

Three-dimensional image-based mechanical modeling for predicting the response of breast cancer to neoadjuvant therapy

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Available online 1 September 2016

Highlights

- Medical imaging data can be used to inform mathematical models of tumor growth.
- Models may assist therapeutic intervention for breast cancer therapeutic response.
- We present a volumetric extension of our data driven mathematical model predictions.
- Inclusion of additional volumetric patient-specific data enhances model predictions.

Abstract

The use of quantitative medical imaging data to initialize and constrain mechanistic mathematical models of tumor growth has demonstrated a compelling strategy for predicting therapeutic response. More specifically, we have demonstrated a data-driven framework for prediction of residual tumor burden following neoadjuvant therapy in breast cancer that uses a biophysical mathematical model combining reaction–diffusion growth/therapy dynamics and biomechanical effects driven by early time point imaging data. Whereas early work had been based on a limited dimensionality reduction (two-dimensional planar modeling analysis) to simplify the numerical implementation, in this work, we extend our framework to a fully volumetric, three-dimensional biophysical mathematical modeling approach in which parameter estimates are generated by an inverse problem based on the adjoint state method for numerical efficiency. In an *in silico* performance study, we show accurate parameter estimation with error less than 3% as compared to ground truth. We apply the approach to patient data from a patient with pathological complete response and a patient with residual tumor burden and demonstrate technical feasibility and predictive potential with direct comparisons between imaging data observation and model predictions of tumor cellularity and volume. Comparisons to our previous two-dimensional modeling framework reflect enhanced model prediction of residual tumor burden through the inclusion of additional imaging slices of patient-specific data.

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Keywords: Tumor; Mechanics; Mathematical; Computational; Oncology; Finite element

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1. Introduction

There is considerable interest in the use of computational methods to study and model the growth and behavioral dynamics of cancer [1,2]. With the significant advancement of imaging methodologies [3], efforts have begun to incorporate imaging data into macroscopic models of tumor cell growth in order to include elements of patient specificity into these modeling efforts [4,5]. We [6,7] and others [8,9] have shown the significant potential of using computational methods in a patient-specific and predictive framework to forecast [10] the response of tumors to therapy. This application is particularly compelling as it offers the opportunity for biologically-inspired computational methodologies to direct therapeutic interventions. In particular, predictive modeling strategies are potentially of great clinical significance in the setting of neoadjuvant therapy (NAT; i.e., therapy before surgery) of breast cancer.

In the neoadjuvant setting, breast cancer patients receive therapy to reduce tumor burden to a size more amenable to surgery. Furthermore, patients whose primary breast tumor responds to NAT are more likely to respond systemically, while a lack of response in the primary tumor makes systemic response unlikely. Importantly, patients who have no residual tumor burden (i.e., a pathological complete response, pCR) at the conclusion of NAT have increased survival; conversely, patients who have residual disease at the conclusion of NAT are at increased risk of early recurrence and death [11,12]. If it could be definitively determined – early in the course of NAT – that a particular therapeutic regimen is unlikely to achieve a pCR, the treating physician could discontinue the ineffective (and potentially toxic) treatment and substitute an alternative regimen. With the numerous options for NAT that have become available, development of a method to predict response early in the course of therapy is a highly relevant clinical objective. Unfortunately, conventional, tissue-based biomarkers of response require invasive biopsy with concomitant pain and morbidity as well as sampling errors due to tumor heterogeneity. Standard tumor size-based methods of evaluating response may lag behind biological response and are therefore not well-suited for predicting eventual response [13–15]. Moreover, size-based techniques may underestimate early efficacy for targeted agents exhibiting predominantly cytostatic rather than cytotoxic effects [14–16]. Accurate model prediction strategies would offer the chance to intervene in the case of prediction of failed therapy and potentially allow for a switch to a more effective regimen. To address this problem, we have recently been focused on models of tumor growth and response to therapy that incorporate important biological aspects of the cancer cell niche, such as coupling with the mechanical nature of the extracellular matrix.

Recent evidence has identified an increasingly important role for the supportive extracellular matrix stroma and demonstrated that surrounding tissue mechanics is active in shaping the growth and response of tumors [17–23]. For example, mechanical aspects of the extracellular matrix, such as tissue density and stiffness, have been linked to cancer cell proliferation and motility [24–28], with strong correlations to the aggressiveness of particular tumors. Extracellular matrix stiffness has been shown to act in a mechanoinhibitory role, whereby increased mechanical stiffness limits the expansion (proliferation and migration) in a cancer cell invasiveness-dependent manner [29–31]. These mechanobiology discoveries have motivated efforts to incorporate mechanical coupling into predictive modeling frameworks to enhance biological specificity [6,7,9,32–39]. In previous work [6,7,40], we introduced and tested the hypothesis that interactions between cancer and tissue mechanics at the macroscopic level could be described by a coupled reactive–diffusive mechanics system, where tumor cells grow and diffuse in space depending on their mechanical environment. While there is considerable data on the use of mathematical modeling to describe tumor growth and response to therapy, previous approaches are often not of the form that can be easily applied to clinical data to generate testable predictions in individual patients. Therefore, rather than use our modeling framework to simulate tumor growth *de novo*, we leverage the rich non-invasive imaging data available to construct and test patient-specific models [5–7]. Thus, we developed a mathematical approach to integrate quantitative *in vivo* imaging data into biophysical mathematical models of tumor growth in order to predict eventual response based on measurements performed before and early in the course of NAT. More specifically, we modified the reaction–diffusion model of tumor growth to include mechanical coupling to the surrounding tissue stiffness, creating a mechanically-restricted cell diffusion model. The combination of quantitative, medical imaging data to initialize and guide a mechanistic understanding provided by mathematical models has demonstrated a compelling strategy for these complex evaluations. In particular, we have previously shown that model predictions of tumor cellularity are correlated with data observations and predictive of eventual response to therapy [6,7].

In this work, we make an important step forward by extending our work to a full three-dimensional representation of the biophysics. In past work, our strategies had been limited to two-dimensional planar assumptions, i.e. we only assessed the imaging slice at the center of the tumor and assumed a plane strain approximation, implying a thick body

where through-plane strain is negated, with similar planar diffusive growth. This dimensionality reduction greatly simplified the computational approach by significantly reducing the degrees of freedom of the forward model as well as allowing for simpler numerical approaches for solution of the inverse parameter estimation problem. However, with full volumetric imaging data readily available, extension to three spatial dimensions is technically feasible and would enhance the amount of patient-specific data included in the model. Thus, in this work we present an important development over our previously described mechanically-coupled, reaction–diffusion modeling framework for predicting the response of breast cancer to NAT using a full three-dimensional volumetric analysis. We present derivations of our forward mathematical model and solution of the inverse parameter estimation problem. We also perform an *in silico* simulation study to assess the ability of our methodology to accurately estimate key model parameters. Finally, we present preliminary modeling results from two patients (one patient that achieved a pCR at the conclusion of NAT and one patient that did not) in order to demonstrate and assess model predictions on clinical data types.

2. Methods

2.1. Mathematical model for tumor growth and response to therapy

We have previously presented a mechanically-coupled reaction–diffusion model to describe the growth and response of tumor cells to NAT [6,7]. The coupled partial differential equations governing this mechanoinhibitory tumor growth behavior are shown in Eqs. (1)–(3):

$$\frac{\partial N(\bar{x}, t)}{\partial t} = \nabla \cdot (D \nabla N(\bar{x}, t)) + k(\bar{x}) N(\bar{x}, t) \left(1 - \frac{N(\bar{x}, t)}{\theta} \right), \quad (1)$$

$$D = D_0 e^{-\gamma \sigma_{vm}(\bar{x}, t)}, \quad (2)$$

$$\nabla \cdot \sigma - \lambda \nabla N(\bar{x}, t) = 0. \quad (3)$$

Eq. (1) describes the change in tumor cell number, $N(\bar{x}, t)$, as a combination of tumor cell diffusion and logistic growth, where D is the apparent cell diffusion coefficient, k is the spatially-dependent growth rate, and θ is the tumor cell carrying capacity. Eq. (1) is coupled to tissue mechanics *via* Eq. (2), which describes the tumor cell diffusion coefficient as a local modification of the global tumor cell diffusion coefficient, D_0 , by the von Mises stress, σ_{vm} , and an empirically derived coupling constant, γ . As discussed in [7,40], in this model the von Mises stress represents local distortional strain energy and is used to represent a description of the loading conditions due to tumor growth. This behavior replicates the mechanoinhibitory effects widely seen from *in vitro* and *in vivo* studies whereby cancer cells exhibit restricted diffusivity when encountering areas of high stress intensity [29–31]. Eq. (3) represents mechanical equilibrium with an expansive force due to tumor cell growth. This tumor cell growth force is determined by changes in cell number and a coupling constant λ . We assume an isotropic Hookean linear elastic constitutive material model:

$$\nabla \cdot G \nabla \vec{u} + \nabla \frac{G}{1-2\nu} (\nabla \cdot \vec{u}) - \lambda \nabla N(\bar{x}, t) = 0, \quad (4)$$

where \vec{u} is the displacement vector, and G and ν are material properties of shear modulus and Poisson's ratio, respectively. We use the Galerkin weighted residual method and linear tetrahedrons to perform spatial integration. Therefore, the coupled displacement (u , v , w represent the x , y , z displacements, respectively) and cell number (N) variables are explicitly approximated as:

$$\begin{aligned} u(x, y, z, t) &\approx \sum_{j=1}^4 u_j \phi(x, y, z, t); & v(x, y, z, t) &\approx \sum_{j=1}^4 v_j \phi(x, y, z, t); \\ w(x, y, z, t) &\approx \sum_{j=1}^4 w_j \phi(x, y, z, t) & N(x, y, z, t) &\approx \sum_{j=1}^4 N_j \phi(x, y, z, t). \end{aligned} \quad (5)$$

Thus, using Eq. (5), expansion of Eq. (1) yields:

$$\left\langle \sum_j \frac{\partial N_j}{\partial t} \phi_j \phi_i \right\rangle + \left\langle D \sum_j N_j \nabla \phi_j \cdot \nabla \phi_i \right\rangle - \left\langle k \sum_j N_j \phi_j \phi_i \right\rangle + \left\langle \frac{k}{\theta} \sum_j N_j^2 \phi_j \phi_i \right\rangle = \int D \nabla N \cdot \hat{n} \phi_i ds \quad (6)$$

which can be condensed into:

$$[M] \left\{ \frac{\partial N_j}{\partial t} \right\} + [L] \{N_j\} + [H] \{N_j^2\} = \int D \nabla N \cdot \hat{n} \phi_i ds \quad (7)$$

where,

$$M_{ij} = \langle \phi_j, \phi_i \rangle \quad (8)$$

$$L_{ij} = \langle D \nabla \phi_j \cdot \nabla \phi_i \rangle - k M_{ij} \quad (9)$$

$$H_{ij} = \frac{k}{\theta} M_{ij}. \quad (10)$$

Using a fully explicit forward Euler solution in the time domain we capture,

$$[M] \{N^{k+1}\} = [M - \Delta t L] \{N^k\} - [\Delta t H] \{N^k\}^2, \quad (11)$$

which can be condensed as:

$$[A_N] \{N^{k+1}\} = \{b_N^k\}. \quad (12)$$

Combining the cell number expression, Eq. (12), with the linear elastic mechanical equilibrium expression, Eq. (3) (weighted residual expression not shown), the full weighted residual expression for the system of equations can be written as:

$$[A] \{\bar{x}\}^{t+\Delta t} = \{b(\bar{x})^t\}, \quad (13)$$

where the coupled solution vector, \bar{x} , is given as,

$$\{\bar{x}\} = \begin{Bmatrix} u \\ v \\ w \\ N \end{Bmatrix} = \begin{Bmatrix} x \text{ displacement} \\ y \text{ displacement} \\ z \text{ displacement} \\ \text{cell number} \end{Bmatrix}, \quad (14)$$

and the local stiffness matrix, A is:

$$[A_{ij}] = \begin{bmatrix} G \left\langle \frac{2(1-\nu)}{1-2\nu} \delta_{xx} + \delta_{yy} + \delta_{zz} \right\rangle & G \left\langle \frac{2\nu}{1-2\nu} \delta_{yx} + \delta_{xy} \right\rangle & G \left\langle \frac{2\nu}{1-2\nu} \delta_{zx} + \delta_{xz} \right\rangle & \lambda \langle \delta_x \rangle \\ G \left\langle \frac{2\nu}{1-2\nu} \delta_{xy} + \delta_{yx} \right\rangle & G \left\langle \delta_{xx} + \frac{2(1-\nu)}{1-2\nu} \delta_{yy} + \delta_{zz} \right\rangle & G \left\langle \frac{2\nu}{1-2\nu} \delta_{zy} + \delta_{yz} \right\rangle & \lambda \langle \delta_y \rangle \\ G \left\langle \frac{2\nu}{1-2\nu} \delta_{xz} + \delta_{zx} \right\rangle & G \left\langle \frac{2\nu}{1-2\nu} \delta_{yz} + \delta_{zy} \right\rangle & G \left\langle \delta_{xx} + \delta_{yy} + \frac{2(1-\nu)}{1-2\nu} \delta_{zz} \right\rangle & \lambda \langle \delta_z \rangle \\ 0 & 0 & 0 & A_N \end{bmatrix}. \quad (15)$$

The right hand side forcing conditions for the previous solution step and boundary conditions, b , are:

$$[b_i] = \begin{bmatrix} \vec{x} \cdot \int \sigma(t + \Delta t) \cdot \hat{n} \phi_i ds \\ \vec{y} \cdot \int \sigma(t + \Delta t) \cdot \hat{n} \phi_i ds \\ \vec{z} \cdot \int \sigma(t + \Delta t) \cdot \hat{n} \phi_i ds \\ b_N + \Delta t \int D \nabla N(t + \Delta t) \cdot \hat{n} \phi_i ds \end{bmatrix}. \quad (16)$$

This system of equations is solved *via* a fully explicit finite difference method in the time domain and finite element method (FEM) for the spatial domain. The simulation software is custom built and utilizes PETSc as a sparse matrix solver library [41–43]. To solve, parameter values (D_0 , k , θ , λ , γ , E , ν) are assigned and the stiffness matrix $[K]$ is assembled. As the solution is fully explicit in time, the stiffness matrix can be pre-computed to accelerate the model solution as it does not change during iterative time stepping as all terms with respect to changing cell numbers are applied as a right hand side forcing function. Boundary conditions are applied and the solution is iteratively calculated in a time stepping loop where we assemble b (parameter values, boundary conditions, and data from previous time steps, N^k and σ_{VM}^k), solve for $\{x^{k+1}\}$, partition the solution into displacements and cell numbers, calculate von Mises stress, and then continue to the next time step.

2.2. Solving the inverse problem (parameter estimation)

The key model parameters, which include a global tumor cell diffusion coefficient (that is locally modified by von Mises stress) and spatially-dependent proliferation coefficients at each node in the FE mesh, are estimated using the model and tumor cell data at two time points. The model is initialized by data obtained at the pre-treatment time point and an initial guess of parameter values, and parameter estimation is performed by iteratively minimizing the objective function using a Polak–Ribiere conjugate gradient algorithm [44,45] using tumor cell data acquired after the first cycle of NAT. The objective function is the sum of squared differences between the model prediction and the data observation at the second time point, and is given by,

$$\psi = \sum_{i=1}^n (N_{data}(i) - N_{model}(i))^2, \quad (17)$$

where N_{data} is the experimentally estimated number of tumor cells and N_{model} is the model predicted number of tumor cells at each node.

The adjoint method is used to evaluate the gradient at each optimization step [9,46–48]. The inverse problem estimates a global tumor cell diffusion coefficient and proliferation coefficients at each node, with approximately 25,000–30,000 nodes in a FE mesh for a representative patient. However, due to the size and spatial position of the tumor at the time points used for parameter estimation, approximately 3000–5000 proliferation parameters are changed from the initial guess for a representative patient. Use of the adjoint method allows a more efficient calculation of parameter sensitivity over traditional finite-difference based gradient evaluation methods, and is necessary due to the large number of parameters to be estimated. A finite difference based gradient evaluation would require a time-prohibitive amount of forward model evaluations per optimization step (number of parameters + 1) as opposed to the adjoint method which requires only two forward model solves per optimization step [49].

While a brief description of the adjoint state and parameter sensitivity equations follow, for a complete derivation of the adjoint method the interested reader is referred to one of several Refs. [9,46–48]. With Eq. (13) as the primary problem and Eq. (17) as the objective function, the adjoint state equation can be written with respect to the unknown parameter vector, P , as,

$$A(P)\phi = \left(\frac{\partial \psi(N, P)}{\partial N} \right)^T, \quad (18)$$

where ϕ is the adjoint state variable. The parameter sensitivity of the objective function (i.e., the gradient) can be written as,

$$\frac{d\psi(N, P)}{dP} = \frac{\partial \psi(N, P)}{\partial P} + \phi^T \left(\frac{\partial b(P)}{\partial P} - \frac{\partial A(P)}{\partial P} N \right). \quad (19)$$

Parameters are estimated though an optimization scheme that iteratively minimizes Eq. (17) using Eq. (19) to calculate the parameter sensitivity. The optimization is terminated by satisfying a relative error tolerance condition where the change in error relative to the previous iteration is less than 1×10^{-5} .

2.3. Populating the model with patient-specific data

The model is populated and parameterized using patient-specific tumor cell data by processing quantitative magnetic resonance imaging (MRI) data. The MRI data sources used to populate the model are: diffusion weighted MRI (DW-MRI), dynamic contrast enhanced MRI (DCE-MRI), and anatomical THRIVE (T_1 high resolution isotropic volume examination). Briefly, DCE-MRI collects images before, during, and after contrast agent infusion [50]. While pharmacokinetic modeling of DCE-MRI data can be used to extract compartmental model parameters describing blood vessel perfusion/permeability and volume fractions, in this work we simply use this data for region-of-interest assignment. We do note, however, that we are actively working to incorporate DCE-MRI data to inform other aspects of tumor growth and response to therapy [51]. DW-MRI collects data that can be used to map the apparent diffusion coefficient (ADC), which describes the rate of thermally induced self-diffusion in tissue [52]. This parameter depends on the amount of microscopic barriers that a diffusing water molecule encounters and thus is used as a correlate to tissue cellularity [52–71]. Finally, THRIVE collects data weighted by the T_1 relaxation properties of tissue. It is used in this work to aid segmentation of breast fibroglandular tissue from adipose tissue, as these tissues exhibit contrast in T_1 relaxation properties.

Imaging data is acquired at three time points: prior to initiating NAT, after one cycle of NAT (typically one to two weeks after the initial time point), and at the conclusion of NAT (typically eight to twelve weeks after the initial time point) as described previously [6,7]. Briefly, MR images for each patient are longitudinally co-registered across all time points to the initial time point using an adaptive basis algorithm non-rigid registration with a tumor volume preserving constraint [72,73]. Registered DCE-MRI data is used to define a tumor region-of-interest (ROI) by manually segmenting the difference image of pre- and post-contrast images. Areas within the ROI with greater than 80% signal intensity increase between the pre- and post-contrast infusion data are defined as tumor; this value has previously been empirically determined to provide a high degree of correlation to the tumor volume as measured on the excised tissue obtained at the time of surgery [74].

ADC values were obtained by fitting the DW-MRI data on a voxel-by-voxel basis to Eq. (20):

$$ADC = \frac{\sum_{i=x,y,z} \ln(S_0/S_i) b_i}{3}, \quad (20)$$

where i is the direction of diffusion-weighting, b_i is the amount of diffusion-weighting, S_0 is the signal intensity in the absence of diffusion gradients, and S_i is the signal intensity in the presence of the diffusion-sensitizing gradient. ADC data for areas defined as tumor were transformed to estimate tumor cell number, $N(\bar{x}, t)$ as described in [57,75],

$$N(\bar{x}, t) = \theta \left(\frac{ADC_w - ADC(\bar{x}, t)}{ADC_w - ADC_{min}} \right), \quad (21)$$

where θ is the carrying capacity (total number of tumor cells that can be contained within a voxel), ADC_w is the ADC of free water at 37 °C (set to $3 \times 10^{-3} \text{ mm}^2/\text{s}$), $ADC(\bar{x}, t)$ is the ADC value at a given position, and ADC_{min} is the minimum ADC value observed, corresponding to the voxel with the largest number of cells. Further details on this approach have been previously reported [57,76].

The full model parameter set (D_0 , k , θ , E , ν , λ , and γ) contains model parameters that are determined by inverse model fitting, assumptions based on literature values, or otherwise empirically determined. Specifically, D_0 and k are determined from model fits. The carrying capacity, θ , is calculated by assuming spherical tumor cells with a packing density of 0.7405 [77] and a nominal tumor cell radius of 10 μm . Material properties of Young's modulus, E , are assumed as 2 and 4 kPa for adipose and fibroglandular tissue, respectively [78]. Anatomical T_1 -weighted MR images are used to segment the breast into fibroglandular and adipose tissue for assigning tissue-based mechanical properties. Poisson's ratio, ν , is assumed as 0.45, reflecting the nature of soft tissue and avoiding numerical instabilities due to incompressibility within a linear elastic model solution. Coupling constants, λ and γ , are empirically determined as 2.5×10^{-3} and $2 \times 10^{-3} \text{ Pa}^{-1}$, respectively, and were empirically determined to yield physiologically realistic deformation fields and von Mises stress maps in prior work [6,7]. The time step, Δt , was selected as 0.25 days and was determined based on an empirical stability analysis. Von Neumann stability analysis [79] of a finite difference forward Euler linear diffusion equation in three dimensions was used to guide the initial selection for the time step based on constraints established on the ratio of the product of the diffusion coefficient and time step to the square of

the spatial discretization length ($D \cdot dt / h^2$). For the most restrictive stability conditions, we select a maximal allowable diffusion coefficient of $1 \times 10^{-7} \text{ m}^2/\text{day}$ and find the minimum spatial step size observed in the discretized mesh for all subjects. Following initial selection of step size, forward simulations of the model were analyzed for numerical oscillations and the step size was adjusted as necessary such that the approximate solution remained bounded without amplification of numerical errors. This stability requirement is necessary due to the selection of a conditionally-stable explicit solution algorithm (Section 2.1). While these do constrain the speed of time evolution, they do have significant advantages in algorithmic simplicity. For mesh discretization, an empirical selection of nominal element edge length was used to guide mesh refinement using forward simulations of the model on a mesh with incrementally increasing refinement. Comparison of solutions at prescribed locations within the mesh was assessed for convergence using a tolerance of less than 1% difference in solution from successive simulations at increasing refinement levels (and decreasing nominal element edge length).

2.4. *In silico* simulation study

We performed an *in silico* simulation study by assuming an initial tumor seed of 10% of the cell carrying capacity and placed centrally within a realistic breast model domain at a single node. A finite element mesh was generated using custom-built mesh generation software [80] and was composed of four-node tetrahedrons with a nominal edge length of 3 mm. Global diffusion, D_0 , was assigned as $1 \times 10^{-8} \text{ m}^2/\text{day}$. Spatial growth rate, k , was assigned at each node as 0.1 day^{-1} . Other model parameters were assigned as described in Section 2.3. The model was used to ‘grow’ the tumor to a second time point, 14 days later. Using the initial and ‘grown’ tumor cellularity at each time point as data observations, we estimate diffusion and growth rate parameters as described in Section 2.2 using initial guesses of $8 \times 10^{-9} \text{ m}^2/\text{day}$ and 0.08 day^{-1} for diffusion and growth rate, respectively. This simulation study represents a controlled system to assess the accuracy of the inverse methodology.

2.5. Experimental study

To demonstrate data-driven model predictions within our full volumetric modeling framework, we performed a limited proof-of-principle experimental study to assess model prediction performance relative to data observations. MRI data from an existing breast NAT patient imaging database [69,81] was retrieved for two patients, one achieving pCR, and one with residual tumor burden at the conclusion of NAT. MRI data sets were collected prior to beginning NAT, after one cycle of NAT (typically 1–2 weeks), and at the conclusion of NAT (typically 8–12 weeks). MRI data acquisition parameters for this study population have been previously reported [69,81]. Briefly, MRI was performed using a Philips 3T Achieva MR scanner (Philips Healthcare, Best, The Netherlands) with a 16-channel double-breast coil. DCE-MRI data was acquired *via* a $192 \times 192 \times 20$ acquisition matrix over a sagittal square field of view (22 cm^2) with slice thickness of 5 mm, one signal acquisition, $TR/TE/a = 7.9 \text{ ms}/1.3 \text{ ms}/20^\circ$, and SENSE factor of 2. Twenty-five time points, with a temporal resolution of 16 s, were collected with contrast agent infusion occurring after collection of the third time point. The contrast agent, 0.1 mmol/kg of gadopentetate dimeglumine (Magnevist, Wayne, NJ) was delivered through an antecubital vein catheter at 2 mL/s, followed by a saline flush, *via* a power injector (Medrad, Warrendale, PA). DW-MRI data was acquired with a single-shot spin echo, echo planar imaging sequence in three orthogonal diffusion-encoding directions with b -values of 0 and 500 or 600 s/mm^2 , 96×96 (reconstructed to 144×144) acquisition matrix over a $19.2 \text{ cm} \times 19.2 \text{ cm}$ field of view using SENSE parallel imaging (acceleration factor = 2) and spectrally-selective adiabatic inversion recovery fat saturation while subjects were freely breathing. Twelve sagittal slices with slice thickness of 5 mm, $TR = 3080 \text{ ms}$, $TE = \text{‘shortest’}$ (41 or 60 ms), $\Delta = 19.8$ or 29 ms, and $\delta = 10.7$ or 21 ms, and 10 signal averages. THRIVE (T_1 high resolution isotropic volume examination) anatomical data was acquired *via* a $400 \times 400 \times 129$ acquisition matrix over a $20 \text{ cm} \times 20 \text{ cm} \times 12.9 \text{ cm}$ transverse field of view, one signal acquisition, $TR/TE/\alpha = 6.43 \text{ ms}/3.4 \text{ ms}/10^\circ$.

For each patient, a finite element mesh was generated from anatomical imaging data using custom-built, semi-automatic software. Briefly, breast MR image volumes were manually segmented at the chest wall using ITK-SNAP [82]. The boundary surface of the breast was generated through a marching cubes extraction [83] that was followed by radial basis function smoothing (FastRBF toolkit, FarField Technology, Christchurch, New Zealand). A finite element mesh was generated [80] and was composed of four-node tetrahedrons with a nominal edge length of 3 mm. Model parameters were estimated as described in Section 2.2, and the model was run forward in time to

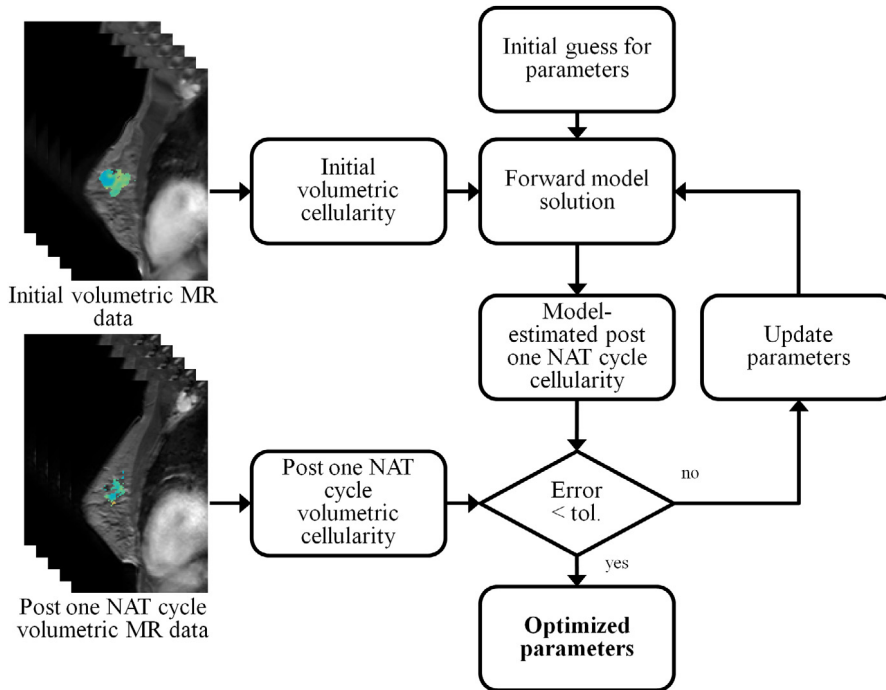


Fig. 1. Schematic of the inverse parameter estimation procedure. The volumetric MR data from the initial and post one NAT cycle is converted to estimate tumor cellularity at each time point. Given an initial guess of the model parameters, the model is initialized by the tumor cellularity at the initial time point and used to generate an estimate of the tumor cellularity at the post one NAT cycle time point. The model estimate is compared to the data observation for the post one NAT cycle cellularity and parameters are iteratively refined until convergence below a preset tolerance to yield an initial estimate of the parameters.

generate predictions of tumor burden at the conclusion of NAT. Model predictions were compared to data observations for both total predicted tumor cellularity and tumor volume. A schematic of this procedure is outline in Fig. 1.

For comparison of the full-volumetric extension to our prior work, we also performed a two-dimensional model prediction for one patient with residual tumor burden at the conclusion of NAT, using a two-dimensional planar assumption that assessed the imaging slice at the center of the tumor, as described previously [6,7].

2.6. Biomechanical constitutive model simulation study

To demonstrate the effects of biomechanical constitutive model selection, we performed a limited study to investigate simulations of tumor growth and response to therapy with respect to changing the assumed biomechanical constitutive model used for mechanical coupling to our mechano-inhibitive, reaction–diffusion model of tumor growth. We compared the results of coupling to a linear elastic biomechanical model, as described in Section 2.1, and a hyperelastic neo-Hookean nonlinear biomechanical model [84] given by the strain energy function:

$$W = \frac{\mu}{2} (I_1 - 3) - \mu \ln J + \frac{\lambda}{2} (\ln J)^2, \quad (22)$$

where I_1 is the first invariant of the right Cauchy–Green deformation tensor, J is the Jacobian of the deformation tensor, and μ and λ are Lamé parameters defined as:

$$\lambda = \frac{\nu E}{(1 + \nu)(1 - 2\nu)} \quad (23)$$

$$\mu = \frac{E}{2(1 + \nu)}. \quad (24)$$

The open-source biomechanical nonlinear finite element solver FEBio [85] with a custom-designed pre-processing interface based on GIBBON [86] was used for neo-Hookean biomechanical model coupling. Estimated model param-

Table 1

Performance results for *in silico* parameter estimation of global diffusion and local proliferation reflect the initial guess, ground truth, parameter estimate, and percent error of the parameter estimate as compared to ground truth.

	Initial guess	Ground truth	Estimated	Error
D_0	8.00×10^{-9}	1.00×10^{-8}	9.74×10^{-9}	2.59%
k	0.0800	0.1000	0.0995	0.5%

Table 2

Tumor cellularity and volumetric results reflect the imaging-based estimation of tumor burden at the initial, post one NAT cycle, and final time points in addition to the model prediction results at the final time point. Cellularity numbers are normalized by 1×10^9 tumor cells and volumes are expressed as cm^3 .

	Initial	Post one NAT cycle	Final	Prediction
Responsive cellularity	1.435	0.406	0.056	0.097
Non-responsive cellularity	3.751	1.425	0.165	0.179
Responsive volume	9.338	2.532	0.302	1.155
Non-responsive volume	23.99	9.355	1.084	2.417

eters were taken from the experimental study (Section 2.5). For the neo-Hookean model, assumed Poisson's ratio was varied between 0.45, as used in the linear elastic biomechanical model, to a value of 0.499. The forward model, with either linear elastic or neo-Hookean biomechanical model coupling, was used to simulate tumor growth and response to therapy from the initial time point to the post one NAT cycle time point and predicted tumor cellularity and volume were compared for the two models.

3. Results

3.1. *In silico* simulation study

We assessed the accuracy of our parameter estimation approach, by performing an *in silico* simulation study. Using the initial and 'grown' tumor cellularity at each time point as data observations, we evaluated the accuracy of our parameter estimation approach. In this controlled study, we found an estimation error less than 3% for diffusion and growth rate. As shown in Table 1, the error in estimating global tumor cell diffusion, D_0 , was 2.59% and the error in estimating the growth rate was 0.5%. Parameter estimation performance is shown in Fig. 2, reflecting the behavior in diffusion, growth rate, and objective function error over approximately 50 iterations prior to termination of iterative estimation due to relative error convergence criteria.

3.2. Experimental study

Figs. 3 and 4 show data from parameter reconstruction and model prediction for a patient achieving pathological complete response at the conclusion of NAT. While we performed a full volumetric analysis, for visualization purposes, Fig. 3 shows data and results at the central slice of the tumor. After converting MRI data (Fig. 3, top row) to tumor cellularity data (Fig. 3, middle row), we used the model to estimate global diffusion and local growth rate parameters (Fig. 3, bottom row, left). Following parameterization, we used the model to estimate the residual tumor burden at the end of NAT (Fig. 3, bottom row, right). While this patient was evaluated pathologically as achieving complete response (i.e., no residual tumor burden), the imaging data at the final time point reflected a slight focal residual tumor burden. Fig. 4 shows three-dimensional volumetric representations of the tumor volume at each data observation time point (Fig. 4, top row) as well as the model prediction (Fig. 4, bottom row). The model is seen to accurately predict a small residual tumor burden, with similar total cellularity (see Table 2). However, the model is shown to overestimate the volume as well as incorrectly identify the location of the residual tumor burden, placing the predicted residual tumor slightly inferior to the observed location (approximately 1.3 cm).

Similarly, Figs. 5 and 6 show data for a patient identified pathologically with residual tumor burden at the end of NAT. MR data (Fig. 5, top row) is converted to estimate tumor cellularity (Fig. 5, middle row), and the model is used

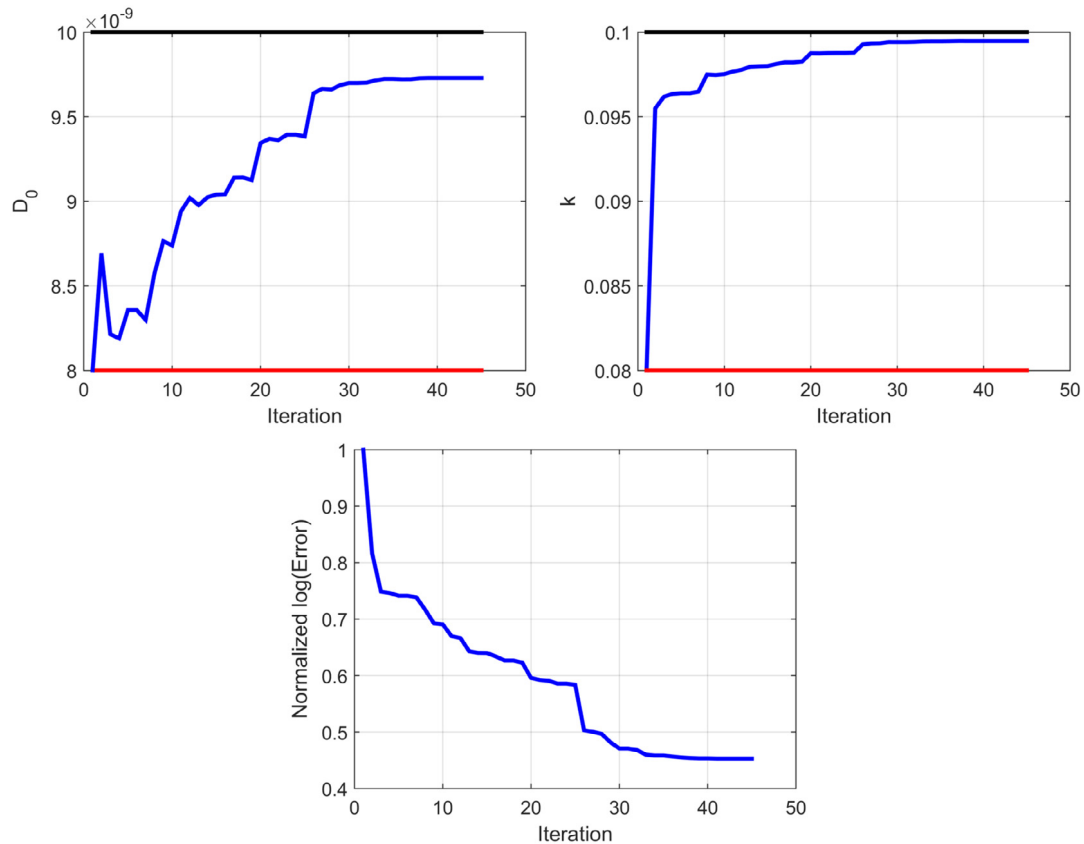


Fig. 2. Parameter estimation performance for the *in silico* study. Top: Tumor cell diffusion, D_0 , and local proliferation, k , are initialized at the first iteration with a guess, given by the value at the red line, and the iterative parameter estimation continues until convergence, given by the blue line. The parameter estimation procedure is shown to exhibit accurate performance in generating parameter estimates as compared to ground truth, given by the value at the black line. Bottom: Log transformed objective function error is normalized to the value at the initial iteration and shows iterative minimization throughout the parameter estimation procedure. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

Table 3

Model estimated parameter results reflect the mean, maximum, and minimum spatial proliferation rates and tumor cell diffusion coefficients for the responsive and non-responsive patients.

	Mean proliferation	Max proliferation	Min proliferation	Tumor cell diffusion
Responsive	−0.1179	0.1398	−0.2860	5.06×10^{-9}
Non-responsive	−0.1158	0.1671	−0.3186	7.91×10^{-8}

to estimate parameters and predict the residual tumor burden at the final time point (Fig. 5, bottom row). Again, while we performed a full volumetric analysis, the central slice plane of imaging data is shown for visualization purposes. The model is shown to accurately predict total tumor cellularity (Table 2), but is seen to overestimate the volume and misidentify the spatial location of the residual tumor (Fig. 6). While a portion of the predicted residual tumor is located within the area of the observed tumor burden, the model predicts additional focal lesions outside that of the area observed in the data observation.

Fig. 7 shows merged and smoothed volumetric renderings reflecting the time course of tumor volumetric changes in both imaging data observation and model prediction during the course of NAT for patients with both pCR and non-pCR tumors. Model estimated parameter results shown in Table 3 reflect slight differences in proliferation and tumor cell diffusion parameters between tumor types. The patient with a pCR tumor is seen to have a slightly lower mean and maximum proliferation and tumor cell diffusion than the patient with a non-pCR tumor.

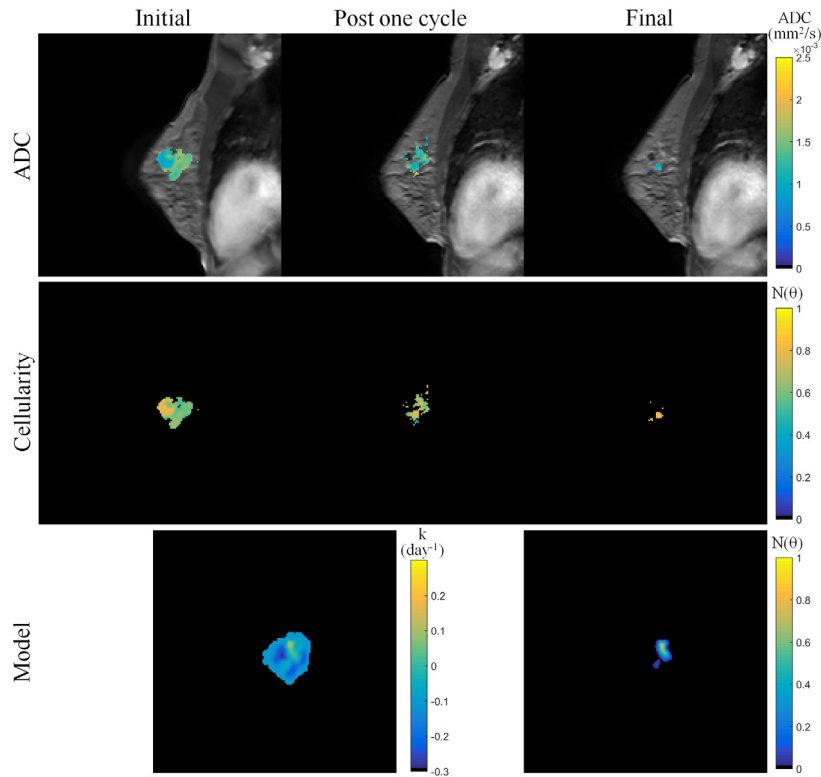


Fig. 3. MR imaging data and mathematical model prediction results for a patient achieving a pathological complete response at the completion of NAT. ADC data at the initial, post one NAT cycle, and final time points (top row) is converted to estimates of tumor cellularity (middle row) using Eq. (21). Global diffusion and spatial proliferation (bottom row, left) model parameters are estimated and the model is used to predict tumor cellularity at the final time point (bottom row, right) and compared to data observations. Though the patient was evaluated at surgery as having achieved a pathological complete response, the imaging data and model predictions at the final time point reflect a very small focal residual tumor. (Note that although we performed a volumetric analysis, for visualization purposes, this figure displays data and results at the central slice of the tumor; see Figs. 4 and 7 for volumetric renderings.)

Comparison of model prediction results from the reduced spatial dimensionality model, presented in prior work, to the full volumetric model presented within this study are shown in Fig. 8 for a patient with residual tumor burden at the conclusion of NAT. Tumor cellularity (Fig. 8, top row) predicted from the two-dimensional model reflects a significant overestimation of cellularity and volume as compared to the imaging data observation, whereas the model prediction from the three-dimensional model more closely matches imaging data. Comparison of the estimated proliferation maps (Fig. 8, bottom row and Table 4) reflects large differences in spatial pattern and magnitude between the two models.

3.3. Biomechanical constitutive model simulation study

We assessed the effects of constitutive biomechanical model selection within our tumor growth model in a limited simulation study. Total tumor cellularity and volume results are compared in Table 5 with a tumor growth model based on coupling to a biomechanical model with a linear elastic or a neo-Hookean nonlinear hyperelastic assumption. Three different values of Poisson's ratio are used for the neo-Hookean model, representing a gradually more incompressible behavior in hyperelasticity. When varying the biomechanical model from linear elastic to nonlinear hyperelastic, both predicted total tumor cellularity and volume reflect a maximum difference of less than 10% in this simulation scenario.

4. Conclusions

We have presented an extension of our image-based mechanically-coupled mathematical model of tumor response to NAT into three spatial dimensions. Using an *in silico* study, we used the model to simulate the growth of a tumor, given an initial tumor 'seed' location. The predictive modeling framework demonstrates accuracy in estimation of the

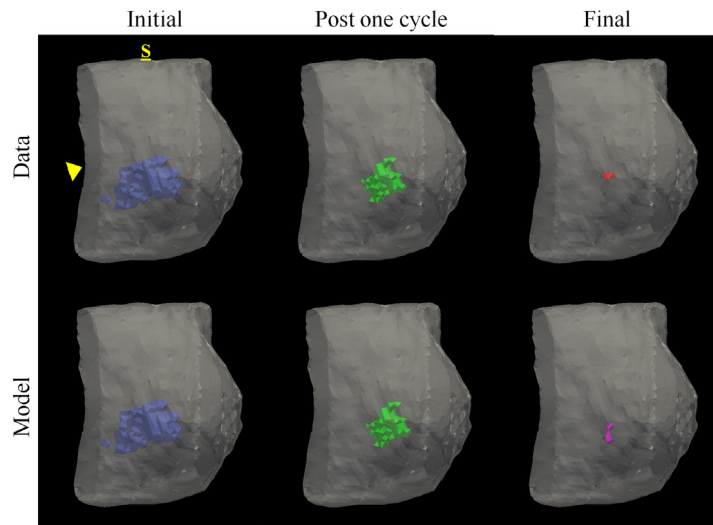


Fig. 4. Volumetric rendering of the tumor cell volumes taken from imaging data and model prediction are displayed within the breast volume, segmented at the chest wall (arrowhead), for a patient achieving a pathological complete response. Data is collected at the initial (blue), post one NAT cycle (green), and final (red) time points. The model is informed with data at the initial time point and parameters are estimated to minimize model/data error the post one NAT cycle time point. The model is then used to predict the residual tumor burden at the final time point (purple). The label, S, designates the superior position of the breast. This patient was evaluated at surgery as having achieved pCR, but imaging data and model prediction at the final time point reflect a very small focal residual tumor. Similar to imaging observation, the model predicts a small residual tumor burden; however, the model incorrectly predicts the location of the residual tumor burden as slightly inferior to the observed location. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

Table 4

Model estimated parameter results reflect the mean, maximum, and minimum spatial proliferation rates for the non-responsive patient assessed with the two-dimensional and three-dimensional (assessed at the same central imaging slice) modeling frameworks.

	Mean proliferation	Max proliferation	Min proliferation
2D	−0.2012	0.3229	−0.8762
3D (central slice)	−0.1094	0.0398	−0.2707

model parameters of diffusion and growth rate with less than 3% error when compared to the ground truth parameters used to simulate tumor growth. In a preliminary experimental study to demonstrate predictive performance, we used our modeling approach to predict the response to NAT for two patients, one that achieved pathological complete response and one that did not. The model accurately predicts the residual tumor cellularity as compared to observed quantitative imaging data estimates of tumor cellularity. In our preliminary data, however, the model is challenged with accurate spatial placement of the predicted location of residual tumor burden, with one patient showing slight misalignment and one patient with more pronounced misalignment demonstrating additional focal tumor lesions that were found to have completely resolved in follow-up imaging. However, it is important to note that the prediction framework is based on a very challenging, but clinically significant, timeframe of imaging data acquisition in which imaging data is collected before and after only a single cycle of NAT and is used to predict eventual tumor burden. Patients continue to receive additional cycles of NAT, approximately 8–12 total, prior to data observations at the final time point. Therefore it is important to recognize the value of predictive modeling for guidance of suspicion for residual tumor burden within the context of clinical goals for interventional therapy (i.e. discriminating between pCR and non-pCR), rather than exact prediction of tumor spatial placement. While a predictive framework that could accurately resolve exact tumor location would be ideal, there are additional nonlinear effects currently unaccounted for in the model that occur between the final time point of data used to inform predictive models and the time point at which model prediction is compared to data observation. We return to this limitation below.

We also present a comparison of our results from the full volumetric model predictions of this study to that of the reduced spatial dimensionality modeling framework presented in prior work. We show that incorporating additional

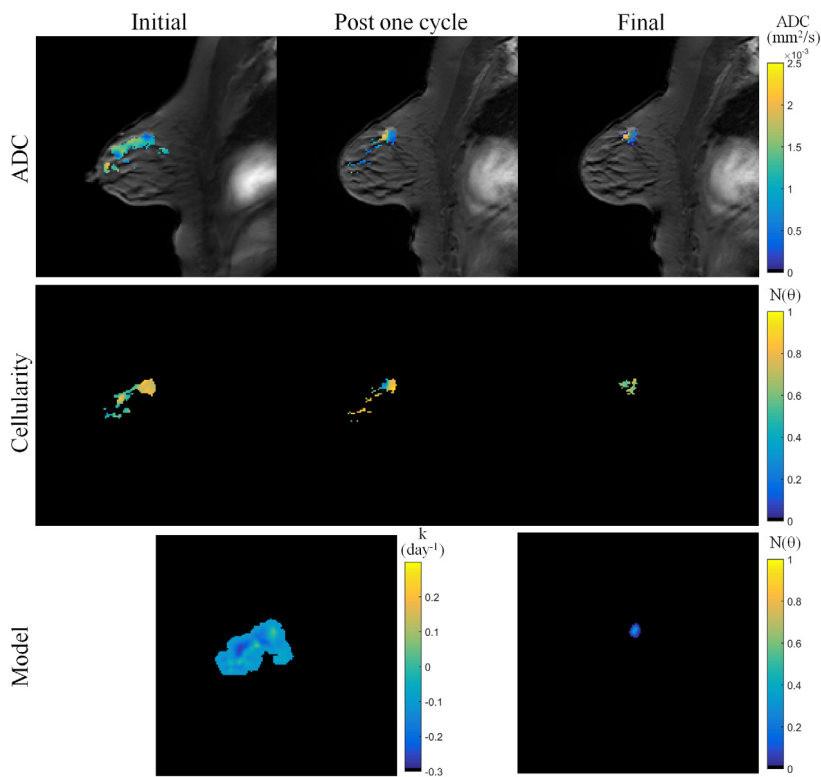


Fig. 5. MR imaging data and mathematical model prediction results for a patient with residual tumor burden at the completion of NAT. ADC data at the initial, post one NAT cycle, and final time points (top row) is converted to estimates of tumor cellularity (middle row) using Eq. (21). Global diffusion and spatial proliferation (bottom row, left) model parameters are estimated and the model is used to predict tumor cellularity at the final time point (bottom row, right) and compared to data observations. (Note that although we performed a volumetric analysis, for visualization purposes, this figure displays data and results at the central slice of the tumor; see Figs. 6 and 7 for volumetric renderings.)

Table 5
Tumor cellularity and volumetric results for constitutive model simulation study comparing linear elastic and neo-Hookean hyperelastic biomechanical coupling. Cellularity numbers are normalized by 1×10^9 tumor cells and volumes are expressed as cm^3 . Percent difference is expressed relative to the linear elastic model.

	Tumor cellularity	% Difference	Tumor volume	% Difference
Linear elastic	0.3674	n/a	3.572	n/a
Neo-Hookean ($\nu = 0.45$)	0.3657	0.469%	3.563	0.249%
Neo-Hookean ($\nu = 0.49$)	0.3385	7.87%	3.287	7.96%
Neo-Hookean ($\nu = 0.499$)	0.3317	9.70%	3.225	9.70%

patient-specific imaging data results in large changes to the parameter estimates extracted from the model. This additional data is then shown to enhance the accuracy of the model predicted cellularity, more closely matching imaging data observations. In previous investigations that used our simplified reduced spatial dimensionality model, we have demonstrated qualitative and quantitative comparisons demonstrating the improved performance of our model-to-data match using our mechanoinhibitive model of tumor growth [6,7]. In future work we will more fully explore the increased fidelity that can be achieved using the fully three dimensional realization presented herein.

While in this work we have eliminated the assumption of reduced spatial dimensionality, several limitations still remain in our current approach. One such limitation of this study is that we assume an isotropic Hookean linear elastic constitutive mechanical model. However, the true constitutive nature of soft tissue is well-known to exhibit particularly complex non-linear and multi-physics behavior. Therefore, selection of an appropriate constitutive model for use in computational analysis of biological tissue is often controversial and presents unique challenges. Soft tis-

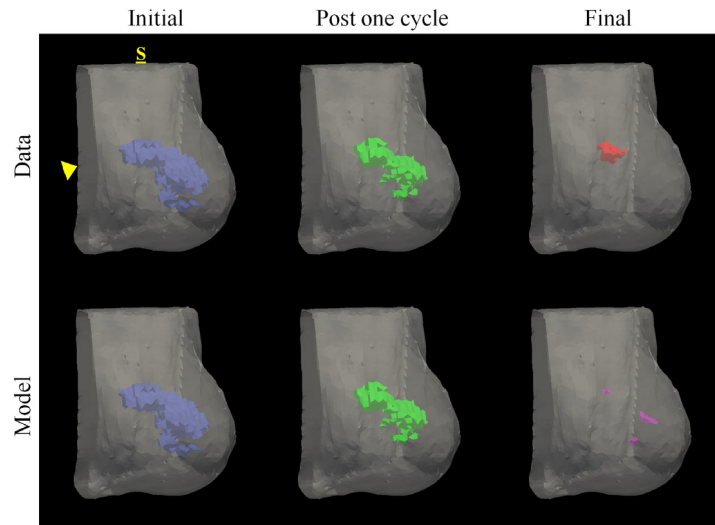


Fig. 6. Volumetric rendering of the tumor cell volumes taken from imaging data and model prediction are displayed within the breast volume, segmented at the chest wall (arrowhead), for a patient with residual tumor burden. Data is collected at the initial (blue), post one NAT cycle (green), and final (red) time points. The model is informed with data at the initial time point and parameters are estimated to minimize model/data error the post one NAT cycle time point. The model is then used to predict the residual tumor burden at the final time point (purple). The label, S, designates the superior position of the breast. While there is significant response to NAT, residual focal tumor burden is observed. A portion of the predicted residual tumor is located within the area of the observed tumor burden, however the model predicts additional focal lesions outside that of the area observed in the imaging data. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

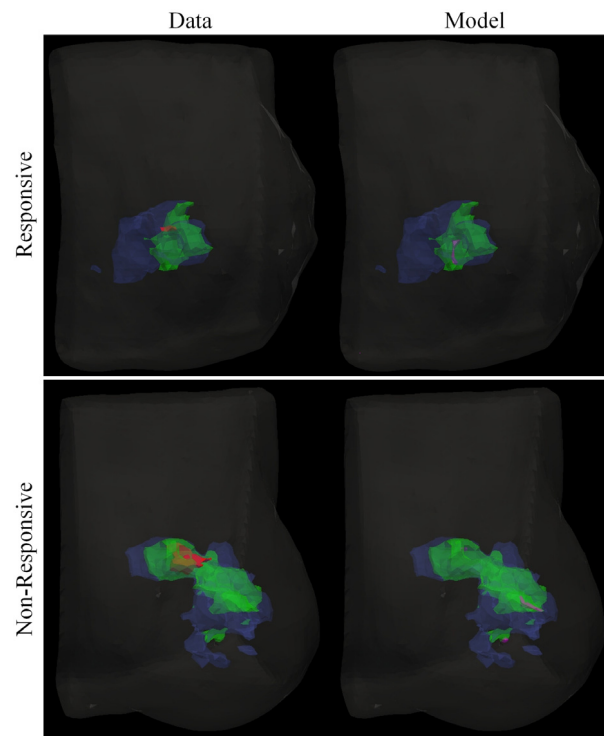


Fig. 7. Merged volumetric renderings of the tumor cell volumes taken from imaging data and model predictions are smoothed and displayed within the breast volume for patients with pCR and non-pCR tumors. Data is collected at the initial (blue), post one NAT cycle (green), and final (red) time points. The model is calibrated using data at the initial and post one NAT cycle time points and used to predict the residual tumor burden at the final time point (purple). (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

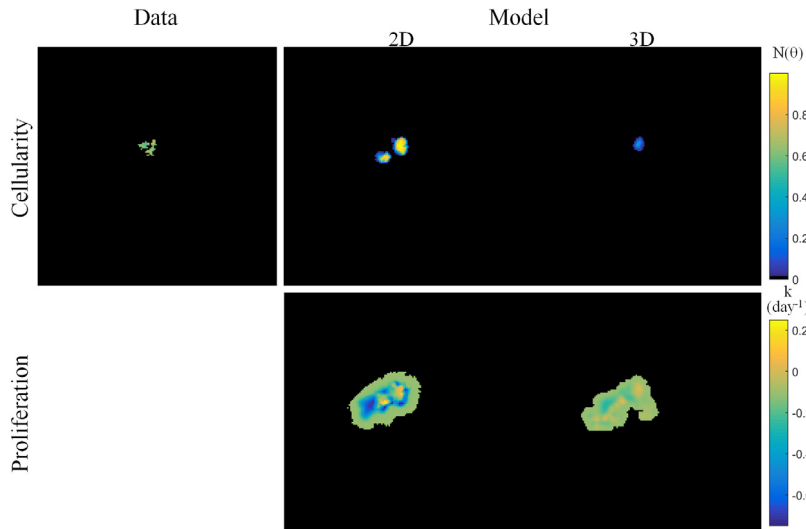


Fig. 8. Comparison of two-dimensional (from prior work) to three-dimensional image-based mathematical model prediction results for a patient with residual tumor burden at the completion of NAT. For two-dimensional model prediction, ADC-based estimates of tumor cellularity at the central imaging slice, as shown in Fig. 5, are used to calibrate the model and predict residual tumor burden. Tumor cellularity at the final time point (top row) are shown for imaging data, two-dimensional model prediction, and three-dimensional model prediction (at the central imaging slice) and reflect that three-dimensional model prediction is seen to more closely match imaging data observations than the two-dimensional model. Spatial proliferation maps (bottom row) reflect differences in estimated model parameters for each model. (Note that although we performed a full volumetric analysis for the three-dimensional model, for comparisons to the two-dimensional case, this figure displays results at the central slice of the tumor.).

sue, unlike conventional materials, exhibits many highly complex intrinsic properties in addition to the computational challenges encountered in implementing, solving, and determining accurate material properties for more sophisticated constitutive models. Investigators have employed anisotropic [87], hyperelastic [88], viscoelastic [89], and poroelastic [90] material models to represent soft tissue in mechanical modeling efforts, among others. In a limited simulation study, we show that the first-order linear elastic simplification we employ in this effort introduces a small degree of numerical model error, but is less than the uncertainty of the underlying ADC-based measurement of observed cellularity [91–93]. This study, while preliminary, suggests that the uncertainty due to biomechanical constitutive model selection is contained within the experimental error of a data-driven tumor growth modeling framework. However, it is important (in future efforts) to more fully understand the true nature of material model assumptions, both simplified and complex, with robust validation studies to further explore the consequences of model selection and reducing first order assumptions from the perspective of clinical and interventional goals for prediction of NAT response. Additional technical model limitations include the limited degrees of freedom associated with the use of linear tetrahedral elements rather than quadratic tetrahedral or hexahedral elements.

Another limitation is the use of ADC from DW-MRI data as a linear correlate to tumor cellularity. ADC can also be shown to depend on several factors other than cellularity, such as extracellular matrix density, edema, and inflammatory infiltrates, among others. While we agree that exact ADC-to-cellularity relationships are likely influenced by other mitigating factors, previous studies confirm its use as a first order approximate relationship. ADC has been shown to correlate inversely with cellularity, and has also demonstrated strong potential as an independent measure for assessing treatment response for anti-cancer therapy in both preclinical and clinical studies [52–71]. Additionally, pre-clinical validation studies have shown strong significant correlation of ADC to histologically confirmed extracellular space fraction [71].

Lastly, it is important to recognize that predictions are synthesized by a calibration/fitting process that uses only two imaging time points, before and after a single cycle of NAT. The consequences of this limitation are that it implicitly assumes a fixed spatial proliferation rate map in time for subsequent cycles of NAT. Subsequent cycles of NAT will undoubtedly change the spatial proliferation and migration characteristics, however we have no additional imaging data to guide the spatiotemporal evolution of proliferation or migration terms. Additionally, many patients receive multi-agent therapy with temporal delays in the administration of therapeutic agents which may occur after the acquisition of imaging data. Future studies that include additional imaging time points (currently ongoing), or

predictive models that can simulate additional specific therapeutic agent effects *de novo*, will be important for further development of a model that can compensate for this effect.

5. Summary

The modeling methodology presented in this contribution represents a data-driven framework for prediction of residual tumor burden in breast cancer patients following neoadjuvant therapy. Previously, we have taken a commonly used reaction–diffusion tumor growth model and extended it by incorporating known mechanoinhibitory effects that have been previously demonstrated in biological perturbation studies [27–31]. We previously demonstrated that a reduced spatial dimensionality predictive model exhibits significant potential for use as a predictor of response to therapy and indicates the promise of image-driven predictive modeling for therapeutic intervention [6]. In this study, we addressed a fundamentally limiting assumption of two-dimensional central slice analysis by extending our modeling framework to full volumetric analysis. We demonstrated technical feasibility and performed initial studies showing enhanced predictive capacity. Our modeling framework represents a general approach whereby models of tumor growth and response can be extended by gradually incorporating additional model complexity and then validating model predictions by directly comparing against data observations. Other extensions to our model, such as biologically-inspired perturbations, may be similarly incorporated and validated. Using our data-driven approach, model perturbations in the setting of response of breast cancer to NAT can be incorporated to more fully realize the goal of model-assisted therapeutic intervention.

Acknowledgments

We thank the National Institutes of Health for funding through NCI K25CA204599, NCI R01CA186193, NCI U01CA174706, NCI U01CA142565, NIBIB R21EB022380, and CPRIT RR00016. T.E.Y. is a CPRIT Scholar in Cancer Research.

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