Analyzing valve interstitial cell mechanics and geometry with spatial statistics

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ABSTRACT

Understanding cell geometric and mechanical properties is crucial to understanding how cells sense and respond to their local environment. Moreover, changes to cell mechanical properties under varied micro-environmental conditions can both influence and indicate fundamental changes to cell behavior. Atomic Force Microscopy (AFM) is a well established, powerful tool to capture geometric and mechanical properties of cells. We have previously demonstrated substantial functional and behavioral differences between aortic and pulmonary valve interstitial cells (VIC) using AFM and subsequent models of VIC mechanical response. In the present work, we extend these studies by demonstrating that to best interpret the spatially distributed AFM data, the use of spatial statistics is required. Spatial statistics includes formal techniques to analyze spatially distributed data, and has been used successfully in the analysis of geographic data. Thus, spatially mapped AFM studies of cell geometry and mechanics are analogous to more traditional forms of geospatial data. We are able to compare the spatial autocorrelation of stiffness in aortic and pulmonary valve interstitial cells, and more accurately capture cell geometry from height recordings. Specifically, we showed that pulmonary valve interstitial cells display higher levels of spatial autocorrelation of stiffness than aortic valve interstitial cells. This suggests that aortic VICs form different stress fiber structures than their pulmonary counterparts, in addition to being more highly expressed and stiffer on average. Thus, the addition of spatial statistics can contribute to our fundamental understanding of the differences between cell types. Moving forward, we anticipate that this work will be meaningful to enhance direct analysis of experimental data and for constructing high fidelity computational of VICs and other cell models.

1. Introduction

Cell mechanical and geometric properties play an important role in how cells sense and interact with their surrounding environment (Luo et al., 2016). Quantifying these properties is crucial for understanding the specific roles of mechanical forces in biological tissues (Haase and Pelling, 2015; Lejeune and Linder, 2017; Rigato et al., 2015), including quantifying variation between different cell types (Cross et al., 2008). For example, in the mechanically demanding heart valve tissue micro-environment, valve interstitial cells (VICs) respond to mechanical stimuli due to altered hemodynamic conditions and subsequently contribute to tissue remodeling (Ayoub et al., 2011, 2017). Therefore, understanding how VICs feel their surroundings is crucial for robustly predicting how heart valve surgical interventions can lead to tissue remodeling (Rego and Sacks, 2017; Sacks et al., 2017), and measuring mechanobiological variations between VICs derived from different sources provides insight into characteristic physiological and pathological changes (Merryman et al., 2006, 2007).

Atomic force microscopy (AFM) is a popular and well established tool to quantify both cell topography and intrinsic cell mechanical properties on two-dimensional substrates (Gavara, 2017). To measure mechanical properties, an AFM probe is used for micro-indentation. The cantilever tip indents the cell at discrete grid points, and the force response curve for each indentation is used to approximate the value of local cell stiffness, reported as apparent elastic modulus (Merryman et al., 2007; Mathur et al., 2001). To measure cell topography, the scanning probe is dragged over the surface of the cell and height is recorded at densely spaced points along the path of the probe (Merryman et al., 2007; Radmacher et al., 1992). Typically, data from these experiments

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is presented visually, as cell height and stiffness maps, and quantitatively, through the mean and standard deviation of these measured values (Yew et al., 2017). Furthermore, cell indentation by AFM has been modeled extensively (Nguyen et al., 2016). Our group has used AFM in the study of VICs, both directly (Merryman et al., 2007), and in simulations (Sakamoto et al., 2016, 2017).

As AFM data is recorded at multiple points in space, it is natural to turn to the field of spatial statistics to analyze, and ultimately model, the resulting spatially distributed data. Spatial statistics is specifically intended to handle data where the assumption of independence among observations is violated (Ripley, 2005). For example, spatial statistics is used to characterize spatial autocorrelation, the co-variation of properties within space, and detect patterns and anomalies in spatially distributed data (Goodchild, 1986). The field of spatial statistics also covers methods for more sophisticated interpolation techniques to make enhanced predictions at unmeasured locations (Stein, 1999). The most common application of spatial statistics is in analyzing geographic data and creating geographic models, where specialized statistical analysis software packages are used in both research and industry (Steiniger and Hunter, 2012). Though spatial statistics is a natural toolset for handling spatially distributed data, such as that generated by AFM studies of biological materials, to the authors’ knowledge, this application has yet to be explored in detail.

In the present work, we show how the toolset of spatial statistics can be meaningfully applied to the problem of analyzing the spatially mapped data from AFM cell studies. Specifically, we introduce three example methods from the spatial statistics literature: single variable measurements of spatial autocorrelation, empirical semivariograms, and Gaussian process regression. Then, we show the results of applying these methods to AFM studies of aortic valve interstitial cells (AVICs) and pulmonary valve interstitial cells (PVICs). Critically, these methods quantitatively reveal different spatial autocorrelation behaviors in AVIC and PVIC stiffness. These results suggest that differences between AVIC and PVIC stiffness are a result of both intrinsic stress-fiber stiffness, and also how sub-cellular components are spatially distributed within the cell. The results of this study and the methods presented can also be extended to the general AFM studies of cell mechanics.

2. Methods

In this section, we briefly introduce three non-standard methods to spatially interpret AFM data. The first two methods that we introduce, computing Moran’s I and plotting an empirical semivariogram, are useful for understanding spatial autocorrelation in experimentally recorded data. Essentially, they show how similar the measured values are at spatially close locations. The third method that we introduce, Gaussian process regression, is a pragmatic strategy for interpolating spatial data. If statistically significant spatial autocorrelation is detection with the first two methods, then Gaussian process regression can be used to predict unseen values \( y^* \) at unsampled locations \( x^* \) in order to construct robust models of cells with heterogeneous material properties.

2.1. Experimental data source

The AFM data for this work was taken from (Merryman et al., 2007), in which information on the experimental methods has been extensively detailed. Briefly, porcine AVICs and PVICs were isolated from the same animal. The AV and PV leaflets were dissected, leaflet surfaces were scraped to remove endothelial cells, and the leaflets were digested and strained to isolate the VICs. Both populations were plated separately in 250 ml culture flasks in complete media. A monolayer of seeded VICs from each valve type were structurally mapped by AFM. VIC stiffness was measured in the so-called tapping mode and the AFM probe was modeled as a conical tip with a probe opening angle of 35°. The apparent stiffness of the cell was calculated based on the Hertzian model. For each VIC type, multiple indentations at different points were made over the surface of the cell and each force-indentation depth curve was fit to determine the apparent stiffness \( E \). In summary, the resulting AFM data contains scalar values of interest \( y \) (height, stiffness) measured at discrete positions \( x_k = (x_k^1, x_k^2) \). In the examples presented in Section 3, the scalar values \( y \) are apparent Young’s modulus \( E \) (Fig. 1a) and cell height \( h \) (Fig. 1b). It should be noted that these techniques are sufficiently general such that they can be directly applied to other experimentally measured scalar quantities of interest.

2.2. Moran’s I

One standard single variable measurement of spatial autocorrelation in a data set is Global Moran’s I (Mitchel, 2005; Moran, 1950). Given a single cell with stiffness sampled by AFM indentation at \( n \) locations, \( I \) is defined as

\[
I = \frac{n}{S_0} \sum_{i=1}^{n} \sum_{j=1}^{n} w_{ij} (y_i - \bar{y})(y_j - \bar{y}) \tag{1}
\]

where \( y_i \) is the scalar variable of interest, \( \bar{y} \) is the mean of all \( y_i \), \( w_{ij} \) is defined as a matrix of weights in which immediate neighbors have value 1 with \( w_{kk} = 0 \), and \( S_0 \) is defined as

\[
S_0 = \sum_{i=1}^{n} \sum_{j=1}^{n} w_{ij} \tag{2}
\]

If there is no spatial autocorrelation, then \( I = 0 \). Negative spatial correlation corresponds to \(-1 < I < 0 \) and positive spatial autocorrelation corresponds to \( 0 < I < 1 \). To interpret \( I \), we test for statistical significance by comparing the measured value of \( I \) to the null hypothesis that the values in the data set are not spatially correlated (Mitchel, 2005; Moran, 1950). To do this, we compute the expected value \( E[I] \) and variance \( V[I] \) for the uncorrelated case

\[
E[I] = -\frac{1}{n - 1} \quad V[I] = E[I^2] - E[I]^2 \tag{3}
\]

where \( E[I^2] \) is computed as

a)

b)

Fig. 1. (a) an AFM probe is used to measure cell stiffness by indenting the cell at discrete grid points; (b) an AFM probe is used to measure cell height by tracing over the surface of the cell.
\[
J^2 = \frac{n(n^2 - 3n + 3)S_1 - nS_2 + 3S_0^3}{(n - 1)(n - 2)(n - 3)S_0^3}
\]

where the number of points in a bin is \(N(h \pm \delta)\) and \(y_i\) is the scalar quantity measured at each point. Through this equation, \(\tilde{\gamma}\) captures the average difference between sample pairs at a fixed separation distance. When data is spatially correlated, the semivariogram will show that values of \(y\) are more similar when the distance between them is smaller (Fig. 2). When spatial autocorrelation is present, \(\frac{1}{2}||y_i - y_j||^2\) approaches zero as \(||x_i - x_j||\) approaches zero.

For additional analysis, a semivariogram model can be fitted to an empirical semivariogram when there is sufficient data present. In a semivariogram model, there are three key parameters: the range, the sill, and the nugget. The range indicates the distance over which data points are spatially autocorrelated, the sill indicates the value on the \(y\) axis of the semivariogram at the \(x\) value range, and the nugget (often set to 0) captures measurement error and variations over distances smaller than the minimum sampling distance. The interested reader is referred to the literature for more information on semivariogram models (Cressie, 1993; Mälicke and Schneider, 2018).

### 2.4. Gaussian process regression

The two methods introduced in Sections 2.2 and 2.3 are useful for determining if spatial correlation is present in AFM data. When the data is spatially correlated, it is appropriate to develop a model that can be used to predict the measured values \(y\) between the locations where the experimental data is recorded. Gaussian process regression (GPR), often referred to as kriging, is a non-parametric kernel-based supervised learning method that is commonly used to predict spatially correlated measurements (Rasmussen and Williams, 2006). With the AFM data, this means using position \(x_i\) to predict location specific scalar value \(y_i\). This allows for the prediction of \(y_i\) at locations where \(x_i\) has not been measured.

Here we describe an established pragmatic strategy to conduct Gaussian process regression. First, we define a training set \(\mathcal{D}\) of \(n\) observed data points, \(\mathcal{D} = \{x_i, y_i\}_{i=1}^n\) where \(x_i\) denotes an input vector, in this case the position of a sample, \(y_i\) denotes a scalar output, in this case either measured height or stiffness. We define an unseen or “test” data set \(\mathcal{D}^*\) of \(n^*\) points with input \(x^*\) and output \(y^*\). A Gaussian process is defined via a mean function (in this case we scale our data and assume zero mean) and positive definite covariance matrices that relates values with features \(x\) that are close together. We define \(K\) as the \(n \times n\) matrix of covariances evaluated at all pairs of training points, define \(K^*\) as the \(n^* \times n^*\) matrix of covariances evaluated at all pairs of training and test points, and define \(K_{\cdot \cdot}\) as the \(n \times n^*\) matrix of covariances evaluated at all pairs of test points. We assume that observations \(y\) are potentially noisy, with \(y = f(x) + \epsilon\) where \(\epsilon\) is additive independent.

![Fig. 2. A semivariogram contains information about correlations in spatially distributed data. Left: scalar value \(y_i\) is measured at each location \(x_i\) marked by an orange cross; Right: the value \(\frac{1}{2}||y_i - y_j||^2\) is plotted with respect to the distance between \(x_i\) and \(x_j\) for each location pair \(x_i\) and \(x_j\). The example illustrates spatially correlated data because the value \(\frac{1}{2}||y_i - y_j||^2\) approaches 0 as \(||x_i - x_j||\) approaches 0.](image)
identically distributed Gaussian noise. Then, we write the joint distribution of the test and training locations under the prior as

$$\begin{bmatrix} y \\ f_\cdot \end{bmatrix} \sim \mathcal{N} \left( \begin{bmatrix} 0 \\ K + \sigma^2_n \mathbf{I} \\ K^T_n \\ K_n \end{bmatrix} \right)$$

(11)

where $\mathcal{N}$ is a multivariate normal distribution. We use this equation to derive the predicted value of $f_\cdot$ in terms of the values which are known as

$$f_\cdot | x, y, x_\cdot \sim \mathcal{N} \left( f_\cdot, \text{cov}(f_\cdot) \right).$$

(12)

If we consider a single test point $x_\cdot$, where $K$ is the vector of covariances between the test point and the $n$ training points, the predicted mean and variance of $f_\cdot$ are

$$\tilde{f}_\cdot = K^T_n (K + \sigma^2_n \mathbf{I})^{-1} y$$

(13)

and

$$\text{var}(f_\cdot) = K(x_\cdot, x_\cdot) - K^T_n (K + \sigma^2_n \mathbf{I})^{-1} K_n.$$  

(14)

Essentially, the training data set is used to construct a Gaussian field which is then used to make predictions on the test data set and subsequent unmeasured locations (Fig. 3).

Given this structure, the GPR model is fully specified by choosing the kernel function and the optimal kernel hyperparameters. A pragmatic way to choose the kernel and hyperparameters is to separate the available data into test and training data sets and select the GPR model that performs best at predicting the test data given the training data. The model that performs best will have the lowest test error, where test error is a function of the known values $y$ and the predictions $f_\cdot$. Additional information on separating the kernel function and the optimal kernel hyperparameters is to separate the available data into test and training data sets and select the model that performs best at predicting the test data given the training data. The model that performs best will have the lowest test error, where test error is a function of the known values $y$ and the predictions $f_\cdot$. Additional information on separating the kernel function and the optimal kernel hyperparameters is given in Appendix A. Ultimately, GPR will likely lead to a better approximation of cell properties between the measured data points than strategies such as linear interpolation, illustrated in Appendix B.

3. Results

As introduced in Section 1, AFM is commonly used to make spatially distributed measurements of cell stiffness and cell height. Here, we re-visit an experimental analysis of AVICs and PVICs where it was previously shown using standard methods that the two cell types have different average mechanical properties (Merryman et al., 2007). From our analysis, we obtain novel insights into the characteristics of AVICs and PVICs. First, we report global Moran’s $I$ and associated z-score $z_i$ for measurements of scalar stiffness represented by apparent elastic modulus $E$ for five aortic valve cells and six pulmonary valve cells (Table 1). None of the five AVICs showed statistically significant spatial correlation in stiffness ($p < 0.001$) at the length scale of sampling while six out of six PVICs showed statistically significant spatial autocorrelation. We then show a representative histogram and semivariogram for an AVIC with low spatial autocorrelation and a PVIC with high spatial autocorrelation (Fig. 4). Though the histograms look qualitatively similar, the semivariograms look qualitatively different. This is consistent with the empirical semivariogram plots for the other AVICs and PVICs (Fig. 5). Overall, AVICs show much higher variance for every lag distance, and the decrease in variance at small lag distances is much less pronounced than in the PVICs. Though the sample size is relatively small, the difference between the two cell types is consistent, both in the present study and in related studies.

Next, we show the results of using Gaussian process regression to interpolate both height data and stiffness data. First, we show the interpolated heights for a single cell measured by AFM (Fig. 6). Mean absolute test error under 0.2 $\mu$m, reported in Appendix A, was a good approximation for the error with this method. In addition, we show the interpolated stiffness values for a single cell measured by AFM (Fig. 7). Using leave one out cross validation, where all but one data points are included in the model at a time, we can get an understanding of model error on unseen data. For the cell shown, PV-4, which had high spatial autocorrelation, the mean absolute error was under 3 kPa. Additional analysis of this error is given in Appendix B. For the cells which show no significant spatial correlation, i.e. the AVIC cells, GPR is not a good strategy to model spatial variations in stiffness because stiffness will not be well predicted by position.

4. Discussion

4.1. Major findings

The methods described in this paper are simple yet powerful tools to interpret and model spatial heterogeneity in AFM data. With the simple methods described in Section 2, we are able to both quantify experimentally observed spatial autocorrelation and introduce a strategy towards better capturing spatially varying data. When we compute Moran’s $I$ for stiffness measurements in multiple AVICs and PVICs, following the method in Section 2.2, we are able to show that there are differences in spatial autocorrelation between the two cell types. Creating empirical semivariograms, following the method in Section 2.3, further exposes the fundamental distinction between the two cell types and illustrate the distance over which PVIC stiffness is correlated.

For the experimental measurements that display spatial autocorrelation, Gaussian process regression is a suitable method for interpolating data points. As emphasized in Section 2.4, care must be taken to select the appropriate kernel and hyperparameters to ensure accurate predictions. Additionally, the use of multiple kernel combinations and hyperparameter optimizations can further improve the accuracy of the predictions.
be taken when considering the separation of test and training data. The pragmatic approach put forward here for dealing with the directionally dependent spacing of AFM height data is appealing because of its simplicity. Furthermore, we show an example of GPR being used to interpolate cell stiffness in PVICs. We anticipate that GPR and similar enhanced interpolation strategies will be key to creating realistic computational cell models that account for spatial heterogeneity in cell geometry and properties. When cell properties are not spatially correlated, alternative methods must be introduced to create representative cell models (Ostoja-Starzewski, 2006).

The present findings reinforce and extend previous results that show differences in the mean cell stiffness, αSMA levels, stress fiber stiffness, and extracellular matrix remodeling capabilities between aortic and pulmonary VICs (Merryman et al., 2006, 2007, Sakamoto et al., 2016, 2017). In particular, previous studies have found via micropipette aspiration and AFM indentation that PVICs are less stiff than AVICs, AVICs express higher αSMA levels, intrinsically stronger stress fibers, and will typically remodel collagen gels at an accelerated rate. In this work, it was observed that PVICs exhibit much higher levels of spatial correlation in stiffness than their AVIC counterparts (Fig. 5). Essentially, AVICs have a more heterogeneous stiffness. This indicates that some sub-cellular stress-fiber formation activity is likely quite different between the two cell types. One possible explanation for this is that the stress fiber architecture of AVIC form more focal adhesion compared to the “smoother” spatial features of the PVICs. The underlying mechanism driving this difference could be connected to highly localized stress fiber formation in AVICs. We also note that another mechanisms driving this difference may be that AVICs and PVICs responds differently to cell culture conditions due to different degrees of deviation from their native chemical and mechanical environment. Future research will be needed to clarify these insights and compare the behavior of these isolated cells to cells in the native tissue.

4.2. Broader findings

Putting this in a broader perspective, distinguishing between cells types and detecting changes in cell state and behavior is critical for understanding tissue and organ scale function (Ayoub et al., 2017; Rego et al., 2018). Given that VICs are present in all heart valves, yet likely subject to dramatically different valve-specific mechanical loading conditions throughout their lifetime (Ayoub et al., 2018), quantitative methods for determining the difference between cell types are quite meaningful. With this enhanced...
strategy for quantifying experimental data, it is possible to both detect these differences and more robustly consider their implications in the broader context of multiscale modeling (Lee et al., 2017; Lejeune and Linder, 2018; Lejeune et al., 2019). Furthermore, this clear demonstration of spatial heterogeneity within individual cells shows that experiments with techniques such as micropipette aspiration may be missing critical information about the heterogeneous distribution of mechanical properties within the cell (Sakamoto et al., 2016).

Looking forward, capturing and modeling the spatial variation in cell properties is an important step towards understanding both cell mechanics and cell-substrate/cell-matrix interaction. Fundamentally, this methodology will lead to a better understanding of how sub-cellular components come together and contribute to the mechanical behavior of the whole cell. And, it will contribute to our understanding of how spatially heterogeneous cell behavior ultimately influences tissue function. In addition, information on spatial heterogeneity of cell stiffness will lead to enhanced computational models for calculating the forces that cells experience that, unlike many approaches, will not assume cell stiffness is homogeneous. Computational models will be particularly useful for comparing isolated cells to cells in the native tissue environment. Furthermore, additional understanding of the inherent heterogeneity in biological samples will interface well with methods for rapidly acquiring AFM measurements (Hartman and Andersson, 2017; HHuang and Anderssonuang and Andersson, 2012).

4.3. Conclusions

In conclusion, applying tools from spatial statistics to AFM data will provide substantial new insight and lead to a deeper understanding of the properties of biological cells. We were able to demonstrate that pulmonary valve interstitial cells display higher levels of spatial autocorrelation of stiffness than aortic valve interstitial cells. Thus, the addition of spatial statistics can contribute to our fundamental understanding of the differences between these cell types. The methods presented in this paper are also only a small subset of what is available in the spatial statistics literature. For example, Geary’s C and Getis-Ord Gi are other single variable measurements of spatial autocorrelation (Geary, 1954; Getis and Ord, 1992), and AFM indentation may be used to approximate indentation parameters beyond apparent elastic modulus (Putman et al., 1994). Furthermore, there are extensions of these techniques to understand correlations in both space and time (Sherman, 2011). Biomechanics researchers working with spatially heterogeneous AFM data should consider these approaches in their analysis.

Declaration of Competing Interest

The authors declared that there are no conflicts of interest.

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Appendix A. Separating test and training data for Gaussian Process Regression

A pragmatic strategy for constructing a GPR model is to separate data into a test set and a training set and evaluate model performance as the error on the test set. Much of this workflow is accomplished by a package such as sklearn-learn (Pedregosa et al., 2011). Here we note that it is important to consider what a meaningful split of test and training data is given the nature of the available data. More specifically, consider the AFM height measurements where the scanning probe is moved in a line along the
sample surface (Fig. 1). As the tip passes over the cell, multiple data points are recorded with a fixed y-value so the data is quite dense in the x-direction. The tip then passes over another line with a fixed y-value such that the distance between data points is much greater in the y-direction than in the x. If we first consider the AFM height data in one y plane, where the data available is quite dense and uniformly spaced, separating test and training data is straightforward. To demonstrate this, we show the decrease in mean absolute test error as the number of training samples is increased and the resulting closer match between the predicted height of the GPR model and the measured height (Fig. A.8).

Next, we consider the AFM height data in both the x and y planes. If we separate the test and training data without considering that the data is unevenly spaced, the reported test error is artificially low and the GPR parameters may be tuned such that the model is overfitting (Fig. A.9 left). Instead, to get a more realistic impression of test error and to select the GPR model parameters that will best predict unseen data, we must separate test and training error by grouping entire “strips” of data with identical y-coordinates corresponding to one pass of the AFM tip probe. When we do this, we see that the test error decreases with each additional “strip” of data and is notably higher than with the previous uninformed test and training data separation method (Fig. A.9 right). Furthermore, the best performing kernel may be different than the case where no special consideration is given to the separation of test and training data. The resulting GPR model can then be used to interpolate height data in the regions between each pass of the AFM tip (Fig. 6 right). When the data is evenly spaced in both dimensions, as is the case with the AFM stiffness data available, this additional consideration is not necessary.

### Appendix B. Comparing Gaussian Process Regression to linear interpolation

When the AFM data is quite densely spaced, for example in the x-direction of height measurements, the strategy for inferring values between measured data points is less important. However, when the spacing between data points is larger, for example in the y-direction of height measurements and the grid of stiffness measurements, critically evaluating how to infer these values matters. To illustrate this, we compare the mean absolute error (MAE) in height prediction computed with GPR and linear interpolation with respect to the number of training points (Fig. B.10 left) and the number of training strips (Fig. B.10 right). Though there is no apparent difference between the two methods for the incorrect test and training data separation, discussed in Appendix A, there

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![Fig. A.8.](image1)

**Fig. A.8.** Left: the mean absolute error (MAE) of the test predictions decreases as the number of training points increases; Center: GPR model fit with five sample points compared to raw data; Right: GPR model fit with 50 sample points compared to raw data.

![Fig. A.9.](image2)

**Fig. A.9.** Left: MAE of the test predictions approaches zero as the number of naively selected training points increases; Right: MAE of the test prediction decreases as the number of strips of training points increases.

![Fig. B.10.](image3)

**Fig. B.10.** Left: Comparison between MAE of the test predictions for GPR and linear interpolation as the number of naively selected training points increases; Right: Comparison between MAE of the test predictions for GPR and linear interpolation as the number of strips of training points increases.
is a notable difference for the correct test and training data separation. Interpolation with GPR results in a lower MAE, and the MAE for GPR appears to decrease with respect to the addition of more training data in a way that the linear interpolation strategy does not. Similar behavior is observed for interpolating stiffness, where MAE reliably decreases with the addition of more training data for GPR but not for linear interpolation (Fig. B.11 left). Of course, there are many other potential strategies besides GPR and linear interpolation (Friedman et al., 2001). For future endeavors in computational modeling of cells based on AFM data, we recommend critically evaluating multiple data-specific strategies for interpolating between measured data points. Noting the difference between GPR (Fig. 7) and linear interpolation (Fig. B.11 right) emphasizes this point.

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