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RESEARCH ARTICLE

Numerical knockouts–In silico assessment of factors predisposing to thoracic aortic aneurysms

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Abstract

Myriad risk factors-including uncontrolled hypertension, aging, and diverse genetic mutations-contribute to the development and enlargement of thoracic aortic aneurysms. Detailed analyses of clinical data and longitudinal studies of murine models continue to provide insight into the natural history of these potentially lethal conditions. Yet, because of the coexistence of multiple risk factors in most cases, it has been difficult to isolate individual effects of the many different factors or to understand how they act in combination. In this paper, we use a data-informed computational model of the initiation and progression of thoracic aortic aneurysms to contrast key predisposing risk factors both in isolation and in combination; these factors include localized losses of elastic fiber integrity, aberrant collagen remodeling, reduced smooth muscle contractility, and dysfunctional mechanosensing or mechanoregulation of extracellular matrix along with superimposed hypertension and aortic aging. In most cases, mild-to-severe localized losses in cellular function or matrix integrity give rise to varying degrees of local dilatations of the thoracic aorta, with enlargement typically exacerbated in cases wherein predisposing risk factors co-exist. The simulations suggest, for the first time, that effects of compromised smooth muscle contractility are more important in terms of dysfunctional mechanosensing and mechanoregulation of matrix than in vessel-level control of diameter and, furthermore, that dysfunctional mechanobiological control can yield lesions comparable to those in cases of compromised elastic fiber integrity. Particularly concerning, therefore, is that loss of constituents such as fibrillin-1, as in Marfan syndrome, can compromise both elastic fiber integrity and mechanosensing.

Author summary

Aneurysms are local dilatations of the arterial wall that are responsible for significant disability and death. Detailed analyses of clinical data continue to provide insight into the natural history of these potentially lethal conditions, with myriad risk factors–including uncontrolled hypertension, aging, and diverse genetic mutations–contributing to their development and enlargement. Yet, because of the co-existence of these risk factors in most cases, it has been difficult to isolate individual effects or to understand how they act in combination. In this paper, we use a computational model of the initiation and progression of thoracic aortic aneurysms to contrast key predisposing factors both in isolation and in combination as well as with superimposed hypertension and aging. The present study recovers many findings from mouse models but with new and important observations that promise to guide in vivo and ex vivo studies as we seek to understand and eventually better treat these complex, multi-factorial lesions, with data-informed patientspecific computations eventually the way forward.

Introduction

Thoracic aortic aneurysms (TAAs) are local dilatations of the proximal aorta that often have an increased risk of subsequent dissection and/or rupture. Clinical experience reveals that uncontrolled hypertension and natural aortic aging are key risk factors for the initiation and progression of these lesions [1,2], though many TAAs arise from diverse genetic mutations [3– 5]. Specifically, predisposing mutations affect genes that encode extracellular matrix (e.g., *FBN1*, which encodes fibrillin-1, or *EFEMP2*, which encodes fibulin-4, both of which affect elastic fibers), cytoskeletal proteins (e.g., *ACTA2*, which encodes smooth muscle α -actin, and *MYH11*, which encodes smooth muscle myosin heavy chain), and the cytokine transforming growth factor-beta (*TGFB2*) or its signaling through transmembrane receptors (*TGFBR1,2*, which encode type 1 and 2 TGF β receptors) or intracellular second messengers (e.g., *SMAD2*, which encodes a key downstream signaling molecule SMAD2), which affect collagen turnover among many other biological processes. There is, in addition, increasing evidence that detrimental alterations to either extracellular matrix or intracellular contractile proteins implicate a dysfunctional mechanotransduction axis in the smooth muscle cells as causative of certain TAAs [5–8].

Advances in genetics have provided critical insight into TAAs based on human data, though the complexity of the disease process continues to stimulate countless studies in mice, particularly those having mutations to genes that have been found in humans, including *Fbn1*, *Acta2*, *Myh11*, *Tgfbr1*, and so forth. Mouse models enable the collection of detailed longitudinal data on the molecular and cellular biology, histopathology, biomechanical properties, and in vivo cardiovascular function. Yet, these mouse models are similarly complex, with disease presentation differing with the wild-type background [9], manifesting at different times and to different extents within the same genetic model [10], and in some cases seemingly manifesting differently in mice and humans (cf. [11,12]). Indeed, genetic analyses reveal hundreds of differentially expressed genes even in the case of a targeted mutation to a single gene (e.g., [13,14]), suggesting that additional collateral or compensatory changes can complicate interpretation further.

There is, therefore, a pressing need to synthesize information from different types of studies, including cellular [15] and murine and clinical [16]. It has long been appreciated that the biomechanics is fundamental to understanding these complex lesions [17–19], with continuing suggestions that "biomechanical analyses should be an integral part of aortic disease characterization under different genetic lesions and pharmacological interventions" [2]. We submit, therefore, that data-informed computational models, including systems biological and biomechanical, can contribute to our understanding by synthesizing empirical information and generating and testing hypotheses (e.g., [20,21]). In this paper, we build on a well-established theoretical framework for describing arterial growth (changes in composition) and remodeling (changes in organization) that allows one to predict evolving geometry, microstructure, and mechanical properties. Specifically, we present computational simulations of the development of local dilatations of the murine thoracic aorta resulting from prescribed changes in many of the key risk factors that give rise to TAAs, including relative effects of local losses of elastic fiber integrity, aberrant collagen structure, reduced smooth muscle contractility, and dysfunctional mechanosensing or mechanoregulation of matrix, all of which have been implicated as causative in human and murine studies. In addition, we consider the effects of hypertension and aortic aging, both in isolation and when superimposed on local losses in matrix integrity or cell functionality. An advantage of in silico approaches is that one can study each of the implicated factors in isolation and then in combination. Results reveal that some local insults can have marked effects alone, but in most cases the combination of multiple insults or risk factors is particularly concerning.

Methods

Theoretical framework

The normal thoracic aorta consists of myriad constituents that collectively endow the wall with appropriate resilience and stiffness, which provide biomechanical functionality, and yet sufficient strength, which protects against mechanical failures such as dissection and rupture. Importantly, these constituents can turnover and allow the aorta to change its caliber, wall thickness, and composition, and thus biomechanical properties, to adapt to diverse changes in hemodynamic loading, as in consistent exercise or conversely hypertension [22,23]. In contrast, adverse changes occur in natural aging and most disease processes, which can result from phenotypic modulation of the cells (as in progressive endothelial dysfunction) and changes in matrix composition and organization (as in fibrosis). Hence, a computational modeling framework must account for different structurally significant constituents that can have different material properties and different rates of turnover.

Over a period of years, we developed a "constrained mixture model" that can describe the evolving composition, geometry, and nonlinear material properties of arteries under many conditions (e.g., [20,24,25]). Briefly, this model requires three different constitutive relations for each of the structurally significant constituents, which for convenience we lump into three groups: elastin-dominated matrix, collagen-dominated matrix, and functional smooth muscle cells. The three relationships describe the rates of mass density production $(m^{\alpha}(\tau)>0, \text{ where } \tau)$ is the time at which constituent $\alpha = 1, 2, \dots, n$ is produced), the removal of these constituents due both to their normal half-lives and accelerated losses in disease ($q^{\alpha}(s,\tau) \in [0,1]$, where *s* is the current time of interest), and the multiaxial mechanical properties (with passive behaviors captured by $\hat{W}^{\alpha}(\mathbf{F}_{n(\tau)}^{\alpha}(s)) > 0$, which denotes the stored elastic energy in constituent α that depends on the multiaxial deformation $\mathbf{F}_{n(\tau)}^{z}(s)$ experienced by that constituent at current time s relative to the configuration at τ at which it was incorporated within the extant matrix) of the individual constituents. In particular, elastin is not produced in maturity, but it can become degraded or damaged; collagen turns over continuously, and functional smooth muscle (with passive and active contributions to load bearing) can increase (via cell proliferation), decrease (via apoptosis), or adapt (remodel).

Importantly, because of the mechanobiological control of matrix by intramural cells [26], the functions for rates of production and removal include (potentially evolving) nominal values ($m_N^{\alpha}(\tau) > 0$ and $k_N^{\alpha}(\tau) > 0$, respectively) as well as modulated changes that are proportional to deviations in mechanical stimuli from homeostatic values, here assumed in terms of a mean pressure- and axial force-induced intramural stress (σ) and flow-induced wall shear stress (τ_w). Specifically, we model production (per unit current volume) by

$$m^{\alpha}(\tau) = m_{N}^{\alpha}(\tau)\Upsilon^{\alpha}(\tau) = m_{N}^{\alpha}(\tau)(1 + K_{\sigma}^{\alpha}\Delta\sigma(\tau) - K_{\tau_{w}}^{\alpha}\Delta\tau_{w}(\tau))$$
(1)

where $K_i^{\alpha} > 0$ are gain-type parameters that dictate the sensitivity of the overall deviations in stress ($\Delta \sigma$ and $\Delta \tau_w$) from homeostatic values (σ_o and τ_{wo}), with $\Delta \sigma = ((1-\delta)\sigma-\sigma_o)/\sigma_o$ where $\delta \in$ [0,1] is a newly introduced parameter to account for perfect mechanosensing ($\delta = 0$) or its dysfunction ($\delta \leq 1$) and $\Delta \tau_w = ((1-\xi)\tau_w-\tau_{wo})/\tau_{wo}$ where $\xi \in [0,1]$ similarly accounts for endothelial health ($\xi = 0$) or its dysfunction ($\xi \leq 1$). These relations, generally collected into stimulus functions Υ^{α} , capture, for example, increases in pressure-induced wall stress tending to heighten matrix production through increased angiotensin II and TGF β signaling as well as increases in flow-induced wall shear stress tending to attenuate matrix production via increased nitric oxide, which additionally stimulates vasodilatation [24,27]. In the simulations below, we assume that flow remains constant and that endothelial-mediated effects are confined to the non-aneurysmal segments; hence, we do not solve the full fluid-solid-growth problem here. We similarly assume that the distending pressure is constant, at either normotensive or hypertensive systolic levels, not considering pulse pressure due to the current lack of appropriate G&R relations in terms of pulsatility (cf. [28]).

We model removal by

$$q^{\alpha}(s,\tau) = \exp(-\int_{\tau}^{s} k_{N}^{\alpha}(t)dt) = \exp(-\int_{\tau}^{s} k_{o}^{\alpha}(1+k(\Delta\sigma(t))^{2})dt)$$
(2)

where k_N^{α} are rate-type parameters modulating the role of deviations in intramural stress ($\Delta\sigma$) from homeostatic values, with k_o^{α} the original rate, noting that increased wall stress can both increase the production / activation of proteinases and decrease the vulnerability of normal matrix to proteolytic digestion [29,30]. Production and removal of constituent α determine its homogenized mass density per unit reference volume through the convolution integral

$$\rho_R^{\alpha}(s) = \int_{-\infty}^s m_R^{\alpha}(\tau) q^{\alpha}(s,\tau) d\tau$$
(3)

where $m_R^{\alpha} = Jm^{\alpha}$ are associated referential mass density production rates, with $J = \sum \rho_R^{\alpha} / \rho =$ det **F** the volume ratio computed from the deformation gradient **F** from reference to current (in vivo) configurations for the mixture.

Finally, the biomechanical behavior of the aortic wall is given in terms of the Cauchy stress

$$\boldsymbol{\sigma}(s) = -p(s)\mathbf{I} + \frac{2}{J(s)}\mathbf{F}(s)\frac{\partial W_{R}(s)}{\partial \mathbf{C}(s)}\mathbf{F}^{T}(s) + \boldsymbol{\sigma}^{act}(s)$$
(4)

where *p* is a Lagrange multiplier that enforces isochoric motions during transient loading, $\mathbf{C} = \mathbf{F}^T \mathbf{F}$ the right Cauchy-Green tensor, $W_R = \sum W_R^{\alpha}$, that is, the total stored energy per unit reference volume W_R is the sum of the energies stored in the constituent parts (namely, with constituents α representing elastin- and collagen-dominated matrix as well as mechanically functional smooth muscle), and $\boldsymbol{\sigma}^{act}$ is the contractile active stress generated by the smooth muscle cells. For the constituent-specific passive mechanical properties, we let

$$W_{R}^{e}(s) = \phi_{R}^{e}(s)\hat{W}^{e}(s) = \phi_{R}^{e}(s)\frac{c^{e}}{2}(\operatorname{tr}(\mathbf{F}^{eT}(s)\mathbf{F}^{e}(s)) - 3)$$
(5)

for the elastin-dominated matrix (*e*), where $\phi_R^e = J\phi^e$ represents its referential mass fraction within the mixture, $\mathbf{F}^e = \mathbf{F}\mathbf{G}^e$ is the deformation gradient measured from its fixed natural (stress-free) configuration, with \mathbf{G}^e an effective deposition stretch tensor, and c^e is a shear modulus that can be determined from biomechanical testing data. For the energy stored in multiple fiber families, which turnover continuously, we let

$$W_{R}^{c,m}(s) = \frac{1}{\rho} \int_{-\infty}^{s} m_{R}^{c,m}(\tau) q^{c,m}(s,\tau) \hat{W}^{c,m}(\lambda_{n(\tau)}^{c,m}(s)) d\tau$$
(6)

for collagen (*c*) and passive smooth muscle (*m*) dominated behaviors, where ρ is the mass density of the tissue, $m_R^{c,m} = Jm^{c,m}$ are referential mass density production rates, $\lambda_{n(\tau)}^{c,m}(s)$ the fiber stretches relative to the associated evolving natural configurations, including the extent of the respective deposition stretches $G^{c,m}$ (which result directly from active cell mechanoregulation at the time of deposition), and

$$\hat{W}^{c,m}(\lambda_{n(\tau)}^{c,m}(s)) = \frac{c_1^{c,m}}{4c_2^{c,m}} \left(\exp[c_2^{c,m}((\lambda_{n(\tau)}^{c,m}(s))^2 - 1)^2] - 1\right)$$
(7)

with c_j^i material parameters that can be determined from biaxial mechanical tests on excised arteries. Moreover, we let [31]

$$\boldsymbol{\sigma}^{act}(s) = \phi^{m}(s)T(\Delta\tau_{w}(s))\lambda_{\theta,act}(s) \left(1 - \left(\frac{\lambda_{m} - \lambda_{\theta,act}(s)}{\lambda_{m} - \lambda_{o}}\right)^{2}\right)\boldsymbol{e}_{\theta} \otimes \boldsymbol{e}_{\theta}$$
(8)

where ϕ^m is the mass fraction of smooth muscle, $T(\Delta \tau_w) = T_{max}(1-\exp(-(C_B-C_S\Delta \tau_w)^2))$ is a flow induced wall shear stress mediated level of contractility of the circumferentially oriented smooth muscle cells (with C_B and C_S parameters), which exhibit a nonlinear dependence on the active circumferential stretch ($\lambda_{\theta,act}$), with λ_m and λ_o levels of stretch at which contraction is maximal or minimal. The active circumferential stretch $\lambda_{\theta,act}(s) = a(s)/a_{act}(s)$, with a(s) the current luminal radius and $a_{act}(s)$ an active reference length that describes adaptive shifts in vasomotor tone via rearrangement of smooth muscle cells via $a_{act}(s) = \int_{-\infty}^{s} k_{act}a(\bar{\tau})e^{-k_{act}(s-\bar{\tau})}d\bar{\tau}$ (cf. [27,31]) with k_{act} the associated rate parameter. In particular, $a_{act}(0) = a(0)$ (i.e., $\lambda_{\theta,act}(s \gg 0) \rightarrow 1$) when active remodeling is complete.

Model parameters

Recall that contributors to TAAs are thought to include compromised elastic fiber integrity and remodeled collagen, with altered turnover rates, dysfunctional or lost smooth muscle cells, and compromised mechanosensing and mechanoregulation of matrix in addition to the general risk factors of hypertension and aging. For purposes of illustration, we used biaxial biomechanical data from a normal descending thoracic aorta from a wild-type male mouse [25] to parameterize the baseline biomechanical model. Rather than determining the remaining parameters from specific mouse models of TAAs, to facilitate comparisons across the many different cases we varied parameters over reasonable ranges to model the different disease conditions of interest.

TAAs frequently present as asymmetric lesions and the involved segment need not be of uniform thickness or luminal diameter. Yet, to facilitate comparisons across cases and to increase insight into vessel-level consequences of the different insults, herein modeled constitutively, consider an initially straight cylindrical segment of the murine thoracic aorta of uniform in vivo wall thickness ($h_o = 40 \ \mu m$) and luminal radius ($a_o = 647 \ \mu m$) into which an initial insult can be placed within the central region, to avoid further complications due to end effects, either asymmetrically or axisymmetrically. Indeed, we focused primarily on axisymmetric cases, which enable the most consistent comparisons, with the evolving axisymmetric insult prescribed by

$$\vartheta(z_o) = \vartheta_{end} + (\vartheta_{central} - \vartheta_{end}) \exp\left(-\left|\frac{z_o - l_o/2}{z_{od}}\right|^{v_z}\right)$$
(9)

where $z_o \in [0, l_o]$, with $l_o = 15$ mm, is the axial coordinate in the reference (homeostatic) configuration, $v_z = 5$ and $z_{od} = 3$ mm are respective (baseline) axial exponential decay and deviation parameters, and ϑ_{end} and $\vartheta_{central}$ are values of the associated controlling parameter near the ends $(z_o = 0, l_o)$ or within the central $(z_o = l_o/2)$ regions of the computational aorta. For example, the axial profile for $\vartheta = K_{\tau}^{\alpha} / K_{\alpha}^{\alpha}$, used in Eq (1) for all of the non-uniform simulations, is shown in Fig A in S1 Supporting Information. Values of key model parameters are listed in Table 1 (and associated caption) for both normal and diseased vessels. Based on preliminary finite element simulations, where different model parameters where altered gradually, we defined (Fig B in S1 Supporting Information), for the axisymmetric lesions, an ~30% loss of elastic fiber integrity as mild and an ~60% loss as severe, both captured by reductions in the value of the elastin-associated material parameter c^{e} (which is computationally equivalent to reducing the local mass fraction of elastin, see Eq (5)); this reduction of the value of c^e is comparable to that found via biaxial tests in the *Fbn1*^{mgR/mgR} model of Marfan syndrome [32]. Likewise, we defined mild-to-severe losses of collagen cross-linking by ~10 to 25% reductions in the material parameter c_1^c . Effects of increased collagen degradation relative to collagen production imply an increased rate k^c ; we used a decreased growth-feedback parameter $k^c K_i^c$ [33] or, equivalently, an increased ratio of the rate of smooth muscle-to-collagen turnover $\eta = (k^m K_i^m)/(k^c K_i^c)$ in our computations. Based on our preliminary study (e.g., Fig B in S1 Supporting Information), we defined mild-to-severe increases in collagen degradation relative to collagen production by 5 to 12.5% increments in the parameter η . We also defined mild-to-complete losses of contractile capacity by letting the value of the active stress parameter T_{max} be reduced by 60 to 100%. These preliminary simulations also revealed that the simulated growth and remodeling (G&R) responses were very sensitive to the parameter δ , which controls that percentage of the intramural stress that is mechano-sensed. Hence, we let mild-to-severe degrees of compromised mechanosensing be defined by $\delta = 0.075$ to $\delta =$ 0.185. Compromised mechanoregulation of matrix was modeled by 0.4 to 1.2 percent reductions in the deposition stretch G^c , another highly sensitive parameter, which relates to collagen undulation at the time of deposition.

Table 1. Representative values of parameters for the baseline model, a non-aneurysmal mouse descending thoracic aorta, adapted (homogenized through the thickness) from [25].

Inner radius, thickness, length	a_o, h_o, l_o	0.647mm, 0.040mm, 15mm
Constituent mass fractions	$\phi^e_o,\phi^m_o,\phi^c_o$	0.34, 0.33, 0.33
Collagen relative fractions	$eta^ heta,eta^z,eta^d$	0.056, 0.067,0.877
Diagonal collagen orientation	α_{00}	29.9°
Elastic material parameters	$c^e,c_1^m,c_2^m,c_1^c,c_2^c$	89.71kPa, 261.4kPa, 0.24, 234.9kPa, 4.08
Deposition stretches	$G^e_ heta, G^e_z, G^e_r, G^m, G^c$	$1.90, 1.62, 1/(G^e_ heta G^e_z), 1.20, 1.25$
Combined production-removal	$\eta = \mathrm{K}^m_i / \mathrm{K}^c_i \cdot k^m_o / k^c_o$	1
Shear-to-intramural gain ratio	$\mathrm{K}^m_{ au_w}/\mathrm{K}^m_{\sigma}=\mathrm{K}^c_{ au_w}/\mathrm{K}^c_{\sigma}$	0.35
Dysfunctional mechanosensing	δ,ξ	0,0
Vasoactive parameters	$T_{max}, C_B, C_S, \lambda_m, \lambda_0$	250kPa, 0.8326, <i>C</i> _B /2,1.1,0.4

Superscripts *e*, *m*, *c* denote elastin-dominated, load-bearing smooth muscle, and collagen-dominated; superscripts/subscripts *r*, θ , *z*, *d* denote radial, circumferential, axial, and symmetric diagonal directions. Subscript *o* denotes original homeostatic values; subscripts *i* = σ , τ_w denote intramural and wall shear stress related parameters, respectively. For the severely diseased vessels, we let: $c_{central}^{\epsilon} = 34.1$ kPa, $c_{1|central}^{\epsilon} = 181$ kPa, $T_{max|central} = 0$, $\delta_{central} = 0.185$, $G_{central}^{c} = 1.235$, or $\eta_{central} = 1.05$ for the axisymmetric aneurysms (see Eq (9) and Figs 1, 2, and H in <u>S1 Supporting Information</u>), and $c_{apex}^{\epsilon} = 20.2$ kPa, $c_{1|apex}^{\epsilon} = 111$ kPa, $\delta_{apex} = 0.4$, or $G_{apex}^{c} = 1.22$ for the asymmetric ones (see Eq (10) and Fig 4), with $\vartheta_{end} = \vartheta_{baseline}$ for all cases.

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Finally, for completeness, we considered a few asymmetric simulations with the evolving asymmetric insult (i.e., axially- and circumferentially-nonuniform) prescribed by

$$\vartheta(z_o, \theta_o) = \vartheta_{end} + \left(\vartheta_{apex} - \vartheta_{end}\right) \exp\left(-\left|\frac{z_o - l_o/2}{z_{od}}\right|^{v_z}\right) \exp\left(-\left|\frac{\theta_o - \pi}{\theta_{od}}\right|^{v_\theta}\right) \tag{10}$$

where $\theta_o \in [0,2\pi]$ is the circumferential coordinate (azimuth) in the reference configuration, $v_{\theta} = 5$ and $\theta_{od} = \pi/3$ are additional (baseline) circumferential exponential decay and deviation parameters, and ϑ_{apex} is the value of the associated controlling parameter at the apex ($z_o = l_o/2$, $\theta_o = \pi$) of the potentially developed aneurysm.

Computational method

We have previously presented finite element implementations of our constrained mixture model (e.g., [34,35]), but herein we use a fast, efficient 3-D implementation [36] that is based on the underlying assumption that each G&R state is mechanobiologically equilibrated [27], which holds for cases wherein the characteristic timescale of the remodeling process is shorter than the timescale for the driving stimulus, that is, for fully quasi-static G&R [37,38].

Briefly, this rate-independent formulation eliminates dependence on G&R time *s* by letting responses "relax" towards evolved homeostatic states that satisfy mechanical and mechanobiological equilibrium at any time *s* (which, thus, plays the role of an evolution parameter). This allows equilibrium to be enforced in homeostatic grown and remodeled in vivo states for given external loads and boundary conditions without the need to track the past history of deposition and removal (as, e.g., in the heredity integral Eq (6)) or to integrate evolution equations (as, e.g., in finite kinematic growth approaches [39]), hence resembling an elastic computation algorithmically. In particular, mechanobiological equilibrium in Eq (1) leads to (subscript *h* refers to evolved homeostatic variables)

$$\Gamma_h^{\alpha}(\Delta\sigma_h, \Delta\tau_{wh}) = 1 \tag{11}$$

whereby production balances removal (i.e., $m_h^{\alpha} = m_{Nh}^{\alpha} = k_{Nh}^{\alpha} \rho_h^{\alpha}$; [27]). On other hand, Eq (6) reduces to $W_{Rh}^{c,m} = \phi_{Rh}^{c,m} \hat{W}^{c,m} (G_h^{c,m})$, which, in combination with an equilibrated energy for elastin in Eq (5), leads to a rule-of-mixtures relation for the total energy $W_{Rh} = \sum \phi_{Rh}^{\alpha} \hat{W}^{\alpha}$ with evolved constituent mass fractions and deformations. Consistently, the Cauchy stress in Eq (4) also adopts a rule-of-mixtures expression in terms of evolved constituent-specific passive and active stresses

$$\boldsymbol{\sigma}_{h} = \sum_{\alpha}^{e,c,m} \phi_{h}^{\alpha} \hat{\boldsymbol{\sigma}}_{h}^{\alpha} + \phi_{h}^{m} \hat{\boldsymbol{\sigma}}_{h}^{act} - p_{h} \mathbf{I}$$
(12)

with the active contribution in Eq (8), and $\lambda_{\theta,act,h} = 1$, relaxed as [27]

$$\boldsymbol{\sigma}_{h}^{act} = \phi_{h}^{m} \hat{\boldsymbol{\sigma}}_{h}^{act} (\Delta \tau_{wh}) = \phi_{h}^{m} T(\Delta \tau_{wh}) \left(1 - \left(\frac{\lambda_{m} - 1}{\lambda_{m} - \lambda_{o}}\right)^{2} \right) \boldsymbol{e}_{\theta h} \otimes \boldsymbol{e}_{\theta h}$$
(13)

where both ϕ_h^m and $\Delta \tau_{wh}$ are deformation-dependent, and the equilibrated (also deformationdependent) Lagrange multiplier p_h is determined from the scalar stress-like constraint Eq (11) during the quasi-static G&R evolution. An exact linearization of the formulation consistent with these constraints enables implementation within a finite element framework, where simultaneous solution of mechanical and mechanobiological equilibrium can be ensured efficiently at load steps that capture evolving geometries, compositions, and properties of interest for complex boundary value problems; see [36] for specific details.

Even if the present model considers each G&R state as mechanobiologically equilibrated, with degradation balanced by production, it still inherits useful ratios among relevant parameters from the full model, as, for example, those that control the rates of turnover of the individual constituents through the smooth muscle-to-collagen turnover ratio $\eta = \eta_a \eta_{\rm Y} = (k^m/k^c) \times$ (K_i^m/K_i^c) mentioned above, whereby $\rho_{Rh}^m/\rho_o^m = (\rho_{Rh}^c/\rho_o^c)^{\eta}$. Moreover, note that the deviation in shear stress from its homeostatic value $\Delta \tau_{wh} = (1-\xi)\tau_{wh}/\tau_{wo}-1$ in Eqs (11) and (13) must be assessed pointwise. We assume the analytical expression $\tau_{wh}/\tau_{wa} = Q_h a_o^3/(Q_o a_h^3)$, with a luminal radius and Q blood flow rate, whose value and derivatives may be approximated and computed at Gauss points in terms of local circumferential and axial stretches $\lambda_{\theta h}$ and λ_{zh} from reference to current homeostatic configurations (for cylindrical segments; see Appendix C in [36]), which, in turn, enables additional straightforward computations of Eq (13) and its consistent tangent from the equilibrated second Piola-Kirchhoff active stresses $\mathbf{S}_{h}^{act} = J_{h} \phi_{h}^{m} \mathbf{U}_{h}^{-1} \hat{\boldsymbol{\sigma}}_{Nh}^{act} \mathbf{U}_{h}^{-1}$ (cf. Eq (40) in [36]). For the intramural stress deviation in Eq (11), we consider $\sigma_h \equiv \sigma_{vh} = \frac{1}{3} \operatorname{tr}(\boldsymbol{\sigma}_h)$ as a stimulus, which has repeatedly yielded numerical responses that reflect experimental observations well [25,36,40]. Moreover, for the sake of clarity in our prior presentation of this rateindependent computational framework [36], we did not posit specific evolution laws for the magnitude or orientation of deposition pre-stretches or -stresses; rather, the rotated (i.e., "right") deposition stretch tensors $G_{Nh}^{c,m}$, and hence associated pre-stresses at the constituent level $\hat{\sigma}_{Nh}^{e,m}$, were prescribed during the evolution, which were particularly critical to control the anisotropic enlargement of the simulated aneurysms. Following previous studies on aneurysms (e.g., [34,35]), here we let the (referential) deposition angle α_{0h} of newly synthesized collagen fibers for the two symmetric diagonal families to evolve during G&R according to

$$\tan \alpha_{0h} = \frac{\lambda_{\theta h}}{\lambda_{zh}} \tan \alpha_{0o} \tag{14}$$

with α_{0o} given in <u>Table 1</u>. Rotated Cauchy pre-stress tensors $\hat{\sigma}_{Nh}^{c,m}(\alpha_{0h})$ for the diagonal fiber families reorient accordingly and require additional tangent contributions in terms of $\alpha_{0h}(\lambda_{\theta h\nu}, \lambda_{zh})$ and its derivatives, see <u>S1 Appendix</u>. In this way, the present implementation extends the computational framework in [36] by considering an active stress for smooth muscle cells and reorientation of diagonal collagen fibers, both computed locally under mechanobiological equilibrium conditions.

Finally, many biomechanical metrics provide important insight into the state of aortic health or disease, including biaxial (circumferential and axial) wall stretch, stress, and material stiffness as well as elastic energy storage. Of these, elastic energy storage is a key indicator of mechanical functionality [41,42] while the value of circumferential material stiffness appears to be a particularly important indicator of aneurysmal propensity or presence [32,43]. Neither quantity can be measured directly; they are best computed from an appropriate nonlinear constitutive model of the aortic wall, modeled here with a 3-D finite element geometry. The finite element model for the initially straight cylindrical segment of the thoracic aorta comprised $N_r N_{\theta} N_{\tau} = 1 \times 20 \times 20 = 400$ displacement-based 3-D quadratic elements with full $3 \times 3 \times 3$ Gauss integration, which showed consistent results with other quadratic meshes analyzed (Fig C in S1 Supporting Information) as well as with a denser linear mesh in [36] while performing faster. Each finite element simulation was computed in a modified open source finite element code FEBio and advanced quasi-statically in 10 incremental steps, which showed quadratic rates of convergence during global Newton-Raphson iterations and elapsed total CPU times ~1 min on a single CPU processor at 3GHz in a Workstation Dell Precision 5810 with 32GB RAM. For all the simulations, axial displacements were fixed at the proximal and distal

ends of the computational thoracic aorta, and cardiac output was considered constant (i.e., $Q_h/Q_o = 1$).

Results

Hypertension and aging

Hypertension can arise via many causes and manifest as diverse vessel level changes, often associating with an increased structural stiffening of the central arteries, which in turn can feedback through the hemodynamics to increase pulse pressure further [44,45]. In particular, elevated blood pressure tends to stiffen the aorta structurally due primarily to an increased accumulation of fibrillar collagen [46]. Fig D in S1 Supporting Information shows simulated effects of a uniform elevation of systolic blood pressure of 1.5-fold on our baseline model of the murine thoracic aorta, with remodeling resulting in decreased elastic energy storage as well as a modestly increased circumferential material stiffness with an increased structural stiffness due to wall thickening. Vascular aging is similarly a complicated biological process, manifesting via many vessel level changes. In humans, a conspicuous change is diffuse elastic fiber fragmentation with an associated modest uniform dilatation and stiffening of the wall [47-49]. Fig D in S1 Supporting Information also shows simulated effects of an aging-related uniform loss of 30% of the elastic fibers relative to our baseline aortic model, which reflects well the general empirical observations, including a (irreversible) reduction in elastic energy storage [50]. Finally, the bottom panels in Fig D in S1 Supporting Information show aortic remodeling in response to a sustained elevation of pressure combined with such aging. Importantly, note that the aortic remodeling is diffuse, not localized, in all three of these cases. Effects of hypertension and aging on aortic remodeling are considered further below as possibly aggregating risk factors when combined with any of the five local factors that associate with TAAs and are contrasted herein.

Elastic fiber integrity

Competent elastic fibers endow the aortic wall with an ability to store energy elastically during systole, which can then be used to work on the blood during diastole to augment flow, as, for example, to perfuse the coronary arteries of the left heart. Most histopathological studies reveal local fragmentation or loss of elastic fibers in human and murine TAAs (e.g., [14,51]). Figs 1 and 2 (first rows) show predicted results for locally compromised elastic fiber integrity both without and with the added risk factors of hypertension and aortic aging-these figures show the mechanically-loaded geometry after the lesion has developed, with colorimetric scaling for elastic energy storage (Fig 1) and circumferential material stiffness (Fig 2). As it can be seen, graded localized losses of elastic fiber integrity of ~30% and 60% can result in G&R responses that cause minor-to-marked localized dilatations of the aortic wall (cf. Fig B in S1 Supporting Information). Importantly, when either hypertension (a sustained increase in blood pressure, in this case a 50% elevation above baseline systolic) or aortic aging (arbitrarily taken as a uniform 30% loss in elastin) is superimposed on even the case of a mild localized loss of elastic fiber integrity, the otherwise unremarkable local dilatation increases dramatically. That is, hypertension and aortic aging can dramatically exacerbate the consequences of this underlying initiator of the localized dilatation. Note, too, that the maximal decrease in elastic energy storage capability and the maximal increase in circumferential material stiffness occur in the central, most dilated, region, with energy storage decreasing about 30% and stiffness increasing approximately 2-fold. These predicted changes are consistent with measured changes in the non-aneurysmal descending thoracic aorta in Fbn1mgR/mgR mice, which have compromised elastic fiber integrity [32], though there was no attempt here to refine the model parameters to

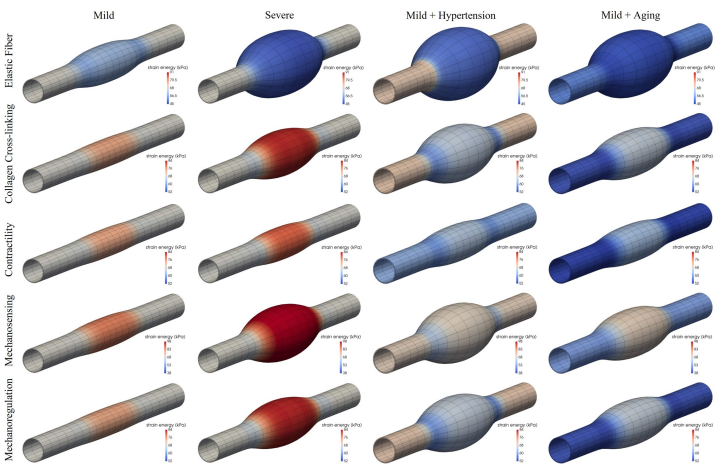


Fig 1. Fully developed, mechanobiologically equilibrated, axisymmetric dilatations of an initially cylindrical aortic segment caused by a mild (first column) or severe (second column) localized (via the respective parameter $\vartheta_{central}$ in Eq (9)) loss of elastic fiber integrity (first row, parameter c^e , mild 32% and severe 62% reduction), decrease in collagen cross-linking (second row, parameter c_1^e , mild 9%, severe 23% reduction), loss of smooth muscle contractility (third row, parameter T_{max} , mild 60%, severe 100% reduction), compromised mechanosensing (fourth row, parameter δ , mild 7.5%, severe 18.5%, absolute), or compromised mechanoregulation (fifth row, parameter G^c , mild 0.4%, severe 1.2%, relative change), with the risk factors of hypertension (third column, 50% increase in blood pressure) and aortic aging (fourth column, uniform 30% loss of elastic fiber integrity) superimposed on the mild insults. Shown are color maps for elastic energy storage per unit current volume for the mechanically-loaded geometries. Note that the original in vivo homeostatic value $W_o = 68$ kPa (light grey, centered mark). Both $K_{\tau_w}^x/K_{\sigma}^x|_{end} > 0$ and $\eta_{end} < 1$ were adjusted via Eq (9) to maintain the distal and proximal segments normal (cf. Figs A and F in S1 Supporting Information).

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fit any particular data set. That the maximal changes co-localized with the region of maximal dilatation is also consistent with experimental (passive) measurements [43].

Finally, note that the loss of elastic fibers tends to associate with reduced vasoconstrictive capability [52], hence we did not initially include effects of smooth muscle contractility (i.e., in results displayed in Figs 1 and 2, first rows). Simulations in Fig E in S1 Supporting Information (left) show a potentially protective role of smooth muscle contraction ($T_{max} = 50$ kPa; see Methods) when combined with a severe localized loss of elastic fiber integrity (~60%). Importantly, however, such an active resistance to dilatation (which would also be mediated by changes in the hemodynamics as the aorta enlarges) is limited, being overcome by an additional 10% localized degradation of elastin (right), thus leading to dilatations similar to those in Fig 1, although with further adverse changes in elastic energy storage and circumferential stiffness. Because all of these simulations depend on particular choices of multiple G&R model parameters, Fig F in S1 Supporting Information (first row) exams potential consequences of different values for the simpler case of a uniform mild (30%) loss of the elastic fiber integrity, which

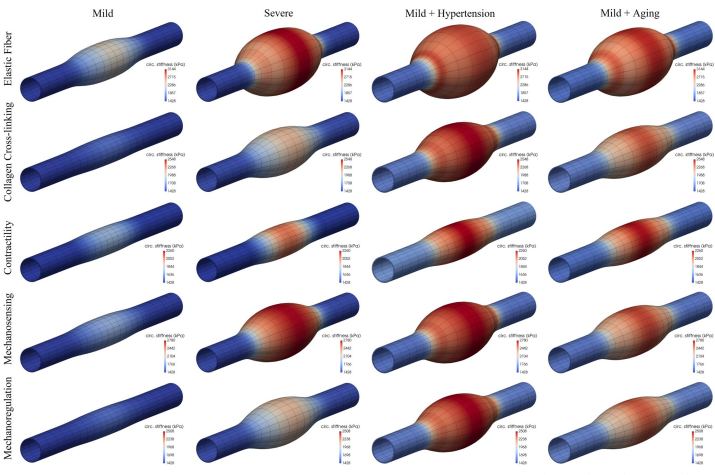


Fig 2. Same as Fig 1 except color maps for circumferential material stiffness for the mechanically-loaded geometries. Note the original in vivo homeostatic value $c_{\theta\theta\theta\thetao} = 1428$ kPa (dark blue, bottom mark) as well as an increased stiffness at the shoulder of some of the lesions resulting from a locally increased α_{0h} (i.e., higher $\lambda_{\theta h} / \lambda_{zh}$, see Eq (14)).

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emphasizes further that (marked) aneurysmal dilatation results from (severe) localized, not uniform, losses in elastic fiber integrity, with various G&R factors simply modulating the specific extent of the response. Likewise, Fig G in S1 Supporting Information (first row) explores effects of size and spatial transition (from normal to diseased regions) of the axisymmetric lesion considered, which can be controlled with the decay (v_z) and deviation (z_{od}) parameters in the Gaussian-like exponential distribution in Eq (9). Perhaps expectedly, for fixed minimal ($c_{e|end}$) and maximal ($c_{e|central}$) levels of compromised elastic fiber integrity, longer aneurysms develop for prescribed lesions that are initiated by greater axial insults (i.e., greater z_{od} for fixed v_z). Conversely, a notably reduced overall dilatation develops for a more gradual decay from maximally to minimally injured locations (i.e., lower v_z for fixed z_{od}). Similar conceptual predictions are obtained for the other insults considered below (second to fourth rows), although with different degrees of maximal aneurysmal dilatation.

Collagen remodeling

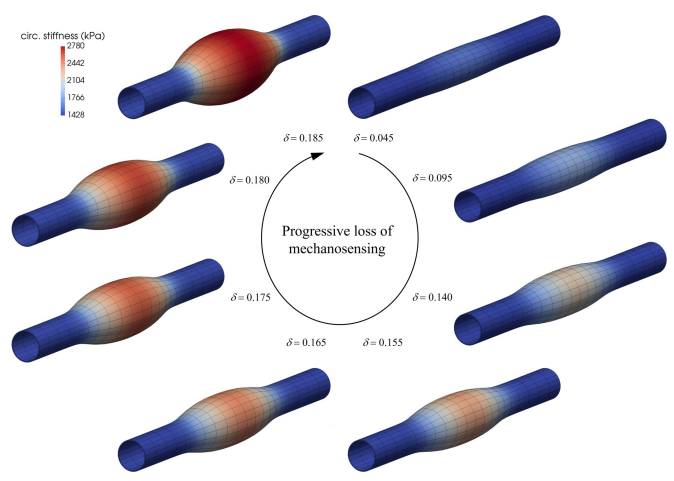
Fibrillar collagens endow the aortic wall with stiffness and strength; they have a normal halflife on the order of months but can turnover more quickly to adapt the wall to changing mechanical loads. Such remodeling can also be maladaptive, however, resulting from or contributing to aneurysmal disease progression [53,54]. Collagen remodeling can manifest in many ways, including differences in cross-linking and rates of degradation. Figs 1 and 2 (second rows) show predicted effects of localized decreases in collagen cross-linking, both in isolation and in combination with hypertension or aging. As it can be seen, localized changes in collagen integrity also result in localized dilatations of the aortic wall, either modest or marked depending on the initiating insult, with all dilatations again exacerbated by hypertension or aging. Proteinase activity is increased in many TAAs [55], hence we similarly considered increased rates of collagen degradation relative to collagen production (via the parameter $\eta = (k^m K_i^m)/(k^c K_i^c)$; see Methods). Fig H in S1 Supporting Information shows predictions for a locally increased ratio η (Table 1) combined with either hypertension or aging, which again results in localized dilatations, particularly when both risk factors are superimposed. Importantly, complementing a prior finding that effects of flow-induced shear stress-mediated matrix turnover contribute to the biaxial G&R near the non-aneurysmal end regions [36], the parameter $\eta > 0$ needed to be adjusted in the non-aneurysmal ends for all simulations to maintain the adjacent non-aneurysmal regions homeostatic (particularly those with uniform effects of aging). Indeed, a uniform moderate decrease of 20% in collagen cross-linking leads to minor-to-modest dilatations and stiffening depending on specific values of these G&R parameters, with a superimposed contractile capacity again proving to be potentially protective (Fig F in <u>S1 Supporting Information</u>, second row).

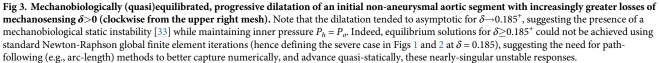
Reduced smooth muscle contractility

Vascular caliber is controlled via multiple mechanisms, both global (central nervous system) and local, with the latter depending on endothelial cells sensing the local wall shear stress and secreting vasodilators or vasoconstrictors that induce smooth muscle relaxation or contraction [56,57]. Reduced smooth muscle contractility thus causes the wall to dilate, transferring loadbearing to the extracellular matrix [58]. Although elastic arteries such as the aorta are thought to have low basal tone compared with muscular arteries, the murine aorta has a significant contractile range. Figs 1 and 2 (third rows) show predicted consequences of mild and severe (total) localized losses of vessel-level smooth muscle tone in our model of the thoracic aorta. As it can be seen, overall effects on central geometry and material properties are relatively modest. Shown, too, is the predicted consequence of a localized loss of smooth muscle contractility with superimposed hypertension or aging. Perhaps surprisingly, overall effects on the geometry and properties are again modest. Albeit not shown, only when both hypertension and aging were superimposed on the case of a mild localized loss of smooth muscle contractility did the aortic dilatation and associated changes in biomechanical properties become dramatic. Aging is often accompanied by hypertension, hence emphasizing the need to consider many factors in combination. Finally, Fig F in S1 Supporting Information (third column) shows different responses given a uniform loss of 80% of smooth muscle tone combined with different G&R parameters, which again yields the most modest changes in geometry and stiffness among all of the considered insults.

Dysfunctional mechanosensing

Intramural cells sense their local mechanical environment, via integrins and competent actomyosin activity, by probing the extracellular matrix that carries most of the hemodynamically imposed load [59,60]. Such mechanosensing of matrix is thus critical to mechanical homeostasis of the aorta. Figs 1 and 2 (fourth rows) show predictions for mild and modest degrees of compromised mechanosensing, which importantly results in localized dilatations of the aortic wall that are exacerbated by hypertension and, to a lesser degree, aortic aging. Again, note that





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the maximal changes in elastic energy storage and circumferential material stiffness manifest within the most dilated region, with quantitative changes similar to those for compromised elastic fiber integrity (cf. first rows). Fig I in <u>S1 Supporting Information</u> shows predicted changes in additional biomechanical metrics–biaxial wall stress and material stiffness as well as local changes in volume ratio–due to the associated localized dilatations, again revealing marked changes. Fig 3 further shows a possible progressive (quasi-equilibrated mechanobiologically) dilatation with increasingly greater losses of mechanosensing. Finally, Fig F in <u>S1 Supporting Information</u> (fourth row) shows different responses given a uniform moderate (12%) loss of mechanosensing combined with different G&R and contractile parameters, with minor-to-modest dilatations and stiffening similar to predictions for a mild loss of elastic fiber integrity (first row).

Dysfunctional mechanoregulation

Functional intramural cells not only degrade old structural constituents and deposit new constituents within the extant matrix, they work on and fashion these new constituents so as to ensure preferred properties under conditions of interest [61,62]. Mechanoregulation of collagen can be captured, in part, via values of the deposition stretches (or, pre-stresses) that are built into the new constituents at the time of matrix production. Figs 1 and 2 (fifth rows) show predicted effects of localized changes in the deposition stretch of the multiple families (i.e., orientations) of fibrillar collagen. Clearly, poorly mechanoregulated collagen within the central region can result in modest-to-marked localized dilatations of the aortic wall, which are again exacerbated by hypertension or aging. Consistent with the stress derived from Eq (7), decreased collagen cross-linking and dysfunctional mechanoregulation, as modeled here, affect the mechanical response of collagen similarly, hence leading to similar computational results when prescribed locally (cf. second rows), but also uniformly (Fig F in <u>S1 Supporting</u> Information, second and fifth rows). Note that it has been shown previously that values of the deposition stretch must be greater than unity to enable potentially optimal tissue maintenance or remodeling [24,63].

Evolving stress-stretch relations

Biomechanical metrics such as material stiffness, energy storage, and contractility are fundamental descriptors of a ortic properties and function [41,64], yet many people intuit changes in stress and stretch more easily. Hence, equibiaxial stress-stretch tests at the apices of the respective aneurysms were simulated, with material properties and mass fractions extracted in postprocessing from our 3-D computational formulation. Fig J in S1 Supporting Information shows (passive) material behaviors after the respective lesions have arisen from marked losses of elastin fiber integrity (recall Figs 1 and 2, first rows), reductions in collagen cross-linking (second rows), decreased vessel-level smooth muscle contractility (third rows), dysfunctional mechanosensing (fourth rows), and dysfunctional mechanoregulation (fifth rows). Comparisons with the initial non-aneurysmal behaviors show a general loss of distensibility and extensibility in all five cases, consistent with the computed increase in material stiffness, even for those insults that do not alter directly the initial (passive) material properties, but rather the mass fractions (such as loss of contractility and dysfunctional mechanosensing). Note that the largest change in material behavior occurs in the circumferential and axial directions for the case of compromised elastic fiber integrity, which shows an initially compliant behavior that stiffens within the region of physiological stresses (~200 kPa). Consistent with the growth and remodeling observed in Figs 1 and 2, the smallest change in passive material behavior occurred for the loss of vessel-level smooth muscle contractility. Effects of reduced collagen cross-linking and compromised mechanosensing and mechanoregulation of matrix yield stiffer responses, although with differently evolved anisotropy. Importantly, the present results, which focus on differences in remodeled properties for aneurysms that develop from a common non-aneurysmal aorta, complement but extend significantly those in [35], which revealed a clear dependency of aneurysmal enlargement on initially different non-aneurysmal aortic properties, hence highlighting the complexity of the disease process.

Asymmetric dilatations

Although axisymmetric simulations are both computationally less expensive (e.g., by allowing one to focus on a single longitudinal section) and easier to interpret, most TAAs are asymmetric. Hence, we considered one class of asymmetric losses of cell or matrix function (Fig 4). Interestingly, findings were similar to those for the axisymmetric case although the degree of the more localized insult needed to be greater to elicit similar localized dilatations. In other words, cases of complete circumferential involvement of the initiator of localized dilatation can give rise to marked dilatations even if the degree of the insult is mild. When truly localized,

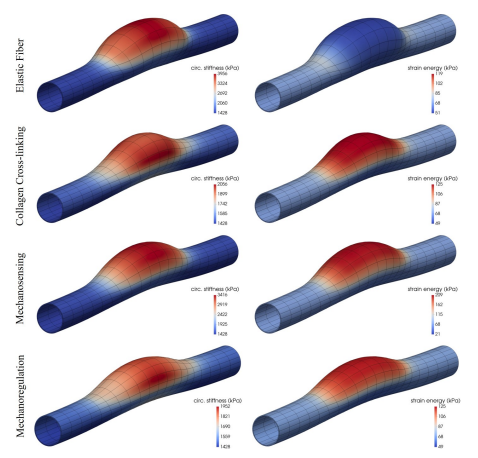


Fig 4. Fully developed, mechanobiologically equilibrated, asymmetric dilatations of an initially cylindrical aortic segment caused by marked locally compromised elastic fiber integrity (first row, 78% decrease in c^e), collagen cross-linking (second row, 53% decrease in c_1^e), mechanosensing (third row, $\delta = 0.4$), or mechanoregulation (fourth row, 2.3% decrease in G^c) via ϑ_{apex} in Eq (10). Shown are color maps for circumferential material stiffness (left) and elastic energy storage per unit current volume (right) for the mechanically-loaded geometries. Note the original in vivo homeostatic values $c_{\theta\theta\theta\theta\sigma} = 1428$ kPa (dark blue, bottom mark) and $W_o = 68$ kPa (light blue, second-from-bottom mark). Both $K_{\tau_w}^z/K_{\sigma}^z|_{end} > 0$ and $\eta_{end} < 1$ were adjusted via Eq (9) to maintain the distal and proximal segments normal (cf. Figs A and F in S1 Supporting Information).

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axially and circumferentially, the surrounding normal tissue appears to lessen or slow the severity of the response. Indeed, Fig K in S1 Supporting Information shows effects of the length and width of the initiating axisymmetric insults considered, controlled in these cases with deviation parameters z_{od} and θ_{od} in the bidirectional Gaussian-like exponential distribution in Eq (10). Similar to the analysis in Fig G in S1 Supporting Information, for fixed minimal (ϑ_{end}) and maximal (ϑ_{apex}) levels of compromised parameters, longer and/or wider asymmetric aneurysms develop for prescribed insults that expand axially and/or circumferentially (i.e., greater z_{od} and/or θ_{od} for fixed v_z and v_{θ}), with damage increased along the cross-sections generally leading to a substantial worsening of the aneurysms. Albeit not shown, the computational model predicted that diagonal fibers of collagen within the damaged area of these asymmetric aneurysms reoriented toward the circumferential direction, via Eq (14), consistent with a prior parametric analysis [36] that showed that asymmetric axial expansions of the central region, hence larger out-of-plane deformations of the aortic centerline associated with increasing realignments toward the circumferential direction (as prescribed therein).

Discussion

Uncontrolled hypertension is a well-known risk factor for thoracic aortopathies and antihypertensive medications are often prescribed [2]. Hypertension increases the pressure-load on the aortic wall, which increases wall stress and the potential for mechanical failure, namely dissection or rupture. In addition, however, increased pressure-loading elicits mechanosensitive responses in normal cells that initially attempt to adapt the wall to offset the increased load, typically via the deposition of additional fibrillar collagen among other changes in composition. This hypertensive response is complex, however, often accompanied by endothelial dysfunction, smooth muscle cell phenotypic modulation, and inflammatory cell infiltration as well as global effects on the central nervous and renin-angiotensin systems [44,65]. Our model of hypertension focused on the initial mechano-adaptive response that seeks to return wall stress toward normal via an increased deposition mainly of collagen, which thickens the wall [25]. Vascular aging is similarly a well-known risk factor for many cardiovascular diseases, including TAAs [1]. Again, the effects of aging on the vasculature are manifold, including endothelial dysfunction, smooth muscle phenotypic modulation or apoptosis, extracellular matrix remodeling, and inflammation [66,67]. Our model of aging focused on the diffuse loss of elastic fiber integrity, as in human though not natural murine aging [50]. Consistent with observations in clinical and animal model studies, our in silico simulations yielded an aortic wall that remodeled in hypertension primarily via an increase in wall thickness and in aging primarily via an increase in caliber, both uniformly and both consistent with an initial mechano-adaptative response (i.e., in the absence of intramural cell dysfunction or inflammation). There were two key observations. First, neither hypertension nor aging alone resulted in localized dilatations; rather their effects were uniform in our idealized model. Second, both of these factors exacerbated each of the five different initiators of localized dilatation, which in many cases (with $\approx 170\%$ and $\approx 100\%$ maximal increases in diameter locally for axisymmetric and asymmetric lesions, respectively) reached the clinical definition of an aneurysm (>50% increase). Our findings are thus consistent with these two conditions being important risk factors for aneurysmal progression, though not aneurysmal initiation consistent with clinical experience of millions of individuals who are hypertensive or of advanced age and yet the much smaller numbers that present with TAAs.

Competent elastic fibers endow the aortic wall with an ability to resist, indeed to recoil against, distending pressures, which helps to control luminal diameter. We considered localized losses of elastic fiber integrity as arise in Marfan syndrome. Studies in the Fbn1^{mgR/mgR} mouse model of Marfan syndrome reveal compromised elastic fiber integrity along the entirety of the aorta, though the greatest losses manifest in the aortic root/ascending aorta where aneurysms first develop [68]. It has been suggested that the different embryonic lineage of smooth muscle cells in this region (from the second heart field and neural crest) and the different state of mechanical loading (biaxial loading and complex hemodynamics) contribute to the predisposition of the proximal aorta to an [69-71]. We did not attempt to model propensity due to differential embryonic lineage or site-specific loading. Rather, we focused on the more general question as to whether a localized loss of elastic fiber integrity can result in aneurysmal dilatation based solely on a mechanobiological response to the insult. Consistent with prior simulations that mimic human abdominal aortic aneurysmal enlargement due to localized losses in elastin [35], we found for the murine thoracic aorta that localized losses of elastic fiber integrity give rise to local dilatations under normotensive conditions, with the initial extent of the insult important in dictating the degree of dilatation over the period studied. Moreover, superimposed hypertension (i.e., uniform increase in pressure) always exacerbated the dilatation, consistent with findings in patients [72], whereas superimposed aging (i.e.,

additional uniform loss of elastin) had differing effects depending on the degree of the initial insult, which appears to be consistent with findings in Marfan patients whereby stiffness differs from age-matched controls in younger but not older Marfan patients [73].

Natural turnover rates of collagen are orders of magnitude higher than those of elastin [26,64], thus collagen remodeling is a ubiquitous part of most aortic adaptations and maladaptations, including TAAs. Importantly, mutations to the gene that encodes lysyl oxidase (*LOX*), which governs enzymatic cross-linking of collagen, result in aneurysms in humans and mice [74–76]. Note that the normal accumulation of mature collagen is prevented in these germline mutations. Allied studies show further that globally blocking lysyl oxidase in mature mice, in which the aortic collagen had matured, results in a modest phenotype unless superimposed on other defects, particularly Marfan syndrome [77] or a model of aortic aneurysm initiated by the degradation of elastin via elastase [78]. Our results reflect but extend these findings. Whereas use of β -aminopropionitrile (BAPN) in animal models results in uniform reductions in collagen cross-linking allowed marked dilatations of the thoracic aorta. Interestingly, superimposing hypertension on the vulnerable aorta increased the localized dilatation more than did superimposing aging, though both exacerbated the situation. Localized collagen degradation had similar effects.

Smooth muscle cell-induced vasoconstriction also enables the aortic wall to resist distension under the action of blood pressure. Multiple mutations predisposing to TAAs, including those encoding fibrillin-1 and fibulin-4 and especially those encoding smooth muscle α -actin and smooth muscle myosin heavy chain, compromise vessel-level contractility [16,52,79,80]. Some have thus implied that loss of vessel-level contractility predisposes to TAAs, yet smooth muscle contractility is also compromised in cases wherein TAAs do not develop, including fibulin 5 null mice [81]. We studied potential effects of localized losses in overall contractility. which resulted in modest localized dilatations in the case of an otherwise normal aortic wall exposed to normal pressure loading. Both hypertension and aging increased these local dilatations, though the changes did not reach aneurysmal levels even when blocking contractility fully. Importantly, albeit not shown, marked dilatations arose when both risk factors were superimposed on the initial smooth muscle insult. This simulation is consistent with a study that showed that pressure elevation augments aortic dilatation in the $Acta2^{-/-}$ mouse, which has reduced vessel-level contractility [82]. Nevertheless, despite some suggestions to the contrary, our simulations suggest that localized losses in vessel-level contractility need not be strong initiators of TAAs. Toward this end, we emphasize that cell-level contractility [80] is controlled by the same actomyosin regulators as is vessel-level contractility [52], with cell-level manifestations affecting both cell migration and the mechanosensing and mechanoregulation of matrix.

Importantly, simulations of potential effects of localized losses of mechanosensing or mechanoregulation of matrix revealed a high propensity toward aneurysmal dilatation. In particular, we modeled compromised mechanosensing by altering the percentage of the intramural stress ($\delta \leq 1$, see Methods) that is transduced to influence matrix production. We have suggested that the propensity of TAAs to form in cases of either fibrillin-1 mutations or smooth muscle cell dysfunction, among others, points to the mechanotransduction axis as a potential initiator of these lesions [6]. Recent studies in cells and different genetically modified mice support this hypothesis [80,83–85]. This in silico study is the first, however, to examine mechanosensing in isolation. The simulations suggest that increasingly greater reductions in localized mechanosensing give rise to increasingly larger localized dilatations (Fig 3), and superimposing aging and especially hypertension increase these dilatations. Because compromised mechanosensing can result from dysfunctional actomyosin activity, comparing these

results (fourth rows in Figs 1 and 2) to those for lost vessel-level contractility (third rows in Figs 1 and 2) suggest that compromised mechanosensing has a more severe effect on the aneurysmal phenotype. Whereas vessel-level contractility affects aortic diameter, and thus wall stress [58], mechanosensing affects the ability of the cells to sense the stress and respond appropriately via matrix turnover [6,61]. Compromised mechanoregulation of the newly deposited collagen, modeled via altered deposition stretches, similarly resulted in marked localized dilatations.

Hence, although actomyosin pathways necessarily overlap for both vessel-level contractility and mechanosensing / mechanoregulation of the matrix, the present simulations suggest that aortic consequences are very different. Our findings are thus consistent with an earlier suggestion that the primary role of actomyosin activity in the aorta is "in managing matricellular interactions and sensing the extracellular environment to control appropriate elaboration and maintenance of extracellular matrix" [3]. It is similarly consistent with a prior computational finding that showed that uniform losses in smooth muscle contractility tend to disrupt arterial mechano-adaptations less than does uniformly compromised mechanobiologically mediated matrix turnover [24]. Importantly, our results are also consistent with recent findings that decreased actomyosin activity in smooth muscle cells from $Acta2^{-/-}$ mice have reduced integrin expression and reduced mechanosensing [86]. In summary, the present simulations further support the hypothesis that dysfunctional mechanosensing is a contributor to TAAs [5–8].

Whereas the present model allows many effects to be studied in isolation, data obtained from actual vessels necessarily reveal intrinsic couplings. To name a few, loss of elastic fiber integrity in a mouse model of Marfan syndrome associates with reduced vasoconstrictive capability [52]; loss of elastic fiber integrity also alters the micro-properties [53] and undulation [87] of the neighboring collagen fibers; deletion of integrin linked kinase alters smooth muscle contractile protein expression, not just cell-matrix interactions [88]; and disruption of fibulin-4, an elastin associated glycoprotein, leads to changes in collagen synthesis and maturation [54] as well as cytoskeletal actin [83,89]. For the purposes of this study, however, we exploited the advantage of in silico simulations and studied in isolation tissue-level consequences of five key genetically predisposed alterations with superimposed hypertension and aging. These simulations support a "multi-hit" hypothesis in that the aorta tends to be mechano-adaptive, and thus fault tolerant, when subjected to mild insults, but it succumbs to severe and especially multiple insults, or hits. Controlling risk factors such as hypertension is critical [90], though some interpretations may suggest otherwise depending on the mouse model used [91].

Notwithstanding the insights gained, there is a need to consider further determinants as well. Endothelial dysfunction can result in reduced nitric oxide bioavailability, hence promoting smooth muscle contractility but also enabling inflammation, which appears to increase with prior cell and matrix damage [92]. Phenotypic modulation of smooth muscle cells from contractile to synthetic can reduce contractility, but also disturb balances in matrix production and removal. Smooth muscle apoptosis necessarily reduces overall vasoconstriction, but also mechanosensing and mechanoregulation of matrix. Losses in elastic fiber integrity reduces wall resilience, but in some cases also the ability of the smooth muscle cells to mechanosense the mechanical state of the wall. There is, therefore, a need to consider many additional combinations of hits, not just superimposed hypertension and aging. We also did not account for initial aortic curvature, which is particularly important in the ascending aorta, or the pulsatility of loading, which includes cyclic distension and extension in the ascending aorta. We did not account for changes in hemodynamics as the aorta enlarges (except indirectly in Figs D, E, F in S1 Supporting Information), though we assumed that wall shear stress was sensed only within the regions that were free of the prescribed insults. Wall shear stress regulation, along with the relative rates of turnover of smooth muscle versus collagen, is critical for maintaining

homeostatic control of the distal and proximal segments. Fluid-solid-growth models [93] will enable broader follow-up studies. Finally, we did not account for different ages at which the insult occurred, the sex of the mice, or observed mouse-to-mouse variability in geometry, composition, or wall properties, which would provide better bounds on expected behaviors, particularly why certain lesions appear earlier or progress more rapidly. Clearly more data will be needed to inform more complete models.

Rather, we established a baseline model using mean data from one particular group of male mice, and insults of interest were introduced consistently in an otherwise initially straight circular segment of a model thoracic aorta, which thereby enabled consistent comparisons across groups and clearer delineation of the effects of the insults in isolation and in combination. Not surprisingly, loss of matrix integrity, via either compromised elastic fibers or poorly remodeled collagen, enables marked dilatations. Perhaps surprisingly, compromised mechanosensing and mechanoregulation can evoke as strong of an aneurysmal propensity whereas loss of vessel-level smooth muscle contractility does not. Hence, it appears that reduced actomyosin activity is more severe in its effects in compromising mechanosensing and mechanoregulation of matrix (cf. [60,86,94]). There is a need for caution, therefore, not to over-interpret consequences of compromised vessel-level contractility, often measured via wire or pressure myography [52,79], even though many if not most mutations predisposing to TAAs (e.g., *Fbn1*, *Acta2*, *Myh11*, *Mylk*, *Tgfbr1*, and *Prkg1* mutations in mice) reduce vessel-level contractility.

Whereas we focused on aortic dimensions and properties appropriate for mice, because myriad mouse models are available to study the natural history of TAAs, there will be a need to repeat these simulations for human lesions and it is possible that contractility plays an even lesser role in the human than murine aorta. We otherwise expect, however, that the basic theoretical framework, computational approach, and functional forms of most constitutive equations should translate from the mouse to the human, thus necessitating primarily a re-parameterization of the model when studying human lesions. Consistent with an earlier suggestion that "Over time, the major challenge in aortic aneurysmal research will likely shift from gene identification to the assessment of gene product function in large vessel homeostasis" [3], we submit that in silico studies (including digital twins) can complement in vivo and ex vivo studies as we seek to understand and eventually better treat these complex, multi-factorial lesions, with data-informed patient-specific computations (e.g., [95]) eventually the way forward.

Supporting information

S1 Appendix. Tangent moduli contribution for reoriented / remodeled diagonal fibers via Eq (14).

(PDF)

S1 Supporting Information. Supplemental Figs A to K. (PDF)

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Visualization: M. Latorre, J. D. Humphrey.

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