Euler\(^2\) we assemble and solve the linear system below at each time step:\(^3\)

\[
\begin{bmatrix}
M - h \frac{\partial f}{\partial v} - h^2 \frac{\partial^2 f}{\partial x^2}
\end{bmatrix} \Delta v = h f_n + h^2 \frac{\partial f}{\partial x} v_n,
\]

\[Ax = b.\]

This linear system is sparse, but can still be quite large in practice. We represent our system in a Block Compressed Row Storage format (BCRS) with \(3 \times 3\) blocks. We use Preconditioned Conjugate Gradient (PCG) to solve our linear system, but handle hard constraints different from the original paper, which we will discuss shortly. We use hard constraints where two-way coupling is not needed. Some attachments and collisions with character geometry, or other kinematic objects, fall into this category. We use penalty forces to handle attachment and collision between dynamic surfaces for two-way coupling, as did the original paper. Later in this chapter, we will also discuss techniques we have found useful to increase performance of the assembly and solution of this system despite its changing structure.

---

\(^1\)If you enjoyed the previous chapters and ever find yourself needing to name a child, I suggest Jacob for the first name, and Ian for the middle name. If you really want to commit, you can also change your last name to Matrix. If that sounds like an improbable last name, just remember John and Jenny Matrix, the Arnold Schwarzenegger and Alyssa Milano characters from Commando (1985).

\(^2\)Fizt also supports a second-order accurate BDF-2 scheme, but for simplicity we will only discuss the first-order accurate scheme.

\(^3\)We only take one Newton step per time step for performance.
Constraint Filtering in Baraff-Witkin Cloth

In this section, we will give a brief overview of the constraint mechanism described in Baraff and Witkin (1998). To precisely control the motion of a particle in any of its Degrees of Freedom (DOFs), a mass modification scheme was developed. Given the current particle state \((x_n, v_n)\), one can easily compute the required \(\Delta v\) to reach a target velocity or position state at the end of the time step. We project out any free DOFs from \(\Delta v\) and following the convention of the original paper, we label this quantity \(z\). To enforce the constraint during the solution of our linear system, the concept of per-particle constraint filters was introduced. Each particle has a filter \(S_i\) with its constrained DOFs as follows:

\[
S_i = \begin{cases} 
I & \text{if zero DOFs are constrained} \\
I - \hat{p}_i \hat{p}_i^T & \text{if one DOF } (\hat{p}_i) \text{ is constrained} \\
I - \hat{p}_i \hat{p}_i^T - \hat{q}_i \hat{q}_i^T & \text{if two DOFs } (\hat{p}_i \text{ and } \hat{q}_i) \text{ are constrained} \\
0 & \text{if all three DOFs are constrained}
\end{cases} \tag{10.1}
\]

These filters are applied within the inner loop of the original \textit{Fizt} solver. At each iteration the solution is projected onto a state which satisfies the constraints, but nothing about how this projection might impede the convergence of the solver was considered.

10.1.1 Pre-filtered Preconditioned Conjugate Gradient (PPCG)

In a later version, creatively named \textit{Fizt2}, we adopted the Pre-filtered Preconditioned Conjugate Gradient (PPCG) method presented in Tamstorf et al. (2015). We are only using the pre-filtering component from §8 in the paper, which already gives a significant speedup. We will touch on preconditioning further in §10.8.

The PPCG formulation eliminates the need to apply the filters in the inner loop of PCG. Unlike the method of Ye (2009), the PPCG method does not change the size of our linear system, which enables us to maintain certain aspects of our system data structures. PPCG also makes it more convenient to send our system to a direct solver like CHOLMOD or Pardiso. We use a direct solver as fallback to handle severely ill-conditioned systems.

The formulation is:

\[
\left( SAS^T + I - S \right) y = Sc \tag{10.2}
\]

\[
y = x - z \tag{10.3}
\]

\[
c = b - Az. \tag{10.4}
\]

\[^4\text{In chapters 10 and 11, we overload the use of the symbol } \hat{\cdot} \text{ to describe a unit vector, where in chapter 4 it was used to describe the cross-product matrix of a vector.} \]
The key equation is the first one. The $\text{SAS}^T$ term is a straightforward projection of the original matrix $A$ into the subspace spanned by the filters $S$.\footnote{Since $S$ is symmetric, we could just write $\text{SAS}$, but the transpose makes the projection a little more visually obvious.} This creates a nearly-rank-deficient matrix in the subspace comprised of the DOFs that were removed from the system. The $\text{I} - S$ term then serves to improve the conditioning of this subspace. Anywhere that filter was applied, the $\text{I} - S$ essentially gives things a boost with a bunch of ones.

This is easiest to see if some particle $i$ was constrained entirely, and $S_i = 0$. Now we are in trouble because we have a $3 \times 3$ block of zeros along the diagonal. The Gerschgorin circle theorem (roughly) states that the eigenvalue corresponding to the row must lie inside some disc. The diagonal entry prescribes the disc’s center ($a_{ii} = 0$), and the radius is the absolute sum of the off-diagonal entries in that row ($r = \sum_{j \neq i} |a_{ij}|$). The fact that the disc must be centered at zero is bad news, because it means that the eigenvalue is close to zero, and almost certainly ruining the conditioning of the matrix. The $\text{I} - S$ fix will paste an identity matrix precisely on top of that zero block, and replace those zeros along the diagonal with ones. The Gerschgorin discs are now centered around one, and eigenvalues will be near one, which is a much better state.

<table>
<thead>
<tr>
<th>Example</th>
<th>Prefiltered PCG</th>
<th>Modified PCG</th>
</tr>
</thead>
<tbody>
<tr>
<td>Miguel</td>
<td>4.05x</td>
<td>2.09x</td>
</tr>
<tr>
<td>Nana</td>
<td>2.95x</td>
<td>2.03x</td>
</tr>
<tr>
<td>Dress</td>
<td>1.95x</td>
<td>1.55x</td>
</tr>
<tr>
<td>Hector</td>
<td>2.09x</td>
<td>1.12x</td>
</tr>
<tr>
<td>Mama</td>
<td>2.92x</td>
<td>1.55x</td>
</tr>
<tr>
<td>Average</td>
<td>2.79x faster</td>
<td>1.67x faster</td>
</tr>
</tbody>
</table>

Figure 10.1.: Speedup of solver stage using Prefiltered PCG from Tamstorf et al. (2015), without multigrid, and Modified PCG from Ascher and Boxerman (2003) over the original Baraff and Witkin (1998) algorithm.

As observed by the authors (page 9, first column, second paragraph), boosting the filtered subspace in this way improves the conditioning of the matrix, so PPCG should converge even faster than the MPCG method in Ascher and Boxerman (2003). We found this to be true in our tests as well. PPCG runs $2.79 \times$ faster than the original Baraff and Witkin (1998) solve, while Ascher and Boxerman (2003) only runs $1.67 \times$ faster (Table 10.1). PPCG gives a 36.0% wall-clock improvement over the 12.6% improvement achieved by Ascher and Boxerman (2003) (Table 10.2). Again, this $2.79 \times$ speedup was achieved even without the multigrid component of the paper.

Finally, we performed a performance breakdown of Fitz using the PPCG formulation (Table 10.3). While we had become accustomed to seeing a roughly even split between the solver, matrix assembly, and collision processing (i.e. CCD) stages, we found that the collision stage was now the tallest nail.
## 10.2 Performance Improvement Techniques

System assembly and solution are critical to overall simulator performance. Parallel implementations of both are essential problem sizes we usually run with *Fizt*. We find
that we are not compute bound on modern CPU architectures, but instead limited by memory bandwidth. Therefore, reducing memory traffic is a worthy avenue of pursuit for gaining performance. In the following sections, we will discuss some techniques that we have used to improve the performance of system assembly and our PPCG solver.

### 10.3 Reverse Cuthill-McKee

At every iteration of the PPCG solver, we multiply our matrix $A$ with a vector. It is trivial to parallelize this operation over the rows of $A$ which does not require any locking. Since $A$ is sparse, the multiplication of the row blocks with a vector does not access the memory of the vector uniformly. We would like to reduce the amount of random access to the vector during this operation.

One way to reduce random access is to reorder the particles in a way that minimizes the matrix bandwidth. The Boost Graph Library has a number of reordering strategies, but Reverse Cuthill-McKee has worked well for our purposes. At the beginning of our simulation, we make a graph of the fixed system connectivity. Since Fizt does not do adaptive re-meshing or tearing, we can exploit the fixed structure of any cloth or volume meshes in the system. Any permanent attachments, such as springs, between dynamic objects are also considered. The graph edges are made from the $ij$ entries above the diagonal of our global Jacobian matrix. One iterate through all the fixed elements and create a unique collection of these edges and use an initial global ordering of the particles. Once the reordering is performed, we assign each simulated particle an index from this reordering which we call the global solver index. Local element and orderings remain unchanged. The global solver index now indicates the row index of the particle in our linear system.
10.4 System Assembly

As mentioned, a considerable amount of structure remains fixed during our simulation. The sparsity pattern of our dynamic objects remain fixed, but entries from collision penalty forces are transient and there may be other spring forces that are also transient. We divide our force elements into these two broad categories of fixed and transient.

Fixed force elements have local storage for the forces and their Jacobians with respect to $x$ and $v$. To support dihedral bending forces and tetrahedral elements, they have fixed storage for configurations of up to four particles. Since our Jacobians are symmetric, we only allocate the storage required for the upper diagonal block matrices. Given local element particle indices $i$ and $j$ such that $i \leq j$ the index of the $3 \times 3$ block in the appropriately sized storage array is given by the following code snippet:

```cpp
uchar FixedElement::localPairToArrayIndex(uchar i, uchar j) {
    // assert is here for the cautious.
    // If you pass negative indices that's another thing you should never do!
    assert(i <= j);
    uchar n = (_numParticlesInElement << 1) - 1;
    return j + ((n - i) * i) >> 1;
}
```

We also create a Fixed Global Sparse Matrix (FGSM) with $3 \times 3$ blocks for the entries of $\frac{\partial f}{\partial v}$ and $\frac{\partial f}{\partial x}$. Each block in the FGSM has a collection of pointers to the blocks of the fixed elements that contribute to it. At the beginning of our simulation, we construct the FGSM and flatten it into a single vector containing all of these blocks. The first step in our system assembly proceeds by traversing the fixed force elements in parallel to compute the forces and Jacobians which are stored locally. In a second pass, we traverse the blocks of the FGSM to accumulate the results of the first pass. We use a similar scheme for accumulating the forces from the elements to a global vector with a $3 \times 1$ entry per particle. Assembly can be done in a single pass and with less memory usage, the advantage of this two-pass approach is that it is lock-free.

Every simulated particle has a unique global solver index that indicates its associated block-row in the linear system. Our global system structures contain block entries...
both above and below the diagonal. While this requires more memory, it allows for simple and coherent traversal when we need to perform per-row operations. When accumulating into the FGSM blocks, we examine the corresponding pair of global solver indices of the block to determine whether we can add the block directly, or need to add its transpose. Some Jacobians have symmetric blocks, but others do not. Making a mistake of incorrectly orienting the block during accumulation can lead to long nights of debugging. The resulting simulations may have odd behavior, or your global system may go indefinite. You have been warned.

A deformable simulator may have many fixed forces, including simple per-particle forces like gravity, drag, goal-springs, etc. Having a clean and well structured element class for each force type might seem natural at first, but this has a downside. The more elements one needs to traverse, the more memory bandwidth will limit your performance. Consider the sparsity pattern of the following cloth example with point, triangle and dihedral forces:

```
<table>
<thead>
<tr>
<th>Stretch/Shear</th>
<th>Dihedral Bend</th>
<th>Per-Particle</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
```

It makes sense to at least group the element forces by their mesh component domains, but we can go further. Instead of having a force element that handles each type of mesh feature, we aggregate overlapping forces onto the same fixed force element. Some elements naturally will be responsible for more force types than others and some will remain isolated.

In the case of cloth, we aggregate stretch/shear, bending and per-particle forces. When accumulating the Jacobian blocks over the whole mesh as depicted in image (a), we must traverse 97 elements and accumulate from 1266 element blocks. These estimates assumes each stretch/shear contributes 6 blocks, each dihedral element contributes 10 blocks, and each particle contributes one block. We are only considering blocks from the upper diagonal of the element force Jacobians and our numbers are doubled to account for both $\frac{\partial f}{\partial v}$ and $\frac{\partial f}{\partial x}$.

By exploiting the aggregation of elements in (b), we only traverse 46 elements and accumulate from 870 element blocks. In this example, the aggregation strategy more than halved the number of elements traversed and reduced our element block access by almost a third. I don’t recommend the aggregation approach for a a first implementation, but if you do implement it, remember that the orientation of the blocks must be consistent at the element level when accumulating the Jacobian blocks from different forces.
Transient force elements are the second source of forces and Jacobians. Unlike fixed force elements, these do not have any local storage. Instead, we define a Transient Global Sparse Matrix (TGSM) which has per-thread row storage of the blocks using a flat-map structure. We traverse our transient elements in parallel, and each one deposits its block contributions into this structure. A parallel per-row reduction step is then performed over the per-thread flat-map structures in the TGSM. We handle force accumulation with a different structure and reduction pass.

We build two matrices to pass to our solver, but many entries from the TGSM overlap with the blocks of the FGSM. In the image below we illustrate the overlap in block entries from a vertex-face contact with the FGSM. We accumulate any overlapping entries from the TGSM into the FGSM in another per-row parallel pass. This process also collects indices to the non-overlapping entries of the TGSM for our next stage of populating our BCRS structures. This process effectively removes the overlapping blocks from the TGSM with respect to our solver.

At this stage, the FGSM and TGSM both contain the accumulated blocks of \( \frac{\partial f}{\partial v} \) and \( \frac{\partial f}{\partial x} \). We have not constructed either side of the linear system needed for our PPCG solver. We perform this construction as we build up two separate BCRS structures. As we populate the blocks of our BCRS structures, we first construct the term for \( A \) and then apply any necessary constraint filters \( S \). When we process a block on system diagonal, we add the \( I - S \) term.
The BCRS layout for the FGSM remains fixed. Only the block values need to be updated, which can be done efficiently in parallel. Further, since the bandwidth of the fixed system has been reduced, we use type `short` to represent our column indices as offsets from the main diagonal for a tiny reduction in memory bandwidth during the sparse-matrix vector multiplies. The BCRS layout for the TGSM is computed at each step from its remaining entries. This task is serial, but requires only one traversal over the rows of the TGSM to determine the storage requirements and rows offsets for the BCRS structure. Once the memory is allocated, these entries are also populated in parallel. We use type `unsigned int` for column indices because an entry may exist in any column from an interaction between two particles. In large systems, a `short` will not have the range for this task. Assuming we have our force accumulation, vector we can also construct the corresponding \( b - Az \) term for our Right Hand Side (RHS) during the traversal. We then only need to apply our constraint filters \( S \) to this term to complete the system assembly. Instead of performing a second pass over the data, we perform the construction of our RHS during our parallel construction of the LHS.

10.5 System Reduction and Boundary Conditions

At the beginning of each frame we examine which particles will remain fully constrained over it\(^6\). Sometimes large regions of our simulated objects become fully constrained and we can exploit this. If all the fixed force elements incident to a particle also have all fully constrained particles we remove the particle from the system so that it does not appear in the rows of our linear system\(^7\). We assign such particles a global solver index < 0. Fully constrained elements are also removed from our system assembly. We reindex the remaining particles global solver index to the correct range. We don’t re-apply our bandwidth reduction for performance reasons. While this won’t reduce the bandwidth further we take comfort in knowing it won’t grow any larger either.

For each fully constrained and removed particle we calculate \( v_k \) and \( \Delta v_k \), using finite differences of the known positions. Even though these particles may have been removed from the system, it is possible that collision forces or soft constraints will require the interaction between a simulated particle and one that has been removed. The simulated particle receives the force, but we still need to express the interaction, with what is essentially now a kinematic particle, through the \( \frac{\partial f}{\partial v} \) and \( \frac{\partial f}{\partial x} \) terms of our system. The force element Jacobian blocks of the between simulated particles including itself, will get accumulated into the global matrices just as before. Jacobian blocks between a simulated and removed particle must be handled differently since the removed particle is not represented in our system.

Instead we add \( [h \frac{\partial f}{\partial v_k} + h^2 \frac{\partial f}{\partial x_k}] \Delta v_k \) from the mixed block contributions of the LHS to

\(^6\)We also examine if any removed particles become unconstrained and if they and any associated elements need to be added back. We only remove particles if the number to be removed exceeds a threshold of 200.

\(^7\)We leave a boundary of fully constrained particles that have both simulated and non-simulated particles in our system presently even though technically this could be avoided.
the RHS. We also add any mix block velocity contribution from the \( h^2 \frac{\partial f_k}{\partial x_k} v_k \) term in the RHS. This technique is also used to handle arbitrary spring forces between points between a dynamic and kinematic object without increasing the size of our system.

### 10.6 PPCG Solver Details

*Fizt* computes almost everything in double precision. The primary exception is our PPCG solver, and the system we pass to it. When we assign the block values to our BCRS structures, we convert to single precision. We perform the same conversion to the RHS after applying our constraint filters. We pass a single-precision version to our PPCG solver to further reduce the memory bandwidth requirements of the sparse matrix vector multiplication.

Our PPCG solver is non-standard in that it takes two separate BCRS structures to represent the Left Hand Side (LHS), but the benefits of exploiting the fixed structure during assembly appears to be worth the awkwardness\(^8\). We have to handle the application of our LHS in this split form. We first determine which rows have terms from the TGSM BCRS and FGSM BCRS. Each row gets a function pointer appropriate to the task of multiplying the necessary entries with a vector. We also utilize loop unrolling of the row traversal with a block size of 4. The ranges and remainders for this unrolling are computed once per solver call solver for the entries of the TGSM BCRS. There are probably more gains to be had by spending more long nights with a profiling tools, a snapshot of our performance gains between early 2015 and Summer 2016 are presented in figure 10.5. The original times per frame do not include the addition of continuous collisions, which will be discussed later. The 2016 timings include the additional overhead continuous collisions.

### 10.7 A Word on Determinism

Much of *Fizt* is made parallel using OpenMP including our PPCG solver. Dynamic scheduling is used, but care must be taken when doing so. Using dynamic scheduling in a loop containing reduction can lead to non-deterministic results because of differences in the order of rounding error accumulation. Having a suite of regression tests that exercise the simulator is a critical part of maintaining production code, but this can only be achieved though if we have deterministic behavior. Otherwise, false positives will have us chasing non-existent bugs.

Other common sources of non-determinacy fall into a handful of categories. Do not store anything as a pointer in an ordered container like a set or map if traversal of the container can impact roundoff accumulation. Contacts are one such category. We find and gather penalty force contacts in a parallel in non-deterministic fashion, but choose not to compute the response on the fly. Instead we sort them before computing and

---

\(^8\)We experimented with using ViennaCL on the CPU with the same convergence tolerance and our custom solver was still faster.
acquiring their force and Jacobian terms which ensures that their contributions are accumulated the same way each time for a given thread count. There is a performance hit for a determinism guarantee. \textit{Fizt} has a deterministic mode for testing and a faster non-deterministic mode for production use.

### 10.8 Preconditioning

\textit{Fizt} uses a $3 \times 3$ Block Jacobi preconditioner that is the inverses of the diagonal blocks from the LHS. While simple, we have found that it works well in practice. More sophisticated preconditioners have been attempted, such as the parallel Incomplete Cholesky of \textit{Chow and Patel} (2015), and the other preconditioners in the ViennaCL library (\textit{Rupp et al.} (2016)). We found that they were not able to consistently accelerate our tests. Multigrid preconditioners like the one of \textit{Tamstorf et al.} (2015) are also a possibility, but their advantages only begin to become significant when meshes have very large vertex counts.

One obvious question is whether $3 \times 3$ is the optimal size for a Block-Jacobi preconditioner. We did indeed run a series of tests to verify this. Table 10.6 shows timings we collected a sandbox quasistatic simulator (\textit{Glove}) in order to study whether a different block size might benefit wider deployment in \textit{Fizt}. The mesh consisted of 70,831 vertices and 141,579 triangles.

For Table 10.6, we ran a quasistatic skin simulation, where the skin was simulated using

<table>
<thead>
<tr>
<th>Example</th>
<th>Simulated Vertices/Faces</th>
<th>Assembly</th>
<th>Solve</th>
<th>Average Frame</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>20834 / 40643</td>
<td>2.07 vs. 0.60</td>
<td>3.57 vs. 0.65</td>
<td>8.50 vs. 3.26</td>
</tr>
<tr>
<td></td>
<td>27384 / 49826</td>
<td>2.45 vs. 0.88</td>
<td>1.77 vs. 0.46</td>
<td>9.47 vs. 4.97</td>
</tr>
<tr>
<td></td>
<td>49647 / 95744</td>
<td>4.84 vs. 1.74</td>
<td>6.86 vs. 1.51</td>
<td>15.9 vs. 6.76</td>
</tr>
<tr>
<td></td>
<td>46359 / 90783</td>
<td>2.98 vs. 1.10</td>
<td>4.90 vs. 1.23</td>
<td>12.0 vs. 5.32</td>
</tr>
</tbody>
</table>

Figure 10.5.: \textit{Fizt} examples circa 2015/2016. All were run with 6 threads at 10 steps per frame. All times in seconds.
Figure 10.6.: Performance of Block-Jacobi Preconditioning over different block sizes. We ran a quasistatic cloth solve for three Newton iterations, and report the number of PCG iterations for each iteration, as well as the wallclock time of the overall solve.

<table>
<thead>
<tr>
<th>Block Size</th>
<th>Iteration 1</th>
<th>Iteration 2</th>
<th>Iteration 3</th>
<th>Total Time</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>981</td>
<td>822</td>
<td>2028</td>
<td>17.03s</td>
</tr>
<tr>
<td>3</td>
<td>625</td>
<td>531</td>
<td>1023</td>
<td>12.47s</td>
</tr>
<tr>
<td>6</td>
<td>654</td>
<td>542</td>
<td>1203</td>
<td>13.30s</td>
</tr>
<tr>
<td>10</td>
<td>721</td>
<td>597</td>
<td>1217</td>
<td>13.98s</td>
</tr>
<tr>
<td>12</td>
<td>628</td>
<td>541</td>
<td>1123</td>
<td>13.71s</td>
</tr>
<tr>
<td>25</td>
<td>666</td>
<td>553</td>
<td>1321</td>
<td>14.71s</td>
</tr>
<tr>
<td>50</td>
<td>598</td>
<td>513</td>
<td>850</td>
<td>13.69s</td>
</tr>
<tr>
<td>100</td>
<td>486</td>
<td>507</td>
<td>826</td>
<td>15.00s</td>
</tr>
</tbody>
</table>

Figure 10.7.: Performance of Block-Jacobi Preconditioning for $100 \times 100$ blocks, using different factorization strategies. The setup is otherwise the same as Table 10.6.

<table>
<thead>
<tr>
<th>Factorization</th>
<th>Iteration 1</th>
<th>Iteration 2</th>
<th>Iteration 3</th>
<th>Total Time</th>
</tr>
</thead>
<tbody>
<tr>
<td>LDL''</td>
<td>486</td>
<td>507</td>
<td>826</td>
<td>15.44s</td>
</tr>
<tr>
<td>Column-pivoting QR</td>
<td>498</td>
<td>509</td>
<td>837</td>
<td>17.42s</td>
</tr>
<tr>
<td>Full-pivoting QR</td>
<td>498</td>
<td>509</td>
<td>837</td>
<td>22.00s</td>
</tr>
<tr>
<td>Householder QR</td>
<td>498</td>
<td>509</td>
<td>837</td>
<td>17.63s</td>
</tr>
<tr>
<td>Partial-pivoting LU</td>
<td>498</td>
<td>509</td>
<td>837</td>
<td>17.24s</td>
</tr>
<tr>
<td>Full-pivoting LU</td>
<td>498</td>
<td>509</td>
<td>837</td>
<td>17.45s</td>
</tr>
</tbody>
</table>

a cloth simulation as described in Kautzman et al. (2018). For measurement purposes, the simulation was run to four digits of precision for three Newton iterations. We report the number of PCG iterations needed, as well as the overall wall-clock time. The $3 \times 3$ block is the clear winner on wall-clock time. The larger block sizes predictably reduce the number of PCG iterations, but the growing cost of computing the block inverses consistently outweigh this reduction.

The next question is whether a faster but less precise block inverse would win back some of the speedup. Again, as a preliminary test we tried a variety of factorizations available in the Eigen (Guennebaud et al. (2010)) library, and the data is shown in Table 10.7. None of the factorization strategies produced dramatically improved results.

To follow this line of inquiry to its conclusion, the last thing to try is an Incomplete Cholesky or LU factorization that approximates the block inverses. However, we halted the study based on the existing data. Even if an incomplete factorization could be found that exactly (and impossibly) matched the full factorization, and ran in the same
wall-clock time as the $3 \times 3$ case (again impossible), it would only reduce the number of PCG iterations by 20%. Since the PCG solver generally consumes 33% of the wall-clock time, the expected speedup to $Fizl$ would only be around 6.7%. Investing more effort in chasing this level of acceleration did not seem promising, so the inquiry was halted.
Chapter 11

Collision Processing

Collision detection and response are critical components of any dynamics simulator. A vast body of work on collision detection exists and a full day course on the topic could not do justice to it. In this chapter we will only discuss the techniques that we employ in our simulator *Fizt*, and acknowledge that there are a number of other ways to approach these problems. That said our combined approach has proven itself able to handle challenging scenarios found in a production environment and often ignored by the literature.

*Fizt* is used to simulate cloth and volumes that must be able to collide with the animated geometry which, we refer to as *kinematic geometry*. Collisions between *dynamic geometry*, including self-collision, must also be handled by our system. All of our collision geometry is described by manifold triangles meshes and in the case of tetrahedral volumes we only collide with the surface mesh. Each triangle mesh has an axis aligned bounding box hierarchy which is updated at each time step, and is typically rebuilt once per frame. The number of separate objects in *Fizt* is typically small, so we do not bother with a broad-phase collision algorithm like those typically found in rigid body simulators.

11.1 Proximity Queries

Proximity queries look at the geometry at a discrete snapshot in time and are used to detect potential collisions between mesh features. The goal of the response is to prevent actual collisions from occurring over a the time step. Our response to proximity-based contacts is encoded in our linear system, and interacts with the all the other forces in our system.

Vertex-face queries between dynamic particles and the faces of our kinematic meshes are performed first. Our kinematic meshes are mostly closed and consistently oriented. We define a collision envelope around the surface by offset and inset distance parameters. To prevent against redundant contact data, we partition the space above and below each mesh face using this envelope. We refer to these partitioned regions as *cells*. The *cells*
provide us with a way to cull many of the undesirable contacts that occur when the mesh resolution is much smaller than the specified collision envelope 1.

To construct the cell regions of the faces, we first define a unique bisector plane for each edge 2. We define the edge normal, \( \mathbf{n}_e \), as the average of the available neighboring face normals. The normal of the bisector plane for this edge is then defined by \( \mathbf{n}_{eb} = \mathbf{n}_e \times \mathbf{e}_{ij} \) with \( \mathbf{e}_{ij} = \mathbf{x}_j - \mathbf{x}_i \). We then normalize to get \( \hat{\mathbf{n}}_{eb} \) and define distance from a point \( \mathbf{x}_p \) to the bisector plane by \( d_{e\text{plane}} = \hat{\mathbf{n}}_{eb} \cdot (\mathbf{x}_p - \mathbf{x}_i) \). We then orient our plane inward toward its first face neighbor, \( f_0 \), by ensuring that the vertex opposite the edge produces a positive distance to the edge bisector plane.

To examine whether a particle is in a face cell, we first check that the position is between all three bisector planes by computing \( d_{e\text{plane}} \) for all three edges. When performing this check, we examine whether the face is \( f_0 \) or \( f_1 \) with respect to the edge. If it is \( f_1 \), we negate the distance returned. This allows for a simple check that \( d_{e\text{plane}} \geq 0 \) for all three edge bisector planes regardless of which face we are querying. Then we compute \( d_{f\text{plane}} = \hat{\mathbf{n}}_f \cdot (\mathbf{x}_p - \mathbf{x}_0) \) to determine if it is within our collision envelope in the normal directions and if it is above or below the face plane.

A particle position that satisfies all of these checks is inside our cell region, but this does not necessarily mean it violates our collision envelope. As an edge becomes more creased, it becomes more probable that the point can be inside the cell, but outside our collision envelope. We illustrate such this in the second image of figure 11.1. After we verify the point is inside the cell, we compute closest point to the face, (Schneider and Eberly (2003)), and compare the distance to our collision envelope.

To create the bounding boxes of our cell regions, we compute the vectors at each vertex defined by the intersection of the incident bisector planes. Using these directions, we enclose the the extruded vertices in both the offset and inset directions. Instead of scaling the direction by the inset or offset directly, we use the magnitude required to reach the inset or offset in the normal direction with respect to the face 3.

Large inset values for relatively thin regions can cause the inset region to extend through the backside of an opposing region of the the surface. This can lead to particles getting an erroneous response that pulls them through the other side of a surface. An inset reduction step is performed each time the simulator receives a new sample of the kinematic geometry. The reduced insets at these samples are then interpolated across the time steps. When we discover a particle that satisfies all of our contact criteria, we create a contact constraint for the particle and add the kinematic face to it. If a particle already has a contact constraint, we add the new face to it.

---

1 Some post processing of the per-particle contact data is still required despite our region partitioning. It is still possible that particle could contact multiple incident faces at a vertex or edge boundary.

2 One should always guard against degenerate cases like collapsed faces, edges or vanishing edge normals.

3 Finding the minimum bounding-box of a face cell requires some attention when the dihedral angle between edges becomes large.
11.2 Filtered Constraints for One-Way Response

Our kinematic meshes have scripted motion that remains unchanged by the simulator. The scripted motion allows us to know the kinematic positions at the end of each time step. One can think of kinematic objects being as made of particles with infinite mass, which means that collision with a dynamic particle is a one-way interaction.

A proximity contact with a single kinematic face has a straightforward response. We define local coordinates for the particle at the current state using the $u$, $v$ coordinates and its normal offset from the face. The $u$ and $v$ are the first two barycentric coordinates of the particle’s projection into the plane, but may lie outside the boundary of the face. Since we know the state of the kinematic triangle at the end of the step, we can construct the particles’s desired world space position at the end of the step. Using the current particle state $x$ and $v$, we can determine the $\Delta v$ needed to reach this desired position. We use the constraint filtering mechanism described in §10.1 to ensure our system solution achieves the target $\Delta v$ in the normal direction at the end of the step. Constraining only the normal direction allows it to slide tangentially. If we included any collision offset correction in our desired target position, it would result in unnecessary energy being added to the system. The technique described so far is only used to match the velocity of the point on the kinematic face. Offset correction is handled by the position alteration technique described section 6.4 of Baraff and Witkin (1998). They compute the $\Delta x$ term in the normal direction at the end of the step that is required to reach the desired collision offset. Then $\Delta x$ is added to the appropriate term multiplied by $\frac{\partial f}{\partial x}$ in the RHS of the system. Finally, $\Delta x$ is added to the position state update. The addition of the term to the RHS allows the connected particles to be informed about the additional displacement during the solve.
Following section 5.4 of Baraff and Witkin (1998), we determine whether a constrained particle that is sliding on a kinematic surface should be released by computing the residual constraint forces as the last step of our linear system solve. If the constraint force was pulling toward the surface, we do not constrain the particle in the next time step. There are more accurate ways to handle this, but as noted in Baraff and Witkin (1998), it is far less expensive than dealing with the combinatorial complexity of handling proper inequality constraints.

When a particle is in proximity of multiple kinematic faces, we employ a variant of the Flypapering technique described in Baraff and Witkin (2003). Each dynamic object in Fizt has an identical mesh representation that has its vertex positions warped by our character rig. Instead of using a weighted average of the target positions from multiple faces, we fully constrain the particle to move relative to one of the faces found by our proximity query. We choose the face with the motion that minimizes the difference in displacement between the constrained motion it would have on the particle and the motion of the warped position during the step. Said another way, we choose the face that would most closely match the velocity produced by the rigged motion.

In recent films we have taken a very different approach to handling self-intersection of our characters. We now run a volume simulation to deform the character geometry in order to leave space for a cloth simulation in a second pass. This is depicted in figure 11.2 and further details can be found in Wong et al. (2018). This technique is relies on a robust and efficient constitutive model, but also heavily depends on our collision detection and response between dynamic surfaces which we will cover next.

### 11.3 Proximity Between Dynamic Meshes

Our second set of proximity queries is for collision, and self collision, between dynamic surfaces. Dynamic surfaces such as cloth are not closed, and do not have the concept of
an inset region. Instead, we have a self-offset separate from our collision offset which is used for collisions between different dynamic meshes. Our vertex-face proximity query is very similar to that of kinematic meshes, but it must be altered to handle open boundaries. When a face has a boundary edge, we extend its bounding box by the maximum of the self-offset and offset values in the appropriate directions. To do this, we examine the sign from the negative of the boundary edge’s bisector normal. The box is extended with the maximum of our two offsets in the appropriate cardinal directions.

Our particle-in-cell test is no longer sufficient, but is still necessary. Without some notion of space-partitioning, a large self-offset on a fine resolution mesh could lead to undesirable self-collisions, which would deform the our mesh in ugly ways. Let’s examine a couple of particle-face cases where our previous check breaks down. In case (a) below the separating plane between the two faces would only allow the particle to collide with one face when it could be inside the collision envelope of both. In case (b) the self-collision between a face and a neighboring particle would be rejected by the bisector plane between the faces.
To address these issues, and to handle boundary edges, we present the new test criteria summarized by the following pseudo-code:

```plaintext
bool InFaceRegion(p, triFace)

// If the projection of the point into the face plane
// resides inside the face boundary
if pproj in triFace:
    return true

distToPlane[3] = {triFace.DistToBisector(0,p),
                  triFace.DistToBisector(1,p),
                  triFace.DistToBisector(2,p)}

// Test if the point is in the cell of the face
if all distToPlane >= 0:
    return true

// Is the point outside the plane of a boundary edge,
// and inside the planes of all non-boundary edges?
if (triFace.IsBoundaryEdge(0) and distToPlane[0] < 0) or
  (triFace.IsBoundaryEdge(1) and distToPlane[1] < 0) or
  (triFace.IsBoundaryEdge(2) and distToPlane[2] < 0):
    for e in [0,2]:
        if not triFace.IsBoundaryEdge(e) and distToPlane[e] < 0:
            return false

return true

// The point is not in the region of the face
return false
```

After a point passes this check, we determine if it is within our collision envelope. When a dynamic contact is found, the four particles and the barycentric coordinates between the closest points are stored.

Edge-to-edge contacts also employ our `InFaceRegion` test, but in a slightly different fashion. We first find the closest point between the two edges, $e_a$ and $e_b$, using the technique described in Möller et al. (2008). If the closest points are on the edge segments, and not at any of the vertices, we examine if the distance between the two closest points are within our collision envelope. After establishing the distance this criteria, we check that the point of contact on $e_a$ passes our `InFaceRegion` test with the available adjacent faces of $e_b$. Similarly, we examine whether the point of contact on $e_b$ passes our `InFaceRegion` test with the available adjacent faces of $e_a$. Only if the test pass on both sides do we create a dynamic contact for later processing.

Certain feature pairs are purposefully omitted. A vertex should never be tested against

\footnote{In this test, we ignore the case when the closest point between edges is an end point, because that will be covered by our vertex-face test. A separate test for when the edges are deemed parallel is also performed.}
any of its incident faces\footnote{The actual code comment reads: "Never collide with your own face. This is always sound advice!"} and two edges that share a vertex should never be tested.

11.4 Penalty Forces for Two-Way Response

For a long time in computer graphics, penalty forces have been frowned upon, and probably still are by many. To be effective, these forces must typically have a large magnitude, which leads to stiff systems will make explicit integrators slow or unstable. We are using implicit integration, so this is not a concern for us. Even so, we still don’t know the magnitude of our penalty force required to make any guarantee that it will prevent a potential collision from occurring. If we wanted to enforce equality constraints, our linear system would have to be augmented with Lagrange multipliers, and still would have the problem of our constraints exhibiting sticking behavior. Handling this properly involves solving large scale optimization problems with inequality constraints. Solving this problem between deformable bodies has come a long way in recent years, but the computation times remain unsuitable for our production use. Penalty forces have served us well in practice, so we will cover some of their implementation details.

Before proceeding further, there is some convention and notation to discuss. Our calculated response force will act in the direction of the collision normal, which we define to point from entity \(b\) to entity \(a\). Particles on the \(a\) and \(b\) side of the contact are assigned signed barycentric weights \(w\) with the convention that \(w_a \geq 0\) and \(w_b \leq 0\). Our penalty contact object stores the signed weights in a single array \(w\) that shares the same ordering as the particles.

This makes computing relative quantities between the points of contact a weighted sum. In the case of vertex-face proximity, we assign a positive weight of one to the vertex particle and the negative of the barycentric coordinates to face particles. We will refer to the points of contact on each side by \(p_a\) and \(p_b\). For now, let us assume that we are not recovering from a tangled situation as described by Baraff and Witkin (2003) and that our penalty spring is in a state of compression. We define \(\Delta x = \sum_{i=0}^{3} w_i x_i\) and \(\Delta v = \sum_{i=0}^{3} w_i v_i\). Our desired collision distance is \(d_{\text{offset}}\). The spring stiffness and damping parameters, which must be positive, are \(k_s\) and \(k_d\). The proximity contact normal is defined as \(\hat{n}_c = \frac{\Delta x}{\|\Delta x\|}\). We can now express the force acting on \(p_a\) as:

\[
\begin{align*}
  f_a &= -[(k_s\|\Delta x\| - d_{\text{offset}}) + k_d(\Delta v \cdot \hat{n}_c)]\hat{n}_c
\end{align*}
\]

We deliberately chose to compute the response at \(p_a\) for our vertex-face example. Distributing the response force to the particles is a scaling of \(f_a\) by the signed weight of each particle. This follows from a simple application of the chain-rule and our sign convention, which conveys that \(f_b = -f_a\):

\[
  f_i = w_i \ast f_a
\]
We must now compute the force gradients of our spring for implicit integration. The spatial gradient block of $f_a$ with $x_{pa}$ is:

$$\frac{\partial f_a}{\partial x_{pa}} = -k_s[\hat{n}_c \hat{n}_c^T] - k_s(\|\Delta x\| - d_{offset})(I - \hat{n}_c\hat{n}_c^T)$$  \hspace{1cm} (11.3)

The full spatial force gradient for a linear spring is given in equation (11.3), but it presents us with a problem. For a spring in compression, the second term can make our system indefinite, as described in Choi and Ko (2002). This term filters motion orthogonal to the direction of our spring response, so we discard it and only keep the outer product term:

$$\frac{\partial f_a}{\partial x_{pa}} = -k_s[\hat{n}_c \hat{n}_c^T]$$  \hspace{1cm} (11.4)

Our signed weight convention make it easy to distribute these Jacobians to the blocks that correspond to our particle pairs. If $i$ and $j$ are the indices of any two particles on in our contact configuration then:

$$\frac{\partial f_i}{\partial x_j} = (w_iw_j)\frac{\partial f_a}{\partial x_{pa}}.$$  \hspace{1cm} (11.5)

The same relationship naturally holds for the analogous velocity gradient block:

$$\frac{\partial f_a}{\partial v_{pa}} = -k_{td}[\hat{n}_c \hat{n}_c^T].$$  \hspace{1cm} (11.6)

The conventions introduced in this section make it trivial to generalize code to handle the response for edge-edge proximity contacts, but it also generalizes further. We can define arbitrary forces to act on any point of a mesh, distribute the forces to the mesh particles, and distribute the Jacobians to the blocks of the linear system. Think of the cool things you might construct with this power!

**11.5 Faux Friction Effects**

Accurate frictional forces are not an option for the performance requirements of Fizt. Instead, for dynamic contacts, we allow artists to control tangential motion though a tangential damping parameter $k_{td}$. Our tangential damping force is defined $f_t = -k_{td}v_{trel}$ which acts opposite the direction of the relative tangential velocity. As before, we use the signed weights $w$ to distribute this force to the particles involved. We add the contribution $-k_{td}[I - \hat{n}_c\hat{n}_c^T]$ to our velocity gradient block $\frac{\partial f_a}{\partial v_{pa}}$ before distributing the weighted blocks of the particle pairs. This approach is cheap and artist friendly, but it can never model static friction behavior because there must be some relative tangential motion present in order for a resisting force to be applied.
Friction effects between dynamic particles and kinematic faces are handed differently. After accumulating all the forces on each particle, if the particle has a contact with a single kinematic face, we use the accumulated force to estimate the normal force. When the normal force is negative, a tangential spring force is then applied. The magnitude of this tangential spring force is set to $\mu ||n_f||$ to approximate dynamic friction behavior.

### 11.6 Debugging Proximity Contact Detection

Writing small unit tests is highly recommended, but one might omit or misunderstand certain cases. In addition to unit tests, I recommend developing the ability to visualize algorithms in action. Pixar’s Universal Scene Description (USD), has proven quite useful for this task. Fitz has the option to write out the geometry and partition cells at sub-frames. We also write out proximity contacts as linear curves that vary in number over time. Using Pixar’s usdview application, this data can be inspected over time. Image (a) below shows the partition cells on a kinematic mesh along with lines that correspond to dynamic particle contacts with it. Image (b) shows the vertex-face and edge-edge self contacts. Similar visualization output also exists for our continuous collision and global intersection analysis algorithms, which will be discussed in later sections.

### 11.7 Continuous Collision Detection

Thin and fast-moving geometry will lead to temporal aliasing issues if a simulator relies solely on proximity queries. Instead of testing discrete states for probable contacts, Continuous Collision Detection (CCD), determines the true collisions that occur between two position states of our geometry. CCD techniques for deformable surfaces were first made popular by Moore and Wilhelms (1988), Provot (1995) and Bridson et al. (2002). A large volume of work followed over the years to make the technique more robust to numerical issues and to improve the performance. However, many papers on CCD emphasize the robustness of their method and make strict guarantees that if things start in a penetration-free state, the algorithm will reliably maintain it. It is our experience

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This uses an explicit guess for the normal force, and does not reflect the true normal force at the point of contact over the time step.
that animation often requires things to intersect at one time or another, and that such guarantees are impossible in practice. Also in a production environment, it is an extra burden for the artists to guarantee a pristine initial state. In the last section of this chapter, we will discuss how coupling CCD with the results of another algorithm can help us handle these issues.

In *Fizt*, CCD is performed after we have integrated to find a new candidate position state. CCD assumes a linear trajectory of the particles over the time step, which is conveniently true for our Backward-Euler integrator as well. Our BVH structures enclose the mesh primitives at both of these states. Candidate vertex-face and edge-edge pairs are then discovered by leaf bounding-box overlaps.

The primitive test for both cases share a common idea. Using the subscripts $s$ and $e$ to denote the start and end positions of a vertex, we parameterize the motion of each vertex with respect to the variable $t \in [0, 1]$ by $x_i(t) = (1 - t)x_is + t(x_{ie})$. We construct a tetrahedron from the four vertices at the begin state, and want to determine if the volume ever goes to zero during the interval. This corresponds to all four points becoming co-planar.

Let us first examine the vertex-face case. We define the face normal over time by:

$$n(t) = (x_1(t) - x_0(t)) \times (x_2(t) - x_0(t))$$ (11.7)

When $n(t) \cdot (x_p(t) - x_0(t)) = 0$ our vertex $v_p$ is coplanar to the face.

A similar test can be made between two edges $e_a$ and $e_b$. Each edge has a two vertex trajectories, $x_0(t)$ and $x_1(t)$. We define the normal by:

$$n(t) = (x_{a1}(t) - x_{a0}(t)) \times (x_{b1}(t) - x_{b0}(t))$$ (11.8)

The equation for our test becomes $n(t) \cdot (x_{a0}(t) - x_{b1}(t)) = 0$.

Both tests result in an equation that is, at most cubic in $t$. Only the real roots $t \in [0, 1]$ are of interest. Even though there is an analytic formula for the roots of a cubic polynomial, robustly finding these roots is not easy. How small can the leading coefficient get before the equation is considered a quadratic? When inserting roots back into the cubic equation, what is an acceptable error? Exact boolean tests for collision have been developed by Brochu et al. (2012) and Tang et al. (2014), and while they are recommended to avoid missing a collision, they do not find the root needed when a collision does occur. We recommend the analytic techniques described in Jim Blinn’s multi-part article, *How to solve a cubic equation*, but found it necessary to polish our roots with a numerical technique. The Boost math library provides an implementation of Schröder’s method, which we’ve found adequate for this purpose. We fully acknowledge that this approach is not as robust as other methods available in the literature.

Geometric tests are performed for each polished root $t \in [0, 1]$ in sorted order. The position of the vertices are first advanced to the time of our candidate root $t$. Numerical roundoff will likely keep the points from being "exactly" coplanar, so this should never
be assumed by the geometric test. If we determine that the point of contact lies within our features, then we can stop our search and record the contact. Otherwise we must check any remaining root candidates.

A good unit testing framework is always your friend, but this can be especially true when developing algorithms can that be affected adversely by numerical errors. One can only begin to have some confidence after procedurally testing millions of collision scenarios that cover various scales and trajectories. Even having taken such measures for our implementation, failures are still possible since we don’t perform the exact methods mentioned. In the rare case of failure, another method which we will describe later will allow us to recover in most cases.

11.8 CCD Response of a Single Collision

After CCD provides us with the collisions that occurred between candidate states, we need to resolve these collisions. Particles that were fully constrained are considered to have infinite mass. The vertices of kinematic geometry features can then be thought of as fully constrained since we cannot alter their motion with our response. Instead of operating directly on the velocities of the vertices, we alter their end positions during each iteration of our response algorithm. Once the entire response algorithm has completed, any particles with altered positions will have their velocity states changed to reflect this to be consistent with these altered positions.

To resolve a single continuous collision, we will follow the notation used to described inelastic projection described from Harmon et al. (2008). Although their work focuses on a multiple simultaneous contacts, the math provides a compact form for resolving a single contacts as well. Our discussion of Harmon et al. (2008) will also vary slightly since we choose to adjust the end positions instead of the candidate velocities. This allows our CCD detection and response module to have knowledge of only the two position states.

Let’s consider the vertex-face collision case. The displacement of each vertex in the time interval is described by \( \Delta x_i = x_e - x_s \). Let’s describe the displacement of all the vertices involved in a collision by the configuration space vector \( \mathbf{q} \). We define a scalar constraint equation to describe the normal relative displacement of our contact points in equation (11.9). We orient the collision normal at the time of impact, \( \hat{n}_c \), so that equation \( C(\mathbf{q}) < 0 \) since we know the relative normal motion must be negative for a collision to occur.

\[
C(\mathbf{q}) = \hat{n}_c \cdot [\Delta x_1 - (u\Delta x_2 + v\Delta x_3 + w\Delta x_4)]
\] (11.9)

The gradient wrt. \( \mathbf{q} \) of our constraint equation becomes:

---

7When processing a single contact, these expressions might appear intimidating, but they reduce to simple summations.
\[ \nabla C = [\hat{n}_c, -u\hat{n}_c, -v\hat{n}_c, -w\hat{n}_c] \]  

(11.10)

Continuing the derivation in Harmon et al. (2008), we arrive at equation (11.11). The right hand side describes the desired change we want in the normal relative motion. Once we solve for \( \lambda \), we update our end positions with equation (11.12). Hard constraints from our dynamics solution can be respected, by multiplying \( M^{-1} \) with \( S_i \) and incorporating it into both of these equations.³

\[ \nabla C M^{-1} \nabla C^T \lambda = \nabla C q \]  

(11.11)

\[ x'_{ie} = x_{ie} - M^{-1} \nabla C^T \lambda \]  

(11.12)

At this point, all normal relative motion between the points in contact will vanish. Let \( x_s \) be a vector of the vertex start positions and \( d_s = \nabla C x_s \) define the initial relative normal distance. We want all relative normal motion to stop when \( d_s \leq d_{offset} \). When \( d_s > d_{offset} \) we must alter the RHS of equation (11.11). If we want the relative normal motion to stop at \( d_{offset} \), we want a post-correction normal displacement magnitude = \( d_{offset} - d_s \). The desired change between our pre-correction and post-correction normal displacement magnitude defines our right hand side and is given by \( \nabla C q + \nabla C x_s - d_{offset} \).

I do not recommend modifying the RHS for trying to achieve post-correction normal offset greater than the initial one. Compensating for the collision offset would add energy, and it is probably best to model a fully dissipative response for deformable bodies. Applying inelastic projection to handle edge-edge collisions is straightforward. If multiple simultaneous contacts are found, this can be extended to the technique in Harmon et al. (2008). It would result in a linear system of size \( N_c \times N_c \) where \( N_c \) is the number of simultaneous contacts.

11.9 Resolving CCD Collisions in Chronological Order

In a single time step, there can be many collisions between our two position states, but one cannot process them all at once. Some of the found collisions won’t occur if others are resolved first and new collisions can be formed by the resolution of others. We present a summary of our resolution scheme in the pseudo-code below:

³If the point of contact on both sides is closer to the constrained particles on each side than the unconstrained ones, the resulting deformation from the response can be quite large. Special care must be taken in these situations.
bool HandleContinuousCollisions(CollisionDetector, maxIters)

CollisionDetector->DetectInitialCollisions()
collisions = CollisionDetector->GetResults()
iter = 0
bool collisionFree = true
bool allRegionsResolved = false
if not collisions.empty():
    collisionFree = false
    GatherCollisionsPerDynamicVertex(collisions)
    CreateContactRegions(collisions)

while !collisionFree and iter < maxIters:
    // Solve first contacts in each disjoint regions
    // and update the end position state marking
    // incident mesh features for the BVH update.
    SolveContactRegions()
    iters++
    allRegionsResolved = AreAllContactRegionsCollisionFree()
    // Steps for incremental detection
    CollisionDetector->UpdateBVHsWithMarkedFeatures()
    CollisionDetector->PerformIncrementalDetection()
    CollisionDetector->GetResults(newCollisions)

    // Determine whether collisions in newCollisions
    // are genuinely new or are already represented
    // in the contact regions. Remove already discovered
    // collisions from newCollisions.
    ProcessContacts(newCollisions)
    if not newCollisions.empty():
        // Contact regions may just expand
        // or expand and merge with others
        ExpandAndMergeContactRegions(newCollisions)
    else if allRegionsResolved:
        collisionFree = true

return collisionFree

HandleContinuousCollisions takes in our collision detection module and a maximum number of iterations to perform. We perform the initial collision detection pass in line 3. If any collisions were found, we gather the collisions per dynamic vertex and store them chronological order in line 10. We create groups of contacts, called ContactRegions, that are connected by shared vertices including ones with kinematic geometry. We compute and apply our response for the earliest contacts in each region in SolveContactRegions on line 17. Contact regions are solved independently in parallel each iteration. After applying the response, AreAllContactRegionsCollisionFree sweeps over the contact regions to see if all of their contained contacts have been resolved. When a collision in the region still occurs we update its Time of Impact(TOI), and barycentric coordinates.

Even when AreAllContactRegionsCollisionFree indicates all of our contacts have been resolved, we are not done because the response may have generated new collisions.
In lines 20-23 an incremental detection pass is then performed to discover any new contacts. Our incremental detection consists of first marking all the leaf level nodes of our BVHs as clean, and then labeling the nodes that contain a feature with a vertex that has moved by the last response iteration as dirty. We propagate the dirty flag up the tree from the leaves when updating the BVH extents. During the incremental detection pass we can then ignore descending down any pairs that are both clean.

Inside ProcessContacts, line 27, we examine if the additional contacts should cause contact regions to be expanded or merged. A new collision should always fall into either of those categories. It is the responsibility of the ContactRegions to keep the unresolved collisions they contained in chronological order. We then repeat this process until all the collisions have bee resolved or we have exceeded our maxIters. A dynamic vertex being pinched between two opposing kinematic surfaces sometimes cannot be resolved with this method. We will discuss our remedy to that problematic situation and others in §11.10.

If all collisions have not been resolved by maxIters, the inelastic projection described in Harmon et al. (2008) can be used to halt all normal relative motion in a contact region. This extra failsafe method is a good idea for systems that do not implement a means for recovery. Fitz has a means of recovery for some intersection cases, which will begin to discuss shortly. While CCD and the associated response can make sure collisions aren’t missed due to temporal aliasing, there is one drawback. The response is completely decoupled from the dynamics of the material model and excessive deformation may occur as a result.

11.10 Global Intersection Analysis

Character animation often leads to situations where the kinematic geometry intersects leaving the simulated surface nowhere to go and can easily lead to constrained situations that can cause it to intersect. To handle these situations, the Flypapering technique was introduced in the Untangling Cloth paper Baraff and Witkin (2003) 9. The Global Intersection Analysis, (GIA) algorithm was also introduced in this paper, which we continue to rely heavily on. We give a brief summary of the GIA algorithm in this section and discuss its application to proximity and CCD in section §11.10.

Edge-face intersection pairs are first found using our BVH. Closed contours of intersection are then computed by traversing the intersections and using mesh connectivity. A flood-fill algorithm then colors the regions on both sides of each contour. The GIA algorithm makes the reasonable assumption that the region with the smaller area is the region of intersection, though this is certainly not true in pathological cases. The algorithm also does not support surfaces that are non-manifold. Since not all intersection configurations lead to a closed contour of intersection, we still label the mesh features that are involved in intersection.

9As previously noted we now run a volume simulation to deform the character geometry, Wong et al. (2018), before running cloth to remove a majority of Flypapering cases.
11.11 Using GIA with Proximity and CCD

Fitz performs GIA before proximity queries and CCD. GIA is applied to all mesh pairs in the simulator, including the kinematic meshes. We also apply it to the kinematic meshes at the end of the time step.

Kinematic objects are assumed to have meshes that are oriented and closed. When a particle of a deformable body is detected beneath the surface of a kinematic object, a natural first response would be to push it to the desired collision offset of the face it is underneath. When objects are thin or penetrations are deep, this can easily result in the particle being pushed out the wrong side of the surface. This will lead to tangling from which a simulation would rarely ever recover. To alleviate this issue, we look to see if the face we plan to eject through has been labeled as being in a GIA contour, or is a part of an intersection. Only if it is labeled do we deliver the response. This heuristic will not handle all cases, but in practice, particles closer to the intersection region are pushed out first. That response then causes connected particles that are in deeper penetration to emerge from the correct side of the surface.

When two dynamic surfaces are intersecting, we use the GIA coloring information to change our penalty response. In the case of cloth, we do not have a notion of an inside and outside. When a colored particle has a proximity contact with a face of the same color, we instruct our penalty force to pull it through to the appropriate offset distance on the other side. This follows the approach to untangling cloth intersections described in Baraff and Witkin (2003).

The untangling response just described will be thwarted by our CCD pass unless it also uses information from GIA. When CCD discovers a contact between two surfaces that have been labeled to be in intersection by GIA, we ignore the collision so that it will not prevent the penalty response from resolving the situation. Once the GIA intersection is resolved, CCD can do its best at preventing intersection from recurring. By coupling GIA with CCD we can safely allow for recovery without compromising the robustness.
of our CCD with any local heuristics\textsuperscript{10}.

There can be scenarios where a dynamic contact cannot be resolved, because it is trapped between kinematic surfaces that are passing through each other. We use the kinematic intersection states of the kinematic object to ignore continuous collisions that involve kinematic features that are pinched, becoming pinched, or becoming unpinched.

Instead of attempting to engineer a collision strategy that is never allowed to fail, \textit{Fizt} has taken an approach of assuming failure is inevitable in the face of production work. What is important is to have the ability to recover from any failures gracefully. While our implementations of GIA, proximity queries and continuous collision could be more robust, by working together they are able to handle many demanding production scenarios.

\section*{11.12 Things Not Covered}

\textit{Fizt} has evolved at Pixar over 20 years and there are many features and topics that have not been discussed. The list that follows is far from complete but is meant to provide some more information and perspective.

\begin{itemize}
  \item A myriad of custom implicit forces and constraints exist to give the artists complex control during asset tailoring and shot work.
  \item We have many controls to change the rest shape of the cloth over time to deal with drastic character stretch/squash and to help facilitate wrinkles in certain regions.
  \item Particle masses are not constant to support a consistent look despite drastic changes in rest shape area. This is one reason why some newer and faster techniques that exploit a single system factorization are not applicable to us.
  \item Our cloth material model is rooted in \textit{Baraff and Witkin} (1998), but has become more complex to allow the artists the control they’ve needed to provide a wide variety of looks over many films\textsuperscript{11}.
  \item \textit{Fizt} has the ability to cope with fast motion by dialing in the amount of rotation and translation experienced by the simulation relative to the motion of a frame on the character.
\end{itemize}

There are still a lot of robustness and performance improvements that can be made to \textit{Fizt} and for many it may seem that we are slow to adopt the latest advances in the literature. If physical accuracy was our primary goal, then this would certainly be true. Our production work however prioritizes stability\textsuperscript{12}, speed and artist directability above physical accuracy. Only physical plausibility is required.

\textsuperscript{10}Many were tried before discovering GIA is a far more robust approach.
\textsuperscript{11}Perhaps in future versions of this course we will discuss it in detail.
\textsuperscript{12}Numerical stability as well as code stability.
There is a lot more to our simulation pipeline than the simulator itself. Our tailoring/asset creation tools exist in Maya, but have had many years of custom development which is still ongoing. Our pre-roll system to simulate characters into their shot pose is another tool that our artists constantly rely on. To evolve and support a system this large requires the hard work of many people from Pixar’s Tools-Sim and Research teams.
Appendix A

Table of Symbols

We adhere as closely as possible to the following convention:

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>$a$</td>
<td>unbolded, lower-case letter is a scalar</td>
</tr>
<tr>
<td>$A$</td>
<td>unbolded, upper-case letter is (still) a scalar</td>
</tr>
<tr>
<td>$\mathbf{a}$</td>
<td>bold, lower-case letter is a vector</td>
</tr>
<tr>
<td>$\mathbf{A}$</td>
<td>bold, upper-case letter is a matrix</td>
</tr>
<tr>
<td>$a \in \mathbb{R}^n$</td>
<td>$a$ is an $n$-dimensional vector</td>
</tr>
<tr>
<td>$A \in \mathbb{R}^{m \times n}$</td>
<td>$A$ is an $m$ by $n$ matrix</td>
</tr>
<tr>
<td>$A \in \mathbb{R}^{m \times n \times o \times \ldots}$</td>
<td>$A$ is a higher-order (3rd and above) tensor</td>
</tr>
</tbody>
</table>

We use the following operators and special matrices:

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>$| \cdot |_2$</td>
<td>2-norm (§C.1.1)</td>
</tr>
<tr>
<td>$| \cdot |_F$</td>
<td>Frobenius norm (§C.1.2)</td>
</tr>
<tr>
<td>$A : B$</td>
<td>Double-contraction of matrices $A$ and $B$ (§C.3)</td>
</tr>
<tr>
<td>$\mathbf{A} : \mathbf{B}$</td>
<td>Double-contraction of higher-order tensor $\mathbf{A}$ and matrix $\mathbf{B}$ (§3.2,§C.3)</td>
</tr>
<tr>
<td>$\mathbf{A} \otimes \mathbf{B}$</td>
<td>A Kronecker product between two matrices (§C.5)</td>
</tr>
<tr>
<td>$\text{tr} \ A$</td>
<td>trace of matrix $A$, i.e. the sum of its diagonal entries, $\sum_{i=0}^{n} a_{ii}$ (§C.2)</td>
</tr>
<tr>
<td>$\text{det} \ A$</td>
<td>determinant of matrix $A$</td>
</tr>
<tr>
<td>$\text{vec}(\cdot)$</td>
<td>flattening or vectorization of a matrix or tensor (§3)</td>
</tr>
<tr>
<td>$0$</td>
<td>The zero matrix, a.k.a. the null matrix. Nothing but zeros</td>
</tr>
<tr>
<td>$I$</td>
<td>The identity matrix. A diagonal matrix of all ones</td>
</tr>
</tbody>
</table>
These symbols are reserved for the following deformation-specific phenomena:

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>(\bar{x})</td>
<td>rest vertex, before deformation</td>
</tr>
<tr>
<td>(x)</td>
<td>deformed vertex</td>
</tr>
<tr>
<td>(\mathbf{F} = \begin{bmatrix} f_{00} &amp; f_{01} &amp; f_{02} \ f_{10} &amp; f_{11} &amp; f_{12} \ f_{20} &amp; f_{21} &amp; f_{22} \end{bmatrix} = \begin{bmatrix} f_0 \ f_1 \ f_2 \end{bmatrix})</td>
<td>deformation gradient, in 3D</td>
</tr>
<tr>
<td>(\mathbf{F} = \begin{bmatrix} f_{00} &amp; f_{01} \ f_{10} &amp; f_{11} \end{bmatrix})</td>
<td>deformation gradient, in 2D</td>
</tr>
<tr>
<td>(\mathbf{C})</td>
<td>right Cauchy-Green tensor, (\mathbf{C} = \mathbf{F}^T \mathbf{F})</td>
</tr>
<tr>
<td>(\mathbf{E})</td>
<td>Green’s strain, (\mathbf{E} = \mathbf{F}^T \mathbf{F} - \mathbf{I})</td>
</tr>
<tr>
<td>(t)</td>
<td>translation</td>
</tr>
<tr>
<td>(a)</td>
<td>anisotropic fiber direction from §8</td>
</tr>
<tr>
<td>(J)</td>
<td>(J = \text{det} \mathbf{F}), the current relative volume of (\mathbf{F})</td>
</tr>
<tr>
<td>(g_1)</td>
<td>flattened gradient of (I_1) invariant from §5.5</td>
</tr>
<tr>
<td>(g_2)</td>
<td>flattened gradient of (I_2) invariant from §5.5</td>
</tr>
<tr>
<td>(g_j \equiv g_3)</td>
<td>flattened gradient of (J \equiv I_3) from §4.2.2.1</td>
</tr>
<tr>
<td>(H_1)</td>
<td>flattened Hessian of (I_1) invariant from §5.5</td>
</tr>
<tr>
<td>(H_2)</td>
<td>flattened Hessian of (I_2) invariant from §5.5</td>
</tr>
<tr>
<td>(H_J \equiv H_3)</td>
<td>flattened Hessian of (J \equiv I_3) from §4.2.2.2</td>
</tr>
<tr>
<td>(\mathbf{R})</td>
<td>rotation matrix from the polar decomposition (\mathbf{F} = \mathbf{RS})</td>
</tr>
<tr>
<td>(\mathbf{S})</td>
<td>scaling matrix from the polar decomposition (\mathbf{F} = \mathbf{RS})</td>
</tr>
<tr>
<td>(\mathbf{U})</td>
<td>left singular vectors of the SVD of (\mathbf{F} = \mathbf{U} \Sigma \mathbf{V}^T)</td>
</tr>
<tr>
<td>(\Sigma = \begin{bmatrix} \sigma_x &amp; 0 &amp; 0 \ 0 &amp; \sigma_y &amp; 0 \ 0 &amp; 0 &amp; \sigma_z \end{bmatrix})</td>
<td>singular values of the SVD of (\mathbf{F} = \mathbf{U} \Sigma \mathbf{V}^T)</td>
</tr>
<tr>
<td>(\mathbf{V})</td>
<td>right singular vectors of the SVD of (\mathbf{F} = \mathbf{U} \Sigma \mathbf{V}^T)</td>
</tr>
<tr>
<td>(\hat{\mathbf{x}} = \begin{bmatrix} 0 &amp; -x_2 &amp; x_1 \ x_2 &amp; 0 &amp; -x_0 \ -x_1 &amp; x_0 &amp; 0 \end{bmatrix})</td>
<td>the cross product matrix of a vector (\mathbf{x})</td>
</tr>
<tr>
<td>(\hat{\mathbf{n}})</td>
<td>a normalized vector in §9 and §10</td>
</tr>
</tbody>
</table>
Appendix B

Useful Identities

Invariants:

\[ I_C = \| \mathbf{F} \|_F^2 \] \hspace{1cm} (B.1)
\[ II_C = \| \mathbf{F}^T \mathbf{F} \|_F^2 \] \hspace{1cm} (B.2)
\[ III_C = \det (\mathbf{F}^T \mathbf{F}) \] \hspace{1cm} (B.3)
\[ IV_C = \mathbf{a}^T \mathbf{C} \mathbf{a} \] \hspace{1cm} (B.4)
\[ V_C = \mathbf{a}^T \mathbf{C}^T \mathbf{a} \] \hspace{1cm} (B.5)
\[ I_1 = \text{tr} (\mathbf{R}^T \mathbf{F}) = \text{tr} \mathbf{S} \] \hspace{1cm} (B.6)
\[ I_2 = I_C = \| \mathbf{F} \|_F^2 \] \hspace{1cm} (B.7)
\[ I_3 = J = \det \mathbf{F} \] \hspace{1cm} (B.8)
\[ I_4 = \mathbf{a}^T \mathbf{S} \mathbf{a} \] \hspace{1cm} (B.9)
\[ I_5 = IV_C = \mathbf{a}^T \mathbf{F}^T \mathbf{F} \mathbf{a} \] \hspace{1cm} (B.10)

Trace and Frobenius identities:

\[ \text{tr} (\alpha \mathbf{A}) = \alpha \text{tr} \mathbf{A} \] \hspace{1cm} (B.11)
\[ \text{tr} (\mathbf{A} + \mathbf{B}) = \text{tr} \mathbf{A} + \text{tr} \mathbf{B} \] \hspace{1cm} (B.12)
\[ \text{tr} (\mathbf{A} - \mathbf{B}) = \text{tr} \mathbf{A} - \text{tr} \mathbf{B} \] \hspace{1cm} (B.13)
\[ \text{tr} (\mathbf{A}^T \mathbf{B}) = \mathbf{A} : \mathbf{B} = \text{vec} (\mathbf{A})^T \text{vec} (\mathbf{B}) \] \hspace{1cm} (B.14)
\[ \| \mathbf{A} + \mathbf{B} \|_F^2 = \| \mathbf{A} \|_F^2 + \| \mathbf{B} \|_F^2 + 2 \text{tr} \mathbf{A}^T \mathbf{B} \] \hspace{1cm} (B.15)
\[ \| \mathbf{A} - \mathbf{B} \|_F^2 = \| \mathbf{A} \|_F^2 + \| \mathbf{B} \|_F^2 - 2 \text{tr} \mathbf{A}^T \mathbf{B} \] \hspace{1cm} (B.16)
B. Useful Identities

Invariant identities and derivatives:

\[
\frac{\partial I_1}{\partial F} = \frac{\partial \text{tr } S}{\partial F} = R \quad \text{(B.17)}
\]

\[
\frac{\partial I_2}{\partial F} = \frac{\partial I_C}{\partial F} = \frac{\partial \| F \|^2}{\partial F} = 2F \quad \text{(B.18)}
\]

\[
\frac{\partial I_C}{\partial F} = \frac{\partial \| F^T F \|^2}{\partial F} = 4FF^T F \quad \text{(B.19)}
\]

\[
\frac{\partial I_3}{\partial F} = \frac{\partial J}{\partial F} = \begin{bmatrix} f_{11} & -f_{10} \\ -f_{01} & f_{00} \end{bmatrix} \quad \text{(In 2D)} \quad \text{(B.20)}
\]

\[
\frac{\partial I_3}{\partial F} = \frac{\partial J}{\partial F} = \begin{bmatrix} f_1 \times f_2 & f_2 \times f_0 & f_0 \times f_1 \end{bmatrix} \quad \text{(In 3D)} \quad \text{(B.21)}
\]

\[
\frac{\partial I_5}{\partial F} = \frac{\partial IV_C}{\partial F} = 2Faa^T \quad \text{(B.22)}
\]

\[
\text{vec} \left( \frac{\partial^2 I_2}{\partial F^2} \right) = H_2 = H_I = 2I_{9 \times 9} \quad \text{(B.23)}
\]

\[
\text{vec} \left( \frac{\partial^2 I_3}{\partial F^2} \right) = H_J = \begin{bmatrix} 0 & 0 & 0 & 1 \\ 0 & 0 & -1 & 0 \\ 0 & -1 & 0 & 0 \\ 1 & 0 & 0 & 0 \end{bmatrix} \quad \text{(In 2D)} \quad \text{(B.24)}
\]

\[
\text{vec} \left( \frac{\partial^2 I_3}{\partial F^2} \right) = H_J = \begin{bmatrix} \mathbf{0}_{3 \times 3} & -\hat{f}_2 & \hat{f}_1 \\ \hat{f}_2 & \mathbf{0}_{3 \times 3} & -\hat{f}_0 \\ -\hat{f}_1 & \hat{f}_0 & \mathbf{0}_{3 \times 3} \end{bmatrix} \quad \text{(In 3D)} \quad \text{(B.25)}
\]

\[
\text{vec} \left( \frac{\partial^2 I_5}{\partial F^2} \right) = \text{vec} \left( \frac{\partial^2 IV_C}{\partial F^2} \right) = 2aa^T \otimes I \quad \text{(B.26)}
\]

\[
\text{vec} \left( \frac{\partial^2 I_C}{\partial F^2} \right) = H_{II} = 4 \left( \mathbf{I}_{3 \times 3} \otimes FF^T + F^T F \otimes \mathbf{I}_{3 \times 3} + D \right) \quad \text{(B.27)}
\]

\[
D = \begin{bmatrix} f_0 f_0^T & f_1 f_0^T & f_2 f_0^T \\ f_0 f_1^T & f_1 f_1^T & f_2 f_1^T \\ f_0 f_2^T & f_1 f_2^T & f_2 f_2^T \end{bmatrix} \quad \text{(B.28)}
\]
B. Useful Identities

\[
\frac{\partial \log(I)}{\partial F} = \frac{1}{I} \frac{\partial I}{\partial F} = \frac{1}{J} \frac{\partial J}{\partial F}
\]

\[
\text{tr}\left(F^T F\right) = \|F\|^2_F = F : F = \sum_i \sum_j F^2_{ij}
\]

\[
F^T R = S
\]

\[
\frac{\partial \|R\|^2_F}{\partial F} = \frac{\partial \text{tr}\left(R^T R\right)}{\partial F} = \frac{\partial \text{tr} I}{\partial F} = 0
\]

\[
\|F^{-1}\|^2_F = \frac{I_2}{(I_3)^2} \quad \text{(In 2D)}
\]

\[
\|F^{-1}\|^2_F = \frac{1}{4}\left(\frac{I_2^2 - I_2}{I_3}\right) - 2\frac{I_1}{I_3} \quad \text{(In 3D)}
\]
Appendix C

Notation

C.1 Norms

A norm is a way to boil the entries of a vector or matrix down to a single scalar, or score. Often it’s impossible to stare at the wall of numbers that constitute a vector or matrix and understand what’s going on, so it’s useful to somehow distill the “essence” of the vector or matrix into a single number. Often you are looking for different “essences” though, so to capture each of these, there are a variety of different norms.\(^1\)

C.1.1 The 2-Norm

The vector norm we use most often is the 2-norm. For some vector \(x\),

\[
x = \begin{bmatrix}
x_0 \\
x_1 \\
x_2 \\
\vdots \\
x_{n-1}
\end{bmatrix}
\]

the 2-norm \(\| \cdot \|_2\) is the square root of the squared sum:

\[
\|x\|_2 = \sqrt{\sum_{i=0}^{n-1} x_i^2}.
\]

If there are only two entries in \(x\), such as \(x = \begin{bmatrix} x_0 \\ x_1 \end{bmatrix}\), then we get the familiar Euclidean distance:

\[
\|x\|_2 = \sqrt{x_0^2 + x_1^2}.
\]

\(^1\)Another way to think of a norm is “average over all the members”. So when we talk of societal norms, it is the overall behavior that we expect from averaging over the members of that society.
The 2-norm is just a generalization of this style of measurement.

### C.1.2 The Frobenius Norm

The matrix norm we will use the most often is the Frobenius norm. It is exactly the same as the 2-norm, except that it is defined over all the entries of a matrix. For some matrix $A$,

$$A = \begin{bmatrix}
a_{00} & a_{01} & \cdots & a_{0(n-1)} \\
a_{10} & a_{11} & \cdots & a_{1(n-1)} \\
\vdots & \vdots & \ddots & \vdots \\
a_{(m-1)0} & a_{(m-1)1} & \cdots & a_{(m-1)(n-1)}
\end{bmatrix}$$

the Frobenius norm $\| \cdot \|_F$ is (again) the square root of the squared sum:

$$\|A\|_F = \sqrt{\sum_{i=0}^{m-1} \sum_{j=0}^{n-1} a_{ij}^2}.$$  

(C.5)

It is ugly to have the square root enveloping everything, and it also introduces pesky $1/2$ factors when you take derivatives, so usually we use the squared Frobenius norm:

$$\|A\|^2_F = \sum_{i=0}^{m-1} \sum_{j=0}^{n-1} a_{ij}^2.$$  

(C.6)

Notationally, this version only differs from $\| \cdot \|_F$ with the 2 in the super-script, $\| \cdot \|^2_F$, so it can be easy to get confused. Just remember that we’re almost always going to use the squared Frobenius norm. In fact, as I write this, I’m having trouble thinking of a single instance where we will need to use the un-squared $\| \cdot \|_F$.

I’ll wrap up discussion of this norm with two observations.

- The 2-norm is literally a vector version of the Frobenius norm. You can write $\|x\|_F$ instead of $\|x\|^2_F$ and they mean the exact same thing.
- The Frobenius norm is weird. With the 2-norm, we have some notion of measuring the magnitude of a vector. With the Frobenius norm, we appear to be arbitrarily smashing all the entries of a matrix together into one scalar.

Doesn’t anybody care which row and column these entries came from? Surely their ordering in the matrix should count for something? Maybe there is a matrix norm out there that cares, but not Frobenius. Despite this apparent blindness, the Frobenius norm gets the job done in many useful scenarios.
C.2 Matrix Trace

The trace of a matrix, $\text{tr } A$, is the sum of its diagonal entries:

$$\text{tr } A = \sum_{i=0}^{n-1} a_{ii}. \quad (C.7)$$

This seems like an arbitrary operation to define for a matrix. So if I swap two of the rows, then I get a different trace? How is this at all useful? While this is indeed odd, the interesting and surprising property of the matrix trace is that $\text{tr } A$ is the sum of the eigenvalues of $A$.

This is not obvious at all! The eigenvalues are supposed to be some fundamental, atomic property that can only be acquired after a perilous journey involving Householder reductions, shifted-QR factorizations, or other exotic numerical analysis weapons. And yet, the sum of these buried jewels can somehow be plucked right off of the surface of the matrix. Huh.

C.3 Matrix Double-Contractions

The double-contraction, denoted with a ‘;’, is a generalized version of the dot product for vectors. With vectors, we multiply together the corresponding entries in vector, and then glob them all together in a summation:

$$x^T y = \begin{bmatrix} x_0 \\ x_1 \\ x_2 \end{bmatrix}^T \begin{bmatrix} y_0 \\ y_1 \\ y_2 \end{bmatrix} = x_0y_0 + x_1y_1 + x_2y_2. \quad (C.8)$$

The double-contraction does the exact same thing, but for the entries of two matrices:

$$A : B = \begin{bmatrix} a_0 & a_2 \\ a_1 & a_3 \end{bmatrix} \begin{bmatrix} b_0 \\ b_1 \end{bmatrix} = a_0b_0 + a_1b_1. \quad (C.9)$$

The dimensions of the two matrices need to be the same, otherwise the operations is undefined (and you screwed up somewhere).

This operation may look odd to you. Similar to the Frobenius norm, it chucks all the entries of the matrices into the same pile, insensitive to whether they have important-looking VIP status like being along the diagonal. It is indeed odd, but this is the definition nevertheless, and despite its status-obliviousness, it is still quite useful.

C.4 Outer Products

You’ve probably been sent here by §4.2.2.1. We’re used to seeing inner products, a.k.a. dot products like $x^T y = \alpha$ where two vectors $x$ and $y$ are smooshed together in such a way
that they’re demoted to the lowly scalar $\alpha$. In other words, inner products take two rank-one quantities (vectors) and bash them down to rank-zero quantity (a scalar).

The outer product goes in the other direction: $xx^T = A$. A pair of rank-one quantities join forces to generate a bigger, rank-two result (a matrix). This product doesn’t appear quite as often as the inner product, so while it probably did show up in your linear algebra class at some point, there’s a good chance that you haven’t seen it since you crammed for the midterm. So, here’s a refresher.

If we have two vectors,

$$x = \begin{bmatrix} a \\ b \\ c \end{bmatrix}, \quad y = \begin{bmatrix} d \\ e \\ f \end{bmatrix},$$  \hspace{1cm} \text{(C.10)}

then their outer product is

$$xy^T = \begin{bmatrix} a \\ b \\ c \end{bmatrix} \begin{bmatrix} d \\ e \\ f \end{bmatrix}^T = \begin{bmatrix} ad & ae & af \\ bd & be & bf \\ cd & ce & cf \end{bmatrix}$$  \hspace{1cm} \text{(C.11)}

We place a copy of $x$ into each column, and in turn, scale the copy by each entry in $y$.

Why is this helpful? Inner products have intuitive interpretations like projecting one vector onto another, but what’s the point of this baroque and arbitrary-looking operation?

The outer product form often shows up when you see what’s really going on inside the primordial morass of numbers that form a matrix. We can look at the eigendecomposition of a symmetric $n \times n$ matrix,

$$A = QQ^T,$$  \hspace{1cm} \text{(C.12)}

where $Q$ is the matrix of eigenvectors, $q_i$, and $\Lambda$ is a matrix of the eigenvalues, $\lambda_i$:

$$Q = \begin{bmatrix} q_0 & q_1 & \cdots & q_{n-1} \end{bmatrix}, \quad \Lambda = \begin{bmatrix} \lambda_0 & & & \\ & \lambda_1 & & \\ & & \ddots & \\ & & & \lambda_{n-1} \end{bmatrix}.$$  \hspace{1cm} \text{(C.13)}

We can see a hint of the $xy^T$ outer product form by looking at $QQ^T$. Both have a transpose all the way on the right hand side. This is not a coincidence, as the eigendecomposition of a matrix can indeed be written as a sum of outer products:

$$A = Q\Lambda Q^T = \lambda_0 q_0 q_0^T + \lambda_1 q_1 q_1^T + \cdots + \lambda_n q_n q_n^T$$  \hspace{1cm} \text{(C.14)}

$$A = \sum_{i=0}^{n-1} \lambda_i q_i q_i^T.$$  \hspace{1cm} \text{(C.15)}

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Thus, one way to look at a matrix $A \in \mathbb{R}^{n \times n}$ is as a scaled sum of $n$ outer products. What does this mean? Say you went ahead and just lopped off the last ten terms in the sum:

$$A_{\text{oops}} = \sum_{i=0}^{n-11} \lambda_i q_i q_i^T.$$  \hspace{1cm} (C.16)

You would still get an $n \times n$ matrix out in the end. However, lurking inside that matrix would be rank-deficiency. You secretly don’t need the full complement of $n$ outer products to produce all the entries of $A_{\text{oops}}$.

In the extreme, you might only need one outer product to produce $A_{\text{oops}}$, making it a rank-one matrix. In that case, you don’t need to store the whole matrix $A_{\text{oops}}$, you can just carry around the one eigenpair, $(q_0, \lambda_0)$, and generate $A_{\text{oops}} = \lambda_0 q_0 q_0^T$ on-the-fly whenever you need it.

Conversely, imagine you’re looking at the eigendecompositions of matrices that appear somewhere in your code. The decomposition keeps telling you that it’s a rank-one matrix. In this case, your job as a numerical archeologist just got easier. Instead of trying to excavate the underlying structure of all $n \times n$ entries, you can now just focus on understanding where this one $q_0 \in \mathbb{R}^n$ came from. Your task just got quadratically easier.

### C.5 Kronecker Products

The Kronecker product is the matrix version of the outer product, though it irritatingly reverses the ordering of the operators. The Kronecker product of $A$ and $B$ is:

$$A \otimes B = \begin{bmatrix} a_{00} & a_{01} \\ a_{10} & a_{11} \end{bmatrix} \otimes \begin{bmatrix} b_{00} & b_{01} \\ b_{10} & b_{11} \end{bmatrix} = \begin{bmatrix} a_{00} B & a_{01} B \\ a_{10} B & a_{11} B \end{bmatrix} = \begin{bmatrix} a_{00} b_{00} & a_{00} b_{01} & a_{01} b_{00} & a_{01} b_{01} \\ a_{00} b_{10} & a_{00} b_{11} & a_{01} b_{10} & a_{01} b_{11} \\ a_{10} b_{00} & a_{10} b_{01} & a_{11} b_{00} & a_{11} b_{01} \\ a_{10} b_{10} & a_{10} b_{11} & a_{11} b_{10} & a_{11} b_{11} \end{bmatrix}$$

Whereas the outer product stacked together a bunch of scaled copies of $x$ from the left, the Kronecker product stacks scaled copies of $B$ from the right. I don’t make the rules here. Already I’m wondering where in these notes I messed up the ordering.

This product commonly appears with an identity matrix. If you have a matrix $A$, and you need to stamp copies of $A$ down the block-diagonal of a matrix, then the Kronecker
product is up to the task:

\[
\mathbf{I}_{2 \times 2} \otimes \mathbf{A} = \begin{bmatrix} 1 & 0 \\ 0 & 1 \end{bmatrix} \otimes \begin{bmatrix} a_{00} & a_{01} \\ a_{10} & a_{11} \end{bmatrix} = \begin{bmatrix} \mathbf{A} & \mathbf{0}_{2 \times 2} \\ \mathbf{0}_{2 \times 2} & \mathbf{A} \end{bmatrix}
\]

= \begin{bmatrix} a_{00} & a_{01} & 0 & 0 \\ a_{10} & a_{11} & 0 & 0 \\ 0 & 0 & a_{00} & a_{01} \\ 0 & 0 & a_{10} & a_{11} \end{bmatrix}.

If you need to spread each entry of \( \mathbf{A} \) across the block-diagonal of a matrix, reversing the two matrices gets the job done:

\[
\mathbf{A} \otimes \mathbf{I}_{2 \times 2} = \begin{bmatrix} a_{00} & a_{01} \\ a_{10} & a_{11} \end{bmatrix} \otimes \begin{bmatrix} 1 & 0 \\ 0 & 1 \end{bmatrix} = \begin{bmatrix} a_{00} \mathbf{I}_{2 \times 2} & a_{01} \mathbf{I}_{2 \times 2} \\ a_{10} \mathbf{I}_{2 \times 2} & a_{11} \mathbf{I}_{2 \times 2} \end{bmatrix}
\]

= \begin{bmatrix} a_{00} & 0 & a_{01} & 0 \\ 0 & a_{00} & 0 & a_{01} \\ a_{10} & 0 & a_{11} & 0 \\ 0 & a_{10} & 0 & a_{11} \end{bmatrix}.

These sorts of patterns tend to appear when you flatten out higher order tensors. So, we see them in §4.2.3 when computing \( \mathbf{H}_{II} \) and in §8.2.2.1 when computing \( \mathbf{H}_{IV} \equiv \mathbf{H}_5 \).
Appendix D

How to Compute \( F \), the Deformation Gradient

First we’ll look at an intuitive, geometric, purely linear-algebraic way to compute \( F \) on triangles and tetrahedra. Then we’ll look at a slightly more obtuse, calculus-based method that does the exact same thing using finite element basis functions. While this method is more involved and less intuitive, it will point the way to computing \( F \) on other primitives, such as hexahedra.

D.1 Computing \( F \) for 2D Triangles

Let’s say we have a triangle with vertices \( \bar{x}_0, \bar{x}_1, \) and \( \bar{x}_2 \). After deformation, these same vertices have become \( x_0, x_1, \) and \( x_2 \). How do we compute a \( \mathbb{R}^{2 \times 2} \) matrix \( F \) that describes the rotation and scaling that occurs to transform the points \( \bar{x}_i \) into \( x_i \)?

We specifically want to pull off any translations, so let’s deal with that first. Both triangles are floating off in space somewhere, so just to establish a common point of reference, let’s pull them both back to the origin. If they’re centered about the origin, then no relative translation needed to transform one into the other, and any remaining difference between the triangles must be due to rotation and scaling.

We’ll pull both triangles back to the origin by explicitly pinning \( \bar{x}_0 \) and \( x_0 \) to the origin (see Fig. D.1). The new vertices of our origin-centered rest-triangle become:

\[
\begin{align*}
\bar{x}_0 & \rightarrow \quad \bar{o}_0 = [0 \\ 0]^T \\
\bar{x}_1 & \rightarrow \quad \bar{o}_1 = \bar{x}_1 - \bar{x}_0 \\
\bar{x}_2 & \rightarrow \quad \bar{o}_2 = \bar{x}_2 - \bar{x}_0.
\end{align*}
\]
D. How to Compute $F$, the Deformation Gradient

Figure D.1.: Let’s eliminate the translation by pulling the rest triangle (left, black), back to the origin (left, red). We do the same thing to the deformed triangle (right, black and red). I have to redraw this figure so that $\bar{e}_s$ is replaced with $o_s$.

The vertices of the origin-centered deformed triangle are correspondingly:

$$x_0 \rightarrow o_0 = [0 \ 0]^T \quad (D.4)$$

$$x_1 \rightarrow o_1 = x_1 - x_0 \quad (D.5)$$

$$x_2 \rightarrow o_2 = x_2 - x_0. \quad (D.6)$$

Now we want to know what matrix $F$ will successfully rotate and scale all of our $\bar{o}_i$ vertices so that they become the $o_i$ vertices. In other words, we want the matrix $F$ satisfies these three equations:

$$F\bar{o}_0 = o_0 \quad F\bar{o}_1 = o_1 \quad F\bar{o}_2 = o_2. \quad (D.7)$$

The first equation, $F\bar{o}_0 = o_0$ is trivially satisfied by any matrix $F$, since both $\bar{o}_0$ and $o_0$ are all zeros.

Really we only need to worry about $F$ covering the other two cases by successfully producing $o_1$ and $o_2$ when provided with $\bar{o}_1$ and $\bar{o}_2$. If we write this desire down
D. How to Compute $\mathbf{F}$, the Deformation Gradient

explicitly, it’s a well-posed linear algebra problem:

$$
\mathbf{F} \begin{bmatrix}
\bar{x}_1 - \bar{x}_0 \\
\bar{x}_2 - \bar{x}_0
\end{bmatrix} =
\begin{bmatrix}
x_1 - x_0 \\
x_2 - x_0
\end{bmatrix}
$$

(D.8)

$$
\mathbf{F} \begin{bmatrix}
\bar{o}_1 \\
\bar{o}_2
\end{bmatrix} =
\begin{bmatrix}
o_1 \\
o_2
\end{bmatrix}
$$

(D.9)

$$
\mathbf{FD}_m = \mathbf{D}_s
$$

(D.10)

Computing the final $\mathbf{F}$ then becomes straightforward:

$$
\mathbf{F} = \mathbf{D}_s \mathbf{D}_m^{-1}.
$$

(D.11)

The $\mathbf{D}_m$ matrix is quite small, i.e. $2 \times 2$ in 2D, so there’s no need to be worried that taking its inverse might be expensive. The $\mathbf{D}_m$ matrix also only depends on the rest pose, so you can precompute its inverse once at startup, and then when you need to compute $\mathbf{F}$ at any later time, no new inverse computation is needed.

Here we follow the naming convention in Teran et al. (2005) and abbreviate things to $\mathbf{D}_m$ and $\mathbf{D}_s$, where $m$ denotes material coordinates and $s$ denotes spatial coordinates. In a perfect world we’d stick with our own naming convention and call these $\mathbf{D}_r$ and $\mathbf{D}_d$, for rest and deformed, but the $m$ and $s$ subscripts are popular enough in other papers that we’re going to stick to them here. Then you won’t get confused when you read these other papers (including some of mine).

One thing you may be wondering: we specifically decided to pin $\bar{x}_0$ to the origin. Could we pick $\bar{x}_1$ or $\bar{x}_2$? Does it matter? Will we get the same $\mathbf{F}$? Is one better than the other? It doesn’t actually matter, though you should work out a few examples to convince yourself of this fact.

One thing you might worry about is the conditioning of these elements, i.e. instead of a nice chubby triangle you have a sadly skinny one that is essentially a line. A poorly-conditioned element is known to ruin the stability of simulations, because divide-by-zeros occur when computing $\mathbf{F}$. This happens because $o_1$ and $o_2$ are nearly coincident and $\mathbf{D}_m^{-1}$ becomes rank-deficient. Picking a different $\bar{x}_i$ to the origin will not save your bacon.

D.2 Computing $\mathbf{F}$ for 3D Tetrahedra

For a 3D tetrahedron, there is a straightforward generalization. In this case, we want a $3 \times 3$ version of $\mathbf{F}$. We translate the rest and deformed versions to the origin again, and

---

1I think. I’m guessing here, but it seems quite likely.
D. How to Compute $F$, the Deformation Gradient

After observing that $F\bar{0}_0 = o_0$ is again trivial, we are left with a $3 \times 3$ formulation:

$$F \begin{bmatrix} \bar{x}_1 - \bar{x}_0 & \bar{x}_2 - \bar{x}_0 & \bar{x}_3 - \bar{x}_0 \end{bmatrix} = \begin{bmatrix} x_1 - x_0 & x_2 - x_0 & x_3 - x_0 \end{bmatrix} \quad (D.12)$$

$$F \begin{bmatrix} \bar{o}_1 & \bar{o}_2 & \bar{o}_3 \end{bmatrix} = \begin{bmatrix} o_1 & o_2 & o_3 \end{bmatrix} \quad (D.13)$$

$$FD_m = D_s \quad (D.14)$$

$$F = D_s D_{m}^{-1} \quad (D.15)$$

D.3 Computing $F$ the Finite Element Way

Now let’s do the exact same thing, but in a different and more opaque way. But why? In the end, there will be an obvious way to extend this approach to computing $F$ on squares (quads) and cubes (hexahedra).

D.3.1 Remember Barycentric Coordinates?

We are going to use the Graphics 101 (see e.g. Marschner and Shirley (2016)) concept of barycentric coordinates for this one. If you’ve done some finite element stuff, you will recognize these as basis or shape functions over the triangle, specifically hat functions. If you’re reading this though, you’re probably a graphics person and not a mechanical engineering person, so we’ll stick to barycentric coordinates. For our purposes, the two concepts are completely equivalent.

A barycentric coordinate system assigns an easy-to-use 2D coordinate to every point on the interior of a triangle. These are usually denoted $u$ and $v$, that are easy-to-use because they are always defined over $[0, 1]$. If we have some barycentric coordinate $b = [u \ v]^T$, we can then reconstruct any point $x(b)$ or $\bar{x}(b)$ on the interior of the triangle using a linear combination of the basis functions $\beta_i$:

$$x(b) = \sum_{i=0}^{2} x_i \beta_i(b) \quad \bar{x}(b) = \sum_{i=0}^{2} \bar{x}_i \beta_i(b) \quad (D.16)$$

The basis functions $\beta_i$ are:

$$\beta_0(b) = 1 - u - v \quad \beta_1(b) = u \quad \beta_2(b) = v \quad (D.17)$$

Each basis function $\beta_i$ is attached to its corresponding vertex $x_i$ (see Fig. D.2). It is equal to 1 right on top of the vertex, and then fades off to 0 as you approach the triangle edge that is opposite to that vertex.
D. How to Compute $F$, the Deformation Gradient

Figure D.2.: The basis functions are linear functions that are equal to one at their corresponding vertices and ramp linearly down to zero as the opposing edge is approached.

D.3.2 Computing $F$ for 2D Triangles, the Basis Function Way

Now let’s use these basis functions to compute the deformation gradient. Remember way back in Eqn. 2.2 that the deformed points were defined as

$$x = F\bar{x} + t,$$  \hspace{1cm} (D.18)

so we can retrieve the deformation gradient by taking $\frac{\partial x}{\partial \bar{x}}$ (we actually wrote it as $\frac{\partial \phi}{\partial \bar{x}}$, but it’s the same). We can now use the basis functions to compute this in a brute-force, no-intuition-to-cling-to, calculus-based way.

Remember from Eqn. D.16 that we can also write both $x$ and $\bar{x}$ in terms of the barycentric coordinate $b$. We can thus expand $\frac{\partial x}{\partial \bar{x}}$ using the chain rule to route it through the $b$-dependent version of $x$ and $\bar{x}$:

$$F = \frac{\partial x}{\partial \bar{x}} = \frac{\partial x}{\partial b} \frac{\partial b}{\partial \bar{x}} = \frac{\partial x}{\partial b} \left( \frac{\partial \bar{x}}{\partial b} \right)^{-1}. \hspace{1cm} (D.19)$$

If we then plug in our basis-function-based definitions of $x$ and $\bar{x}$ (Eqn. D.16), we get:

$$F = \frac{\partial x}{\partial b} \left( \frac{\partial \bar{x}}{\partial b} \right)^{-1}$$

$$= \frac{\partial}{\partial b} \left( \sum_{i=0}^{2} x_i \beta_i(b) \right) \frac{\partial}{\partial b} \left( \sum_{i=0}^{2} \bar{x}_i \beta_i(b) \right)^{-1}$$

$$= \left( \sum_{i=0}^{2} x_i \frac{\partial \beta_i}{\partial b} \right) \left( \sum_{i=0}^{2} \bar{x}_i \frac{\partial \beta_i}{\partial b} \right)^{-1}. \hspace{1cm} (D.20)$$
D. How to Compute $F$, the Deformation Gradient

Since $\mathbf{x}_i$ and $\bar{\mathbf{x}}_i$, don’t actually depend on the value of $b$, we can write the derivative purely in terms of $\beta_i$. This lack of dependence makes sense; we’re generating $\mathbf{x}(b)$ by interpolating over a set of $\mathbf{x}_i$. No matter what $b$ we send down the pipe, the set of $\mathbf{x}_i$ points we’re trying to interpolate over stays the same.

The next trick is that we can write down Eqn. D.20 in matrix form

$$F = \left( \sum_{i=0}^{2} \mathbf{x}_i \frac{\partial \beta_i(b)}{\partial b} \right) \left( \sum_{i=0}^{2} \bar{\mathbf{x}}_i \frac{\partial \beta_i(b)}{\partial b} \right)^{-1} = \mathbf{XH} \, (\mathbf{XH})^{-1}. \tag{D.21}$$

The matrices $\mathbf{X}$ and $\bar{\mathbf{X}}$ are just the deformed and rest vertices of the triangles packed together:

$$\bar{\mathbf{X}} = \begin{bmatrix} \bar{\mathbf{x}}_0 & \bar{\mathbf{x}}_1 & \bar{\mathbf{x}}_2 \end{bmatrix} \in \mathbb{R}^{3 \times 3} \tag{D.22}$$

$$\mathbf{X} = \begin{bmatrix} \mathbf{x}_0 & \mathbf{x}_1 & \mathbf{x}_2 \end{bmatrix} \in \mathbb{R}^{3 \times 3}. \tag{D.23}$$

The non-trivial component here is $\mathbf{H}$, the matrix of partials $\frac{\partial \beta_i(b)}{\partial b}$. We can write the matrix of partials out explicitly as:

$$\mathbf{H} = \begin{bmatrix} \frac{\partial \beta_0}{\partial \varepsilon} & \frac{\partial \beta_0}{\partial \varepsilon} \\ \frac{\partial \beta_1}{\partial \varepsilon} & \frac{\partial \beta_1}{\partial \varepsilon} \\ \frac{\partial \beta_2}{\partial \varepsilon} & \frac{\partial \beta_2}{\partial \varepsilon} \end{bmatrix}. \tag{D.24}$$

Fortunately, the $\beta_i$ functions are quite simple! So, the derivatives are really easy and this matrix works out to:

$$\mathbf{H} = \begin{bmatrix} -1 & -1 \\ 1 & 0 \\ 0 & 1 \end{bmatrix}. \tag{D.25}$$

Finally, and somewhat perversely, we point out that:

$$\mathbf{XH} = \begin{bmatrix} \bar{\mathbf{x}}_0 & \bar{\mathbf{x}}_1 & \bar{\mathbf{x}}_2 \end{bmatrix} \begin{bmatrix} -1 & -1 \\ 1 & 0 \\ 0 & 1 \end{bmatrix} = \begin{bmatrix} \bar{\mathbf{x}}_1 - \bar{\mathbf{x}}_0 & \bar{\mathbf{x}}_2 - \bar{\mathbf{x}}_0 \end{bmatrix} = \mathbf{D}_m \tag{D.26}$$

$$\mathbf{XH} = \begin{bmatrix} \mathbf{x}_0 & \mathbf{x}_1 & \mathbf{x}_2 \end{bmatrix} \begin{bmatrix} -1 & -1 \\ 1 & 0 \\ 0 & 1 \end{bmatrix} = \begin{bmatrix} \mathbf{x}_1 - \mathbf{x}_0 & \mathbf{x}_2 - \mathbf{x}_0 \end{bmatrix} = \mathbf{D}_s. \tag{D.27}$$
and therefore

\[ F = XH (XH)^{-1} \]  
\[ = D_s D^{-1}_m \]

(D.28)  
(D.29)

Argh! It’s exactly the same as the previous method we saw, but with a lot of basis function dreck and unnecessary derivatives to slug through! What was the point?!

Relax. You didn’t go through a bunch of math of nothing. As I said before, there is indeed a point: once you’ve gone through this process once, you can use the exact same method to turn the crank and compute \( F \) for primitives that aren’t triangles or tetrahedra.

### D.3.3 Computing \( F \) for 3D Tetrahedra, the Basis Function Way

Let’s turn the crank for a 3D tetrahedron. It contains 4 vertices, so we add a new index to \( \bar{x}(b) \) and \( \tilde{x}(b) \),

\[ x(b) = \sum_{i=0}^{3} x_i \beta_i(b) \quad \tilde{x}(b) = \sum_{i=0}^{3} \bar{x}_i \beta_i(b), \quad \text{(D.30)} \]

one more barycentric coordinate to \( b = [u \ v \ w]^T \) and one more basis function:

\[ \beta_0(b) = 1 - u - v - w \quad \beta_1(b) = u \quad \beta_2(b) = v \quad \beta_3(b) = w. \quad \text{(D.31)} \]

We’re still looking for

\[ F = XH (XH)^{-1}, \quad \text{(D.33)} \]

where

\[ \bar{X} = \begin{bmatrix} \bar{x}_0 & \bar{x}_1 & \bar{x}_2 & \bar{x}_3 \end{bmatrix} \in \mathbb{R}^{3 \times 4} \quad \text{(D.34)} \]

\[ X = \begin{bmatrix} x_0 & x_1 & x_2 & x_3 \end{bmatrix} \in \mathbb{R}^{3 \times 4}. \quad \text{(D.35)} \]

The matrix of partials becomes predictably larger and more unwieldy-looking,

\[ H = \begin{bmatrix} \frac{\partial \beta_0}{\partial u} & \frac{\partial \beta_0}{\partial v} & \frac{\partial \beta_0}{\partial w} \\ \frac{\partial \beta_1}{\partial u} & \frac{\partial \beta_1}{\partial v} & \frac{\partial \beta_1}{\partial w} \\ \frac{\partial \beta_2}{\partial u} & \frac{\partial \beta_2}{\partial v} & \frac{\partial \beta_2}{\partial w} \\ \frac{\partial \beta_3}{\partial u} & \frac{\partial \beta_3}{\partial v} & \frac{\partial \beta_3}{\partial w} \end{bmatrix}, \quad \text{(D.36)} \]
but as before, it works out to a simple result:

\[
H = \begin{bmatrix}
-1 & -1 & -1 \\
1 & 0 & 0 \\
0 & 1 & 0 \\
0 & 0 & 1 \\
\end{bmatrix}.
\]  
(D.37)

Then, just like before, we end up with

\[
\bar{X}H = \begin{bmatrix}
x_0 & x_1 & x_2 & x_3 \\
\end{bmatrix} \begin{bmatrix}
-1 & -1 & -1 \\
1 & 0 & 0 \\
0 & 1 & 0 \\
0 & 0 & 1 \\
\end{bmatrix} = \begin{bmatrix}
x_1 - x_0 & x_2 - x_0 & x_3 - x_0 \\
\end{bmatrix} = D_m
\]
(D.38)

\[
XH = \begin{bmatrix}
x_0 & x_1 & x_2 & x_3 \\
\end{bmatrix} \begin{bmatrix}
-1 & -1 & -1 \\
1 & 0 & 0 \\
0 & 1 & 0 \\
0 & 0 & 1 \\
\end{bmatrix} = \begin{bmatrix}
x_1 - x_0 & x_2 - x_0 & x_3 - x_0 \\
\end{bmatrix} = D_s
\]
(D.39)

and arrive at the same result:

\[
F = XH (XH)^{-1} = D_s D_m^{-1}.
\]  
(D.40)

\[
= D_s D_m^{-1}.
\]  
(D.41)

### D.3.4 Computing F for 2D Quads, the Basis Function Way

For completeness, should really include this.

### D.3.5 Computing F for 3D Hexahedra, the Basis Function Way

For a cube element, let’s begin by defining everything over a canonical cube that goes from \([1,1,1]\) to \([-1,-1,-1]\). We order the vertices of this cube like this:

\[
a = [-1,-1,-1] \\
b = [1,-1,-1] \\
c = [-1,1,-1] \\
d = [1,1,-1] \\
e = [-1,-1,1] \\
f = [1,-1,1] \\
g = [-1,1,1] \\
h = [1,1,1] \\
\]
The vertices are traversed in counter-clockwise order, starting with the square in the 
\( z = -1 \) plane, and followed by the one in the \( z = +1 \) plane. While it’s not really 
“barycentric”, we have a coordinate \( \mathbf{b} = [u \ v \ w]^T \) that we use to address every point 
inside the canonical space. The shape functions using \( \mathbf{b} \) are then:

\[
\begin{align*}
\beta_0(\mathbf{b}) &= (1 - u)(1 - v)(1 - w) \\
\beta_1(\mathbf{b}) &= (1 + u)(1 - v)(1 - w) \\
\beta_2(\mathbf{b}) &= (1 - u)(1 + v)(1 - w) \\
\beta_3(\mathbf{b}) &= (1 + u)(1 + v)(1 - w) \\
\beta_4(\mathbf{b}) &= (1 - u)(1 - v)(1 + w) \\
\beta_5(\mathbf{b}) &= (1 + u)(1 - v)(1 + w) \\
\beta_6(\mathbf{b}) &= (1 - u)(1 + v)(1 + w) \\
\beta_7(\mathbf{b}) &= (1 + u)(1 + v)(1 + w)
\end{align*}
\]

The matrix of partials is a real eyesore this time around,

\[
\mathbf{H} = \begin{bmatrix}
\frac{\partial \beta_0}{\partial u} & \frac{\partial \beta_0}{\partial v} & \frac{\partial \beta_0}{\partial w} \\
\frac{\partial \beta_1}{\partial u} & \frac{\partial \beta_1}{\partial v} & \frac{\partial \beta_1}{\partial w} \\
\frac{\partial \beta_2}{\partial u} & \frac{\partial \beta_2}{\partial v} & \frac{\partial \beta_2}{\partial w} \\
\frac{\partial \beta_3}{\partial u} & \frac{\partial \beta_3}{\partial v} & \frac{\partial \beta_3}{\partial w} \\
\frac{\partial \beta_4}{\partial u} & \frac{\partial \beta_4}{\partial v} & \frac{\partial \beta_4}{\partial w} \\
\frac{\partial \beta_5}{\partial u} & \frac{\partial \beta_5}{\partial v} & \frac{\partial \beta_5}{\partial w} \\
\frac{\partial \beta_6}{\partial u} & \frac{\partial \beta_6}{\partial v} & \frac{\partial \beta_6}{\partial w} \\
\frac{\partial \beta_7}{\partial u} & \frac{\partial \beta_7}{\partial v} & \frac{\partial \beta_7}{\partial w} \\
\end{bmatrix}
\tag{D.42}
\]

and unlike before, this time it does not work out to a matrix of constants:

\[
\mathbf{H} = \begin{bmatrix}
-(1 - v)(1 - w) & -(1 - u)(1 - w) & -(1 - u)(1 - v) \\
+(1 - v)(1 - w) & +(1 - u)(1 - w) & -(1 - u)(1 + v) \\
-(1 + v)(1 - w) & -(1 - u)(1 + w) & +(1 - u)(1 - v) \\
+(1 + v)(1 - w) & +(1 + u)(1 - w) & -(1 + u)(1 + v) \\
-(1 - v)(1 + w) & -(1 - u)(1 + w) & +(1 - u)(1 - v) \\
+(1 - v)(1 + w) & +(1 - u)(1 + w) & +(1 - u)(1 - v) \\
-(1 + v)(1 + w) & +(1 + u)(1 + w) & +(1 + u)(1 - v) \\
+(1 + v)(1 + w) & +(1 + u)(1 + w) & +(1 + u)(1 + v)
\end{bmatrix}
\tag{D.43}
\]

Typing this equation into C++ is an irritating and error-prone process, so I have listed 
my implementation of in Fig. D.3. Let me know if you find any typos.

If we want to compute \( \mathbf{F} = \mathbf{XH(\bar{XH})^{-1}} \), we first build out the \( \mathbf{X} \) and \( \bar{\mathbf{X}} \) matrices, which
now contains all eight vertices of the cube:

\[
\mathbf{X} = \begin{bmatrix} x_0 & x_1 & x_2 & x_3 & x_4 & x_5 & x_6 & x_7 \end{bmatrix} \in \mathbb{R}^{3 \times 8} \tag{D.44}
\]

and

\[
\bar{\mathbf{X}} = \begin{bmatrix} \bar{x}_0 & \bar{x}_1 & \bar{x}_2 & \bar{x}_3 & \bar{x}_4 & \bar{x}_5 & \bar{x}_6 & \bar{x}_7 \end{bmatrix} \in \mathbb{R}^{3 \times 8}. \tag{D.45}
\]

Since \( \mathbf{H} \) is not constant, we have to select \( b \), the location of the quadrature point. In the case of triangles and tetrahedra (with linear basis functions) that we saw before, \( \mathbf{F} \) was uniquely defined over the entire primitive. With a hexahedron, \( \mathbf{F} \) is different depending on where you are in the cube, so you need to specify the exact location you want to sample \( \mathbf{F} \) at.

Just like with tetrahedra, you can pre-cache \( \mathbf{D}^{-1}_m \) at startup, but in this case, you can actually do a little better. In the entire expression \( \mathbf{F} = \mathbf{XH}(\bar{\mathbf{X}}\mathbf{H})^{-1} \), the only matrix that will vary at runtime is \( \mathbf{X} \), so why not go ahead and cache \( \mathbf{H}(\bar{\mathbf{X}}\mathbf{H})^{-1} = \mathbf{H}\mathbf{D}^{-1}_m \)? It will save you a matrix multiply every time. An example of what I mean is in Fig. D.4.
D. How to Compute $\mathbf{F}$, the Deformation Gradient

```cpp
// Shape function derivative matrix
static Matrix8x3 ComputeH(const Vector3& b) {
    const Scalar b0Minus = (1.0 - b[0]);
    const Scalar b0Plus = (1.0 + b[0]);
    const Scalar b1Minus = (1.0 - b[1]);
    const Scalar b1Plus = (1.0 + b[1]);
    const Scalar b2Minus = (1.0 - b[2]);
    const Scalar b2Plus = (1.0 + b[2]);

    Matrix8x3 H;

    H(0,0) = -b1Minus * b2Minus;
    H(0,1) = -b0Minus * b2Minus;
    H(0,2) = -b0Minus * b1Minus;

    H(1,0) = b1Minus * b2Minus;
    H(1,1) = -b0Plus * b2Minus;
    H(1,2) = -b0Plus * b1Minus;

    H(2,0) = -b1Plus * b2Minus;
    H(2,1) = b0Minus * b2Minus;
    H(2,2) = -b0Minus * b1Plus;

    H(3,0) = b1Plus * b2Minus;
    H(3,1) = b0Plus * b2Minus;
    H(3,2) = -b0Plus * b1Plus;

    H(4,0) = -b1Minus * b2Plus;
    H(4,1) = -b0Minus * b2Plus;
    H(4,2) = b0Minus * b1Minus;

    H(5,0) = b1Minus * b2Plus;
    H(5,1) = -b0Plus * b2Plus;
    H(5,2) = b0Plus * b1Minus;

    H(6,0) = -b1Plus * b2Plus;
    H(6,1) = b0Minus * b2Plus;
    H(6,2) = b0Minus * b1Plus;

    H(7,0) = b1Plus * b2Plus;
    H(7,1) = b0Plus * b2Plus;
    H(7,2) = b0Plus * b1Plus;

    return H / 8.0;
}
```

Figure D.3.: Shape derivative matrix for Eqn. D.43.
D. How to Compute $F$, the Deformation Gradient

```cpp
// quadrature point locations along the inside
// of a canonical cube
static const Scalar invSqrt3 = 1.0 / std::sqrt(3.0);
static const std::array<Vector3, 8> restGaussPoints =
{
  invSqrt3 * Vector3(-1, -1, -1),
  invSqrt3 * Vector3( 1, -1, -1),
  invSqrt3 * Vector3(-1, 1, -1),
  invSqrt3 * Vector3( 1, 1, -1),
  invSqrt3 * Vector3(-1, -1, 1),
  invSqrt3 * Vector3( 1, -1, 1),
  invSqrt3 * Vector3(-1, 1, 1),
  invSqrt3 * Vector3( 1, 1, 1)
};

// cache DmInv at startup
std::array<Matrix8x3, 8> ComputeHDmInv(const Matrix3x8& Xbar)
{
  std::array<Matrix8x3, 8> HDmInv;
  for (int x = 0; x < 8; x++)
  {
    // compute Dm
    const Matrix8x3 H = ComputeH(restGaussPoints[x]);
    const Matrix3 Dm = Xbar * H;
    const Matrix3 DmInverse = Dm.inverse();
    // cache it left-multiplied by H
    HDmInv[x] = H * DmInverse;
  }
  return HDmInv;
}
```

Figure D.4.: Caching $\mathbf{H}\mathbf{D}_m^{-1}$ for hexahedra.
Appendix E

All the Details of $\frac{\partial F}{\partial x}$

Following on from §3.4.1 we want to compute

$$\frac{\partial F}{\partial x} = \frac{\partial D_s}{\partial x} D_m^{-1}. \quad (E.1)$$

This is a 3rd-order tensor containing twelve matrices. First, we will show the form of each of the twelve scalar derivatives $\frac{\partial D_s}{\partial x}$, and then we'll show each combined $\frac{\partial D_s}{\partial x_i} D_m^{-1}$.

Again, the entries of $D_m$ and $D_s$ are:

$$D_s = \begin{bmatrix} x_1 - x_0 & x_2 - x_0 & x_3 - x_0 \end{bmatrix}, \quad D_m = \begin{bmatrix} \bar{x}_1 - \bar{x}_0 & \bar{x}_2 - \bar{x}_0 & \bar{x}_3 - \bar{x}_0 \end{bmatrix}, \quad (E.2)$$

and we want the twelve resulting matrices from taking the derivative with respect to $x = \begin{bmatrix} x_0 \\ x_1 \\ x_2 \\ x_3 \end{bmatrix}$. These then get arranged like cheese slices on a cutting board:

$$\frac{\partial D_s}{\partial x} = \begin{bmatrix} \frac{\partial D_s}{\partial x_0} \\ \frac{\partial D_s}{\partial x_1} \\ \vdots \\ \frac{\partial D_s}{\partial x_{11}} \end{bmatrix}, \quad (E.3)$$
Finally, here are the all the actual entries of $\partial D_s/\partial x$:

$$
\begin{align*}
\frac{\partial D_s}{\partial x_0} &= \begin{bmatrix} -1 & -1 & -1 \\ 0 & 0 & 0 \\ 0 & 0 & 0 \end{bmatrix}, &
\frac{\partial D_s}{\partial x_1} &= \begin{bmatrix} 0 & 0 & 0 \\ -1 & -1 & -1 \\ 0 & 0 & 0 \end{bmatrix}, &
\frac{\partial D_s}{\partial x_2} &= \begin{bmatrix} 0 & 0 & 0 \\ -1 & -1 & -1 \\ 0 & 0 & 0 \end{bmatrix} \\
\frac{\partial D_s}{\partial x_3} &= \begin{bmatrix} 1 & 0 & 0 \\ 0 & 0 & 0 \\ 0 & 0 & 0 \end{bmatrix}, &
\frac{\partial D_s}{\partial x_4} &= \begin{bmatrix} 0 & 0 & 0 \\ 1 & 0 & 0 \\ 0 & 0 & 0 \end{bmatrix}, &
\frac{\partial D_s}{\partial x_5} &= \begin{bmatrix} 0 & 0 & 0 \\ 0 & 0 & 0 \\ 1 & 0 & 0 \end{bmatrix} \\
\frac{\partial D_s}{\partial x_6} &= \begin{bmatrix} 0 & 1 & 0 \\ 0 & 0 & 0 \\ 0 & 0 & 0 \end{bmatrix}, &
\frac{\partial D_s}{\partial x_7} &= \begin{bmatrix} 0 & 0 & 0 \\ 0 & 1 & 0 \\ 0 & 0 & 0 \end{bmatrix}, &
\frac{\partial D_s}{\partial x_8} &= \begin{bmatrix} 0 & 0 & 0 \\ 0 & 0 & 0 \\ 0 & 1 & 0 \end{bmatrix} \\
\frac{\partial D_s}{\partial x_9} &= \begin{bmatrix} 0 & 0 & 1 \\ 0 & 0 & 0 \\ 0 & 0 & 0 \end{bmatrix}, &
\frac{\partial D_s}{\partial x_{10}} &= \begin{bmatrix} 0 & 0 & 0 \\ 0 & 0 & 1 \\ 0 & 0 & 0 \end{bmatrix}, &
\frac{\partial D_s}{\partial x_{11}} &= \begin{bmatrix} 0 & 0 & 0 \\ 0 & 0 & 0 \\ 0 & 0 & 1 \end{bmatrix}.
\end{align*}
$$

All ones and zeros! There are a lot of them, but they are indeed quite simple. To write down the full $\partial F/\partial x$ expression, we now need to label the rows and columns of $D^{-1}_m$, which we do thusly:

$$
D^{-1}_m = \begin{bmatrix} \frac{r_0}{r_2} \\ \frac{r_1}{r_2} \end{bmatrix} = \begin{bmatrix} c_0 & c_1 & c_2 \end{bmatrix}.
$$

Now we can list the results of multiplying through each $\frac{\partial F}{\partial x_i} = \frac{\partial D_s}{\partial x_i} D^{-1}_m$:

$$
\frac{\partial F}{\partial x_0} = \begin{bmatrix} -s_0 & -s_1 & -s_2 \\ 0 & 0 & 0 \\ 0 & 0 & 0 \end{bmatrix}, &
\frac{\partial F}{\partial x_1} = \begin{bmatrix} 0 & 0 & 0 \\ -s_0 & -s_1 & -s_2 \\ 0 & 0 & 0 \end{bmatrix}, &
\frac{\partial F}{\partial x_2} = \begin{bmatrix} 0 & 0 & 0 \\ -s_0 & -s_1 & -s_2 \\ 0 & 0 & 0 \end{bmatrix} \\
\frac{\partial F}{\partial x_3} = \begin{bmatrix} r_0 \\ 0 & 0 & 0 \\ 0 & 0 & 0 \end{bmatrix}, &
\frac{\partial F}{\partial x_4} = \begin{bmatrix} r_1 \\ 0 & 0 & 0 \\ 0 & 0 & 0 \end{bmatrix}, &
\frac{\partial F}{\partial x_5} = \begin{bmatrix} r_2 \\ 0 & 0 & 0 \\ 0 & 0 & 0 \end{bmatrix} \\
\frac{\partial F}{\partial x_6} = \begin{bmatrix} r_0 \\ 0 & 0 & 0 \\ 0 & 0 & 0 \end{bmatrix}, &
\frac{\partial F}{\partial x_7} = \begin{bmatrix} r_1 \\ 0 & 0 & 0 \\ 0 & 0 & 0 \end{bmatrix}, &
\frac{\partial F}{\partial x_8} = \begin{bmatrix} r_2 \\ 0 & 0 & 0 \\ 0 & 0 & 0 \end{bmatrix} \\
\frac{\partial F}{\partial x_9} = \begin{bmatrix} r_0 \\ 0 & 0 & 0 \\ 0 & 0 & 0 \end{bmatrix}, &
\frac{\partial F}{\partial x_{10}} = \begin{bmatrix} 0 & 0 & 0 \\ 0 & 0 & 0 \\ 0 & 0 & 0 \end{bmatrix}, &
\frac{\partial F}{\partial x_{11}} = \begin{bmatrix} 0 & 0 & 0 \\ 0 & 0 & 0 \\ 0 & 0 & 0 \end{bmatrix}.
$$

In the first row, $s_i$ denotes the sum of the entries in $c_i$. Again, there’s a lot of stuff here, but each individual piece is not very complicated. The final matrix entries are all just rows and columns from $D^{-1}_m$.
As a big birthday present to you (and Future Me), the code for the flattened version of this tensor, $\text{vec}\left( \frac{\partial F}{\partial x} \right)$ is given in Fig. E.1. Finally, it is important to point out that this code only applies to tetrahedra. Hexahedra and other polyhedra are another can of worms.
E. All the Details of $\frac{\partial F}{\partial x}$

```cpp
static Matrix9x12 ComputePFPx(const Matrix3& DmInv)
{
    const Scalar m = DmInv(0, 0);
    const Scalar n = DmInv(0, 1);
    const Scalar o = DmInv(0, 2);
    const Scalar p = DmInv(1, 0);
    const Scalar q = DmInv(1, 1);
    const Scalar r = DmInv(1, 2);
    const Scalar s = DmInv(2, 0);
    const Scalar t = DmInv(2, 1);
    const Scalar u = DmInv(2, 2);

    const Scalar t1 = - m - p - s;
    const Scalar t2 = - n - q - t;
    const Scalar t3 = - o - r - u;

    Matrix9x12 PFPx;
    PFPx.setZero();
    PFPx(0, 0) = t1;
    PFPx(0, 3) = m;
    PFPx(0, 6) = p;
    PFPx(0, 9) = s;
    PFPx(1, 1) = t1;
    PFPx(1, 4) = m;
    PFPx(1, 7) = p;
    PFPx(1, 10) = s;
    PFPx(1, 10) = t1;
    PFPx(2, 2) = t1;
    PFPx(2, 5) = m;
    PFPx(2, 8) = p;
    PFPx(2, 11) = s;
    PFPx(3, 0) = t2;
    PFPx(3, 3) = n;
    PFPx(3, 6) = q;
    PFPx(3, 9) = t;
    PFPx(4, 1) = t2;
    PFPx(4, 4) = n;
    PFPx(4, 7) = q;
    PFPx(4, 10) = t;
    PFPx(5, 2) = t2;
    PFPx(5, 5) = n;
    PFPx(5, 8) = q;
    PFPx(5, 11) = t;
    PFPx(6, 0) = t3;
    PFPx(6, 3) = o;
    PFPx(6, 6) = r;
    PFPx(6, 9) = u;
    PFPx(7, 1) = t3;
    PFPx(7, 4) = o;
    PFPx(7, 7) = r;
    PFPx(7, 10) = u;
    PFPx(8, 2) = t3;
    PFPx(8, 5) = o;
    PFPx(8, 8) = r;
    PFPx(8, 11) = u;

    return PFPx;
}
```

Figure E.1.: C++ code to compute the flattened version of $\frac{\partial F}{\partial x}$. 
Appendix F

Rotation-Variant SVD and Polar Decomposition

The problem described in §2.3.4.4 is that the traditional polar decomposition, which we will call $F = \hat{R}\hat{S}$, only requires $\hat{R}$ to be unitary. A troublesome reflection can then lurk in $\hat{R}$ that prevents it from being a pure rotation. We will now describe a simple way to compute the rotation variant of the polar decomposition. Along the way, we will also show how to do the same for the SVD.

A straightforward way to compute the polar decomposition of $F$ is to first compute its SVD,

$$F = \hat{U}\hat{\Sigma}\hat{V}^T,$$  \hspace{1cm} (F.1)

and then recombine its components into $\hat{R}$ and $\hat{S}$:

$$\hat{R} = \hat{U}\hat{V}^T$$ \hspace{1cm} (F.2)

$$\hat{S} = \hat{V}\hat{\Sigma}\hat{V}^T$$ \hspace{1cm} (F.3)

Since $\hat{U}$ and $\hat{V}$ are also unitary, these representations are all trivially equivalent:

$$\hat{R}\hat{S} = \hat{U}\hat{\Sigma}\hat{V}^T\hat{V}\hat{\Sigma}\hat{V}^T$$ \hspace{1cm} (F.4)

$$= \hat{U}\hat{\Sigma}\hat{V}^T\hat{\Sigma}\hat{V}^T$$  \hspace{.5cm} (Used unitary property)

$$= \hat{U}\hat{\Sigma}\hat{V}^T = F.$$ \hspace{1cm} (Definition of SVD)

What we want is to generate purely rotational versions of $\hat{U}$ and $\hat{V}$, i.e. $U$ and $V$. Then we can combine them into another pure rotation, $R = UV^T$.

One easy way to test whether $\hat{R}$ is secretly harboring a reflection is to look at the determinant of $\hat{R}$, i.e. $\det\hat{R}$. The determinant is the product of the singular values of a matrix, so if $\hat{R}$ was a pure rotation, all its singular values would be $\Sigma = \begin{bmatrix} 1 & 0 & 0 \\ 0 & 1 & 0 \\ 0 & 0 & 1 \end{bmatrix}$.
and \( \det \hat{R} = 1 \). However, if it contains a reflection, one singular value must be \( -1 \),
\[
\Sigma = \begin{bmatrix}
1 & 0 & 0 \\
0 & 1 & 0 \\
0 & 0 & -1
\end{bmatrix}, \quad \text{and} \quad \det \hat{R} = -1.
\]

There’s no such thing as a double reflection, only a single reflection. A double-reflection is the same as rotating by \( \pi \), which is a valid rotation. In that case, the rotation would have been loaded into \( \hat{U} \) or \( \hat{V} \).

With this in hand, we can now build a reflection matrix:
\[
L = \begin{bmatrix}
1 & 0 & 0 \\
0 & 1 & 0 \\
0 & 0 & \det(\hat{U}\hat{V}^T)
\end{bmatrix}.
\]  

(F.5)

Computing the rotation variant of \( \Sigma \) is now straightforward:
\[
\Sigma = \hat{\Sigma}L.
\]  

(F.6)

If there were no reflections lurking anywhere, then \( L = I \), so no harm done. However, we have to be a little careful when computing the new \( U \) and \( V \). We need to check whether the reflection is lurking in either \( \hat{U} \) or \( \hat{V} \), and only apply \( L \) to that one:
\[
U = \begin{cases} 
\hat{U}L & \text{if} \ \det \hat{U} < 0 \ \text{and} \ \det \hat{V} > 0 \\
\hat{U} & \text{otherwise}
\end{cases}
\]  

(F.7)

\[
V = \begin{cases} 
\hat{V}L & \text{if} \ \det \hat{U} > 0 \ \text{and} \ \det \hat{V} < 0 \\
\hat{V} & \text{otherwise}.
\end{cases}
\]  

(F.8)

With the rotation-variant SVD in hand, computing the rotation-variant polar decomposition is easy:
\[
R = UV^T
\]  

(F.9)

\[
S = V\Sigma V^T.
\]  

(F.10)

Complete Matlab/Octave code for the 3D case is given in Figs. F.1 and F.2

Historically, the need for the rotation-variant SVD and polar decomposition has been observed by a variety of authors (Twigg and Kačić-Alesić (2010); Higham (2008); Irving et al. (2004)). The version we describe here with the slick determinant/reflection matrix trick is from Sorkine-Hornung and Rabinovich (2017).
function \[U \Sigma V\] = svd_rv(F)
\[
[U \Sigma V] = \text{svd}(F);
\]

% reflection matrix
L = \text{eye}(3,3);
L(3,3) = \text{det}(U \ast V');

% see where to pull the reflection out of
detU = \text{det}(U);
detV = \text{det}(V);
if (detU < 0 && detV > 0)
    U = U \ast L;
elseif (detU > 0 && detV < 0)
    V = V \ast L;
end

% push the reflection to the diagonal
Sigma = Sigma \ast L;
end

Figure F.1.: Matlab code to compute the rotation-variant SVD.

function [R S] = polar_decomposition_rv(F)
\[
[U \Sigma V] = \text{svd_rv}(F);
R = U \ast V';
S = V \ast \Sigma \ast V';
end

Figure F.2.: Matlab code to compute the rotation-variant polar decomposition.
Bibliography


Bibliography


