Growing skin: Tissue expansion in pediatric forehead reconstruction

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Abstract Tissue expansion is a common surgical procedure to grow extra skin through controlled mechanical over-stretch. It creates skin that matches the color, texture, and thickness of the surrounding tissue, while minimizing scars and risk of rejection. Despite intense research in tissue expansion and skin growth, there is a clear knowledge gap between heuristic observation and mechanistic understanding of the key phenomena that drive the growth process. Here, we show that a continuum mechanics approach, embedded in a custom-designed finite element model, informed by medical imaging, provides valuable insight into the biomechanics of skin growth. In particular, we model skin growth using the concept of an incompatible growth configuration. We characterize its evolution in time using a second-order growth tensor parameterized in terms of a scalar-valued internal variable, the in-plane area growth. When stretched beyond the physiological level, new skin is created, and the in-plane area growth increases. For the first time, we simulate tissue expansion on a patient-specific geometric model, and predict stress, strain, and area gain at three expanded locations in a pediatric skull: in the scalp, in the forehead, and in the cheek. Our results may help the surgeon to prevent tissue over-stretch and make informed decisions about expander geometry, size, placement, and inflation. We anticipate our study to open new avenues in reconstructive surgery, and enhance treatment for patients with birth defects, burn injuries, or breast tumor removal.

Key words growth, residual stress, finite element analysis, skin, tissue expansion, reconstructive surgery

1 Motivation

One percent of neonates is born with congenital melanocytic nevi, dark-colored surface lesions present at birth [10]. Congenital nevi may vary in size, shape, texture, color, hairiness, and location, but they have one thing in common: their high malignant potential [23]. Birthmarks larger than 10 cm in diameter are classified as giant congenital nevi and have a prevalence of one in 20,000 infants [45]. Because giant congenital nevi place the child at an increased risk to develop skin cancer, surgical excision remains the standard treatment option [20]. Cosmetic deformity, significant aesthetic disfigurement, and severe psychological distress are additional compelling reasons for nevus removal, especially in the craniofacial region [23].

To reconstruct the defect, preserve function, and maintain aesthetic appearance, tissue expansion has become a major treatment modality in the management of giant congenital nevi [36]. Tissue expansion was first proposed more than half a century ago to reconstruct a traumatic ear and has since then revolutionized reconstructive surgery [43]. Today it is widely used to repair birth defects [5], correct burn injuries [3], and reconstruct breasts after tumor removal [46]. Tissue expansion is the ideal strategy to grow skin that matches the color, texture, hair bearing, and thickness of the surrounding healthy skin, while minimizing scars and risk of rejection [49].

Figure 1, left, shows a one-year old boy who presented with a giant congenital nevus concerning 25 percent of his forehead, extending to the right temporal scalp and cheek [21]. To resurface the nevus region and stimulate in situ skin growth, three simultaneous forehead, cheek, and scalp expanders are used. They are implanted in subcutaneous pockets adjacent to the defect, where they are gradually filled with saline solution. The amount of filling is controlled by visual inspection of skin color and capillary refill [49]. Multiple serial inflations stretch the skin and stimulate tissue growth over a period of several weeks [63]. Once enough skin is created, the expanders are removed, the nevus is excised, and the newly grown skin flaps are advanced to close the defect zone. Figure 1, right, shows the boy at age three, after completed fore-
Fig. 1 Tissue expansion for pediatric forehead reconstruction. The patient, a one-year-old boy presented with a giant congenital nevus involving 25 percent of his forehead, extending to the right temporal scalp and cheek. Simultaneous forehead, cheek, and scalp expanders were implanted for in situ skin growth. This technique allows to resurface large anatomical areas with skin of similar color, quality, and texture. The follow-up photograph shows the boy at age three, after completed forehead, scalp, and cheek reconstruction.

Figure 2 shows a schematic sequence of the mechanical processes during tissue expansion. Initially, at biological equilibrium, the skin is in a natural state of resting tension [54]. When the expander is implanted and inflated, skin is loaded in tension. Stretch beyond a critical level triggers a series of signaling pathways eventually leading to the creation of new skin [57]. On the cellular level, mechanotransduction affects a network of several integrated cascades including growth factors, cytoskeletal rearrangement, and protein kinases [11]. On the tissue level, skin growth induces stress relaxation and restores the state of resting tension [54]. The cycle of expander inflation, stretch, growth, and relaxation is repeated multiple times, typically on a weekly basis [63]. As demonstrated by immunocytochemistry, the expanded tissue undergoes normal cell differentiation and maintains its characteristic phenotype [61]. Skin initially displays thickness changes upon expansion, however, these changes are fully reversible upon expander removal [59]. When the expander is removed, the skin retracts and reveals the irreversible nature of skin growth, associated with growth-induced residual stresses [18, 41]. Figure 3 shows a commonly used tissue expander to grow skin in reconstructive surgery.

To predict stress, strain, and area gain during tissue expansion in pediatric forehead reconstruction, we adopt a novel mechanistic approach [8, 9], based on the continuum framework of finite growth [50]. Originally developed for isotropic volumetric growth [14, 37], finite growth theories are based on the multiplicative decomposition of the deformation gradient into an elastic and a growth part [15, 38], a concept adopted from finite...
plasticity [34]. Depending on the format of their growth tensor, continuum growth theories have been refined to characterize isotropic [19, 32], transversely isotropic [48, 56], orthotropic [17], or generally anisotropic growth [40, 42], either compressible [40] or incompressible [51]. Recent trends focus on the computational modeling of finite growth [22], typically by introducing the growth tensor as an internal variable within a finite element framework [16, 25], a strategy that we also adopt here. A recent monograph that compares different approaches to growth and summarizes the essential findings, trends, and open questions in this progressively evolving new field [2]. Despite ongoing research in growing biological systems, the growth of thin biological membranes remains severely understudied. Only few attempts address the growth of thin biological plates [12] and membranes [40]. Motivated by a first study on axisymmetric skin growth [55], we have recently established a prototype model for growing membranes to predict skin expansion in a general three-dimensional setting [8]. This study capitalizes on recent developments in reconstructive surgery, continuum mechanics of growing tissues, and computational modeling, supplemented by medical image analysis. It documents our first attempts to model and simulate skin expansion in pediatric forehead reconstruction using a real patient-specific geometry.

2 Methods

2.1 Continuum modeling of skin growth

We adopt the kinematics of finite deformations and introduce the deformation map \( \varphi \), which, at any given time \( t \) maps the material placement \( X \) of a physical particle in the material configuration to its spatial placement \( x \) in the spatial configuration, \( x = \varphi (X, t) \). We choose a formulation which is entirely related to the material frame of reference, and use \( \nabla \{ \varphi \} = \partial x \{ \varphi \} \) and \( \text{Div} \{ \varphi \} = \partial x \{ \varphi \} \) to denote the gradient and the divergence of any field \( \{ \varphi \}(X, t) \) with respect to the material placement \( X \) at fixed time \( t \). Here, \( I \) is the material identity tensor. To characterize finite growth, we introduce an incompatible growth configuration, and adopt the multiplicative decomposition of the deformation gradient

\[
F = \nabla X \varphi = F^e \cdot F^g
\]  

(1)

into a reversible elastic part \( F^e \) and an irreversible growth part \( F^g \). This multiplicative decomposition, reminiscent of the decomposition of the elastoplastic deformation gradient [34], was first used to describe growth of biological tissues in [50]. Similarly, we can then decompose the total Jacobian

\[
J = \det (F) = J^e \cdot J^g
\]  

(2)

We adopt the kinematics of finite deformations and in-

2.2 Computational modeling

and simulate skin expansion in pediatric forehead recon-

struction in an elastic part \( J^e = \det (F^e) \) and a growth part \( J^g = \det (F^g) \). We idealize skin as a thin layer character-ized through the unit normal \( n_0 \) in the undeformed reference configuration. The length of the deformed skin normal \( n = \text{cof}(F) \cdot n_0 = J F^{-t} \cdot n_0 \) introduces the area stretch

\[
\vartheta = || \text{cof}(F) \cdot n_0 || = \vartheta^e \vartheta^g
\]  

(3)

which we can again decompose into an elastic area stretch \( \vartheta^e = || \text{cof}(F^e) \cdot n_0 || /|| n_0 || \) and a growth area stretch \( \vartheta^g = || \text{cof}(F^g) \cdot n_0 || \) [8]. Here, \( n^g = \text{cof}(F^g) \cdot n_0 = J^g F^g - t \cdot n_0 \) denotes the grown skin normal, and \( \text{cof}(\varphi) = \det(\varphi) (\varphi)^{-t} \) denotes the cofactor of the second order tensor (\( \varphi \)). As characteristic deformation measures, we introduce the right Cauchy Green tensor \( C \) in the undeformed reference configuration

\[
C = F^e \cdot F = F^{e t} \cdot F^e \cdot F^g
\]  

(4)

and its elastic counterpart \( C^e = F^{e t} \cdot F^e = F^{g - t} \cdot C \cdot F^{g - 1} \) in the intermediate configuration. This allows us to rephrase the total area stretch as \( \vartheta = J [n_0 \cdot C^{-1} \cdot n_0]^{1/2} \). Finally, we introduce the pull back of the spatial velocity gradient \( t = F \cdot F^{-1} \) to the intermediate configuration,

\[
F^{-1} \cdot t \cdot F^e = L^e + L^g
\]  

(5)

which obeys the additive decomposition into the elastic velocity gradient \( L^e = F^{e t} \cdot F^e \) and the growth velocity gradient \( L^g = F^{g t} \cdot F^g \). Here, \( \{ \dot{\varphi} \} = \partial_t \{ \varphi \} \) \( X \) denotes the material time derivative of any field \( \{ \varphi \}(X, t) \) at fixed material placement \( X \).

We characterize growing tissue using the framework of open system thermodynamics in which the material den-

sity \( \rho_0 \) is allowed to change as a consequence of growth [26, 28]. The balance of mass for open systems balances its rate of change \( \rho_0 \) with a possible in- or outflux of mass \( R \) and mass source \( \mathcal{R}_0 \) [29, 44].

\[
\dot{\rho}_0 = \text{Div} (R) + \mathcal{R}_0
\]  

(6)

Similarly, the balance of linear momentum balances the density-weighted rate of change of the velocity \( \rho_0 \dot{v} = \rho_0 \dot{\varphi} \), with the momentum flux \( P = F \cdot S \), and the moment-

source \( \rho_0 b \),

\[
\rho_0 \dot{v} = \text{Div} (F \cdot S) + \rho_0 b
\]  

(7)

here stated in its mass-specific form [27]. \( P \) and \( S \) are the first and second Piola-Kirchhoff stress tensors. Last, we would like to point out that the dissipation inequality for open systems

\[
\rho_0 D = S : \frac{1}{2} C - \rho_0 \dot{\varphi} - \rho_0 S \geq 0
\]  

(8)

typically contains an extra entropy source \( \rho_0 S \) to ac-

count for the growing nature of living biological sys-

tems [26, 41]. Equations (7) and (8) represent the mass-
specific versions of the balance of momentum and of the
dissipation inequality which are particularly useful in the context of growth since they contain no explicit dependencies on the changes in mass [27].

To close the set of equations, we introduce the constitutive equations for the mass source $R_0$, for the momentum flux $S$, and for the growth tensor $F^g$, assuming that the mass flux $R = 0$, the momentum source $b = 0$, and the acceleration $a = 0$ are negligibly small. On the cellular level, immunocytochemistry has shown that expanded tissue undergoes normal epidermal cell differentiation [61]. On the organ level, mechanical testing has confirmed that the newly grown skin has the same material properties as the initial tissue [63]. Accordingly, we assume that the newly grown skin has the same density as the initial tissue. This implies that the mass source

$$R_0 = \rho_0 \text{tr}(L^g)$$

(9)

can be expressed as the density-weighted trace of the growth velocity gradient $\text{tr}(L^g) = \dot{F}^g : F^g^{-1}$ [22]. We model skin as a hyperelastic material characterized through the Helmholtz free energy $\psi = \psi(C, F^g)$, which we use to evaluate the dissipative moduli $\mathbf{\mathcal{L}}^g$.

$$\rho_0 \mathbf{\mathcal{D}} = \left[ S - \rho_0 \frac{\partial \psi}{\partial C} \right] : \frac{1}{2} \dot{C} + M^e : L^g - \rho_0 \frac{\partial \psi}{\partial \rho_0} \rho_0 S_0 \geq 0$$

(10)

We observe that the Mandel stress of the intermediate configuration $M^e = C^e \cdot S^e$ is energetically conjugate to the growth velocity gradient $L^g = \dot{F}^g \cdot F^g^{-1}$. From the dissipation inequality (10), we obtain the definition of the second Piola Kirchhoff stress $S$ as thermodynamically conjugate quantity to the right Cauchy Green deformation tensor $C$.

$$S = 2 \rho_0 \frac{\partial \psi}{\partial C} = 2 \frac{\partial \psi}{\partial C^e} : \frac{\partial C^e}{\partial C} = F^g^{-1} : S^e \cdot F^g^{-1}$$

(11)

According to this definition, the first derivative of the Helmholtz free energy $\psi$ with respect to the elastic right Cauchy Green tensor $C^e$ introduces the elastic second Piola Kirchhoff stress $S^e$, while the second derivative defines the elastic constitutive moduli $\mathbf{\mathcal{L}}^e$.

$$S^e = 2 \rho_0 \frac{\partial \psi}{\partial C^e} \text{ and } \mathbf{\mathcal{L}}^e = 2 \frac{\partial S^e}{\partial C^e} = 4 \rho_0 \frac{\partial^2 \psi}{\partial C^e \otimes \partial C^e}$$

(12)

To focus on the impact of growth, rather than adopting a sophisticated anisotropic material model for skin [9,30], we assume a classical Neo-Hookean free energy $\rho_0 \psi = \frac{\lambda}{2} \ln^2(J^e) + \frac{\mu}{2} \left[ C^e : I - 3 - 2 \ln(J^e) \right]$, introducing the elastic second Piola Kirchhoff stress $S^e = [ \lambda \ln(J^e) - \mu ] C^e^{-1} + \mu \mathbf{I}$, and the elastic constitutive moduli $L^e = \lambda C^e^{-1} \otimes C^e^{-1} + [ \mu - \lambda \ln(J^e) ] \left[ C^e \otimes C^e + C^e \otimes C^e \right]$. Motivated by clinical observations [49], we classify skin growth as a strain-driven, transversely isotropic, irreversible process. It is characterized through one single growth multiplier $\vartheta^g$ that reflects the irreversible area increase perpendicular to the skin normal $n_0$.

$$F^g = \sqrt{\vartheta^g} \mathbf{I} + [ 1 - \sqrt{\vartheta^g} ] n_0 \otimes n_0$$

(13)

For this particular type of transversely isotropic growth, for which all thickness changes are reversibly elastic [59], area growth is identical to volume growth, i.e., $\vartheta^g = \det(F^g) = J^g$. Because of the simple rank-one update structure in (13), we can invert the growth tensor explicitly, $F^g^{-1} = 1/\sqrt{\vartheta^g} \mathbf{I} + [ 1 - 1/\sqrt{\vartheta^g} ] n_0 \otimes n_0$, using the Sherman-Morrison formula. This explicit representation introduces the following simple expression for the growth velocity gradient,

$$L^g = \sqrt{\vartheta^g}/\sqrt{\vartheta^g} \mathbf{I} + [ 1 - \sqrt{\vartheta^g}/\sqrt{\vartheta^g} ] n_0 \otimes n_0$$

(14)

which proves convenient to explicitly evaluate the mass source in equation (9) as $R_0 = \rho_0 \left[ 1 + 2 \sqrt{\vartheta^g}/\sqrt{\vartheta^g} \right]$. Motivated by physiological observations of stretch-induced skin expansion [21], we adopt the following evolution equation for the growth multiplier,

$$\dot{\vartheta^g} = k_g (\vartheta^g)^{\phi (\vartheta^g)}$$

(15)

which follows a well-established functional form [37], but is now rephrased in a strain-driven format [17]. To control unbounded growth, we introduce the weighting function

$$k_g = \frac{1}{\tau} \left[ \frac{\vartheta_{\max} - \vartheta^g}{\vartheta_{\max} - \vartheta_{\crit}} \right]$$

(16)

where $1/\tau$ controls the adaptation speed, the exponent $\gamma$ calibrates the shape of the growth curve, and $\vartheta_{\max} > 1$ is the maximum area growth [22,37]. The growth criterion

$$\phi (\vartheta^g - \vartheta_{\crit}) = \langle \dot{\vartheta}/\dot{\vartheta^g} - \vartheta_{\crit} \rangle$$

(17)

is driven by the elastic area stretch $\varepsilon = \vartheta/\vartheta^g$, such that growth is activated only if the elastic area stretch exceeds a critical physiological stretch limit $\vartheta_{\crit}$. Here, $\langle \rangle$ denote the Macaulay brackets.

Figure 4 displays the constitutive response of the four-parameter growth model in equi-biaxial stretch. At a prescribed piecewise constant total stretch $\vartheta$, the growth stretch $\vartheta^g$ increases gradually while the elastic stretch $\varepsilon$ decreases. This induces stress relaxation. Horizontal dashed lines represent the elastic stretch limit beyond which skin growth is activated $\vartheta_{\crit}$ and the maximum area growth $\vartheta_{\max}$. Increased adaptation speeds $1/\tau$ ↑ and decreased growth exponents $\gamma$ ↓ both accelerate convergence towards the biological equilibrium [22], but do not affect the final equilibrium state [48,51]. At all times, the multiplicative decomposition of the deformation gradient $F = F^e \cdot F^g$ introduced in equation (1) carries over to the multiplicative decomposition of the total area stretch $\vartheta = \vartheta^e \vartheta^g$ of equation (3).

2.2 Computational modeling of skin growth

We solve the coupled biological and mechanical equilibrium for skin growth within an incremental iterative finite element setting [58]. To characterize the growth
process at each instant in time, we introduce the growth multiplier $\vartheta^g$ as an internal variable, and solve the biological equation (15) locally at the integration point level. For the temporal discretization, we partition the time interval of interest $T$ into $n_{dp}$ subintervals, $T = \cup_{n=1}^{n_{dp}} [t_n, t_{n+1}]$ and focus on the interval $[t_n, t_{n+1}]$ for which $\Delta t = t_{n+1} - t_n > 0$ denotes the current time increment. Our goal is to determine the current growth multiplier $\vartheta^g$ for a given deformation state $F$ at time $t$, and a given growth multiplier $\vartheta^g_n$ at the end of the previous time step $t_n$. For the sake of compactness, we omit the index $(n+1)$ for all quantities at the end of the current time step $t_{n+1}$. We evaluate the material time derivative of the growth multiplier $\dot{\vartheta}^g = [\vartheta^g - \vartheta^g_n] / \Delta t$ using a finite difference approximation. In the spirit of implicit time integration schemes, we now reformulate the evolution equation (15), introducing the discrete residual $R^g$ in terms of the unknown growth multiplier $\vartheta^g$.

$$R^g = \vartheta^g - \vartheta^g_n - k^g \vartheta^g \Delta t = 0$$  (18)

We solve this nonlinear residual equation using a local Newton iteration. Within each iteration step, we calculate the linearization of the residual $R^g$ with respect to the growth multiplier $\vartheta^g$.

$$K^g = \frac{\partial R^g}{\partial \vartheta^g} = 1 - \left[ \frac{\partial k^g}{\partial \vartheta^g} \vartheta^g + k^g \frac{\partial \vartheta^g}{\partial \vartheta^g} \right] \Delta t$$  (19)

with the derivatives of the weighting function $\partial k^g / \partial \vartheta^g = - \gamma k / [\vartheta^g_{\text{max}} - \vartheta^g]$ and the growth criterion $\partial \vartheta^g / \partial \vartheta^g = - \dot{\vartheta} / \vartheta^g \Delta t$ introduced in equations (16) and (17). Within each iteration step, we iteratively update the unknown growth multiplier $\vartheta^g \leftarrow \vartheta^g - R^g / K^g$ until convergence is achieved, i.e., until the local growth update $\Delta \vartheta^g = - R^g / K^g$ reaches a user-defined tolerance.

To explore the interplay between growth and mechanics, we discretize the deformation map $\varphi$ as nodal degree of freedom, and solve the mechanical equilibrium (7) globally at the node point level. To solve the quasi-static mechanical equilibrium, $\text{Div} (F \cdot S) = 0$, we cast it into its weak form, $\int_{\Omega} \nabla \varphi : [F \cdot S] \, dV = 0$, through multiplication with the test function $\delta \varphi$ and integration over the domain of interest $\Omega$. For the spatial discretization, we partition the domain of interest $\Omega = \cup_{e=1}^{n_{el}} \Omega^e$ into $n_{el}$ finite elements $\Omega^e$. Our goal is to determine the deformation state $\varphi$ for a given load at time $t$. To approximate the test function $\delta \varphi = \sum_{i=1}^{n_{el}} N^i \delta \varphi_i$, the unknown deformation $\varphi = \sum_{i=1}^{n_{el}} N^i \delta \varphi_i$, and their gradients $\nabla \delta \varphi = \sum_{i=1}^{n_{el}} \delta \varphi_i \otimes \nabla N^i$ and $\nabla \varphi = \sum_{i=1}^{n_{el}} N^i \otimes \nabla N^i$, we apply an isoparametric Bubnov-Galerkin based finite element interpolation, where $N^i$ and $N^j$ are the element shape functions and $i, j = 1, \ldots, n_{el}$ are the element nodes. We now reformulate the weak form of the balance of linear momentum (7) with help of these finite element approximations, introducing the discrete residual $R^g_j$ in terms of the unknown nodal deformation $\varphi_j$.

$$R^g_j = \frac{n_{el}}{A_e} \int_{\Omega^e} \nabla N^j \cdot [F \cdot S] \, dV_e = 0$$  (20)

Here, the operator $A$ symbolizes the assembly of all element residuals at the $j = 1, \ldots, n_{el}$ element nodes to the global residual at the global node points $J = 1, \ldots, n_g$. We evaluate the global discrete residual (20), once we have iteratively determined the growth multiplier $\vartheta^g$ for the given deformation state $F$ and the given history $\vartheta^g_n$ as described in the previous section. Then we successively determine the growth tensor $F^g$ from equation (13), the elastic tensor $F^e = F^g \cdot F^{-1}$ from equation (1), the elastic stress $S^e$ from equation (12), and lastly, the second Piola Kirchhoff stress $S$ from equation (11). Again, we suggest an incremental iterative Newton algorithm to solve the nonlinear residual equation for the unknown deformation (20). The consistent linearization of the residual $R^g_j$ with respect to the nodal vector of unknowns $\varphi_j$ introduces the global stiffness matrix.

$$K_{ij} = \frac{\partial R^g_i}{\partial \varphi_j} = \frac{n_{el}}{A_e} \int_{\Omega^e} \nabla N^i \cdot S \cdot \nabla N^j \, dV_e + [\nabla N^j \cdot F]_{\text{sym}} \cdot L \cdot [F^e \cdot \nabla N^i]_{\text{sym}} \, dV_e$$  (21)

The fourth order tensor $L$ denotes the Lagrangian constitutive moduli which, we can determine directly from the linearization of the Piola Kirchhoff stress $S$ with respect to the total right Cauchy Green tensor $C$ [16].

$$L = 2 \frac{dS}{dC} = 2 \frac{dS}{dC} = 2 \left[ \frac{\partial S}{\partial F} \cdot \frac{\partial F}{\partial \vartheta^g} \right] \otimes \frac{\partial \vartheta^g}{\partial C} |_F$$  (22)
The first term
\[ 2 \frac{\partial S}{\partial C} = \left[ F^k - 1 \otimes F^k - 1 \right] \cdot L^c : \left[ F^k - t \otimes F^k - t \right] \quad (23) \]
represents the pull back of the elastic moduli \( L^c \) introduced in equation (12) onto the reference configuration. Here we have used the abbreviations \( \{ \otimes \} \) for the non-standard fourth order products. The second term
\[ \frac{\partial S}{\partial F^k} = - \left[ F^k - 1 \otimes S + S \otimes F^k - 1 \right] \]
\[ - \left[ F^k - 1 \otimes F^k - 1 \right] \cdot \frac{1}{2} L^c : \left[ F^k - t \otimes C^e + C^e \otimes F^k - t \right] \quad (24) \]
consists of two terms that resemble a geometric and a material stiffness in nonlinear continuum mechanics. The third term
\[ \frac{\partial F^k}{\partial \vartheta^e} = \frac{1}{2 \sqrt{\vartheta}} \left[ I - n_0 \otimes n_0 \right] \quad (25) \]
and the fourth term
\[ \frac{\partial \vartheta^e}{\partial C} = 1 \frac{1}{\vartheta} \left[ \frac{\vartheta^{\text{max}} - \vartheta^e}{\vartheta^{\text{max}} - 1} \right] \frac{1}{\vartheta} \Delta t \]
\[ = \frac{1}{2} \vartheta C^{-1} - \frac{1}{2} \frac{J^2}{\vartheta} [C^{-1}, n_0] \otimes [C^{-1}, n_0] \quad (26) \]
\[ \vartheta^e \}
\[ \begin{aligned}
\text{depend on the particular choice for the growth tensor } F^k \\
\text{in equation (13) and on the evolution equation for the growth multiplier } \vartheta^e \\
\text{in equation (15), respectively. For each global Newton iteration step, we iteratively update the current deformation state } \varphi \leftarrow \varphi - K^e_1 \cdot R^e_1 \\
\text{until we achieve algorithmic convergence. Upon convergence, we store the corresponding growth multipliers } \vartheta^e \\
\text{at the integration point level. Table 1 summarizes the algorithmic treatment of skin growth at the integration point level.} \\
\end{aligned} \]

<table>
<thead>
<tr>
<th>Table 1: Algorithmic flowchart for strain-driven transversely isotropic area growth.</th>
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<tbody>
<tr>
<td>given ( F ) and ( \vartheta^e_0 )</td>
</tr>
<tr>
<td>initialize ( \vartheta^e \leftarrow \vartheta^e_0 )</td>
</tr>
<tr>
<td>local Newton iteration</td>
</tr>
<tr>
<td>check growth criterion ( \vartheta^e = \vartheta^e - \vartheta^e_{\text{crit}} \geq 0 ) ? ( (17) )</td>
</tr>
<tr>
<td>calculate residual ( R = \vartheta^e - \vartheta^e_0 - k^e \vartheta^e \Delta t ) ( (18) )</td>
</tr>
<tr>
<td>calculate tangent ( K = \partial R / \partial \vartheta^e ) ( (19) )</td>
</tr>
<tr>
<td>update growth stretch ( \vartheta^e \leftarrow \vartheta^e - R / K )</td>
</tr>
<tr>
<td>check convergence ( R \leq \text{tol} ) ?</td>
</tr>
<tr>
<td>calculate growth tensor ( F^k ) ( (13) )</td>
</tr>
<tr>
<td>calculate elastic tensor ( F^e = F \cdot F^k ) ( (1) )</td>
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<td>calculate elastic right Cauchy Green tensor ( C^e ) ( (4) )</td>
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<tr>
<td>calculate elastic second Piola Kirchhoff stress ( S^e ) ( (12) )</td>
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<tr>
<td>calculate second Piola Kirchhoff stress ( S ) ( (11) )</td>
</tr>
<tr>
<td>calculate Lagrangian moduli ( L ) ( (22) )</td>
</tr>
</tbody>
</table>

2.3 Geometric modeling of skin growth

To simulate skin growth on an anatomically exact geometry, we create a finite element mesh on the basis of three-dimensional computer tomography scans shown in Figure 5. Figure 6 summarizes the sequence of steps to generate our patient-specific geometric model. First, we identify the skin region by a distinct grey scale value in the computer tomography scans and extract point cloud data of its boundary. Figure 6, left, mimics the discrete nature of the extracted point cloud, with high point densities in the scanning plane and low point densities between the distinct planes. To smooth the data and decreases the overall number of points, we homogenize the point cloud using a median filter. Next, we create a triangular surface mesh from the smoothened point cloud by applying a ball-pivoting algorithm [7]. Ball-pivoting algorithms are particularly suited for surface reconstruction of large data sets. After placing an initial seed element, the ball-pivoting algorithm rotates a sphere over the edges of this element and sequentially creates new elements whenever the sphere touches three data points. However, since our data are based on plane-wise computer tomography scans, ball-pivoting algorithms typically fail to automatically create smooth surfaces. Unfortunately, other fully automated meshing strategies such as convex hull or shrink wrap algorithms are not suitable for non-convex geometries like the face, which possesses several non-convexities in the eye, nose, mouth, and ear regions [24]. Accordingly, we smoothen the triangular surface mesh semi-manually, as illustrated in Figure 6, middle.

From the smoothened surface mesh, we finally create a one-element thick volume mesh of the pediatric skull, discretized with 61,228 nodes, 183,684 degrees of freedom, and 30,889 tri-linear brick elements. As a first approximation, we assume that all eight integration points within each element possess the same skin plane normal \( n_0 \), corresponding to the normal from the initial surface mesh. We virtually implant three tissue expanders as shown in Figure 6, right, motivated by the tissue ex-
Fig. 6 Mesh generation from clinical images. From the computer tomography scans, we extract discrete point cloud data (left), which we filter and mesh using a ball-pivoting algorithm. This generates a triangular surface mesh, which is further smoothed (middle) and turned into a final volume mesh (right). The final discretization of the skull consists of 61,228 nodes, 183,684 degrees of freedom, and 30,889 tri-linear brick elements. We virtually implant three tissue expanders, one in the scalp, discretized with 4,356 nodes, 13,068 degrees of freedom, and 2,088 brick elements (red), one in the cheek, discretized with 2,542 nodes, 7,626 degrees of freedom, and 1,200 brick elements (yellow), and one in the forehead, discretized with 3,782 nodes, 11,346 degrees of freedom, and 1,800 brick elements (blue).

3 Results

We illustrate the impact of tissue expansion at three characteristic locations of the skull, in the scalp, the forehead, and the cheek. For the elastic model, we assume Lamé constants of $\lambda = 0.7141$ MPa and $\mu = 0.1785$ MPa, which would correspond to a Poisson’s ratio of $\nu = 0.4$ and a Young’s modulus of $E = 0.5$ MPa in the linear regime [1,52]. For the growth model, we assume a critical threshold of $\vartheta_{\text{crit}} = 1.1$, a maximum area growth of $\vartheta_{\text{max}} = 4.0$, a growth exponent of $\gamma = 3.0$, and an adaptation speed of $1/\tau = 12$. We gradually pressurize the tissue expanders, $0.0 < t \leq 0.125$, then hold the pressure constant to allow the tissue to grow, $0.125 < t \leq 0.75$, and finally remove the pressure to visualize the grown area, $0.75 < t \leq 1.0$.

3.1 Tissue expansion in the scalp

Figures 7, 8, and 9 illustrate the tissue expansion process in the scalp. Figure 7 displays the temporal evolution of the normalized total area, elastic area, and growth area upon subsequent expander inflation, constant pressure, and expander removal. Once the elastic area stretch reaches the critical threshold of $\vartheta_{\text{crit}} = 1.1$,
slightly before the total pressure is applied, at \( t = 0.125 \), the tissue starts to grow. As the expander pressure is held constant, growth increases gradually causing the total area to increase as well. Then, at \( t = 0.75 \), the pressure is decreased to remove the expander. The elastic area retracts gradually, while the grown area remains constant. The vertical dashed lines correspond to the discrete time points, \( t = 0.225, t = 0.300, t = 0.375 \) and \( t = 0.750 \), displayed in Figure 8. Figure 8 illustrates the spatio-temporal evolution of area growth \( \vartheta^5 \). Growth is first initiated at the center of the expander, where the elastic stretch is largest. As growth spreads throughout the entire expanded area, the initial area of 50.4 cm\(^2\) increases gradually as the grown skin area increases from 70.07 cm\(^2\), to 84.25 cm\(^2\), to 95.73 cm\(^2\), and finally to 121.87 cm\(^2\), from left to right.

This indicates an area change of \( \pm 5\% \) giving rise to residual stresses, left. The area growth of \( 1.0 \leq \vartheta^5 \leq 3.5 \) shows that skin has more than doubled its initial area, right. This is in agreement with the final fractional area gain of \( 2.44 \), corresponding to an area growth in the scalp of 122.8 cm\(^2\).

### 3.2 Tissue expansion in the forehead

Figures 10, 11, and 12 summarize the tissue expansion process in the forehead. Figure 10 displays the temporal evolution of the normalized total area, elastic area, and growth area upon gradual expander inflation, constant pressure, and gradual expander removal.

Similar to the expansion in the scalp, growth begins at stretches beyond the critical threshold level, then increases gradually upon constant pressure, and remains constant upon expander removal. Figure 11 illustrates the spatio-temporal evolution of area growth \( \vartheta^5 \) at four characteristic time points indicated through the vertical dashed lines in figure 10. The growth process starts in the center of the forehead and spreads out throughout the entire forehead area. As it does, the initial area of 48.5 cm\(^2\) increases gradually as the grown skin area increases from 66.56 cm\(^2\), to 76.54 cm\(^2\), to 85.96 cm\(^2\), and finally to 116.55 cm\(^2\), from left to right. Figure 12 displays the remaining deformation upon expander removal. The final fractional area gain during forehead expansion is 2.44, corresponding to an area growth of 118.1 cm\(^2\).

### 3.3 Tissue expansion in the cheek

Figures 13, 14, and 15 document the tissue expansion process in the cheek. Figure 13 summarizes the temporal evolution of the normalized total area, elastic area,
and growth area upon gradual expander inflation, constant pressure, and gradual expander removal. Again, the growth process is initiated once the stretches reach the critical threshold of $\vartheta_{\text{crit}} = 1.1$. Upon constant pressure, growth increases gradually. Upon pressure removal, growth remains constant displaying the irreversible nature of the growth process. Figure 14 illustrates the spatio-temporal evolution of area growth $\vartheta$ in the cheek. Again, growth begins in center of cheek, where the elastic area stretch is largest. As the growth process spreads out throughout the entire cheek area, the initial area of 29.3 cm$^2$ increases gradually as the grown skin area increases from 42.74 cm$^2$, to 52.03 cm$^2$, to 59.39 cm$^2$, and finally to 76.86 cm$^2$, from left to right. Figure 15 summarizes the outcome of the expansion in the cheek with a final fractional area gain of 2.64, corresponding to an area growth of 77.4 cm$^2$.

4 Discussion

Tissue expansion is one of the basic treatment modalities in modern reconstructive surgery. Inducing controlled tissue growth through well-defined overstretch, it creates skin that matches the color, texture, hair bearance, and thickness of the surrounding healthy skin, while minimizing scars and risk of rejection [21]. Despite its widespread use, the choice of the appropriate tissue expander is almost exclusively based on the surgeon’s personal preference, and the discrepancy between recommended shapes, sizes, and volumes remains enormous [36]. The current gold standard for expander selection is to predict tissue growth by calculating the difference between the inflated and non-inflated expander surface [13, 53]. From an engineering point of view, it is quite intuitive, that this purely kinematic approach severely overestimates the net gain in surface area [60]. With a discrepancy of up to a factor four, these models assume that the entire deformation can be attributed to irreversible growth, completely neglecting the elastic deformation, which is reversible upon expander removal [36]. In an attempt
to account for this error, empirical correction factors of 6.00, 3.75, and 4.50 have been proposed for circular, rectangular, and crescent-shaped expanders [60]. This demonstrates the vital need to rationalize criteria for a standardized device selection.

Motivated by a first study on axisymmetric skin growth [55], we have recently established a prototype model for growing membranes to simulate tissue expansion in a general three-dimensional setting [8]. We have applied our model to quantitatively compare four commonly available tissue expander geometries, round, square, rectangular, and crescent [9], however, only on initially flat geometries. Here, for the first time, we demonstrate the potential of the model during tissue expansion in pediatric forehead reconstruction using a real patient-specific model. To embed the solution into a nonlinear finite element environment, we discretize the governing equations for in-plane area growth in time and space. To solve the nonlinear set of equations, we apply an incremental iterative Newton-Raphson solution strategy based on the consistent algorithmic linearization. The resulting algorithm is remarkably efficient, stable, and robust. It is capable of predicting tissue expander inflation, tissue growth, and expander deflation at different locations of a human skull within the order of minutes on a standard laptop computer. Because of its geometric flexibility, our general algorithm could also be adapted to predict tissue expansion in the trunk [4] or in the upper and lower extremities [20]. Although the proposed model for skin growth represents a significant advancement over the axisymmetric growth

Fig. 10 Tissue expansion in the forehead. Temporal evolution of normalized total area, elastic area, and growth area upon gradual expander inflation, $0.0 < t \leq 0.125$, constant pressure $0.125 < t \leq 0.75$, and deflation $0.75 < t \leq 1.0$. The final fractional area gain is 2.44, corresponding to 118.1 cm$^2$. Vertical dashed lines correspond to the time points displayed in Figure 11.

Fig. 12 Tissue expansion in the forehead. Remaining deformation upon expander removal. The elastic area strain of $0.95 \leq \varphi^e \leq 1.05$ indicates an area change of $\pm 5\%$ giving rise to residual stresses (left). The area growth of $1.0 \leq \varphi^g \leq 3.5$ shows that skin has more than doubled its initial area (right). The final fractional area gain is 2.44, corresponding to 118.1 cm$^2$.

Fig. 13 Tissue expansion in the cheek. Temporal evolution of normalized total area, elastic area, and growth area upon gradual expander inflation, $0.0 < t \leq 0.125$, constant pressure $0.125 < t \leq 0.75$, and deflation $0.75 < t \leq 1.0$. The final fractional area gain is 2.64, corresponding to 77.4 cm$^2$. Vertical dashed lines correspond to the time points displayed in Figure 14.

Fig. 15 Tissue expansion in the cheek. Temporal evolution of normalized total area, elastic area, and growth area upon gradual expander inflation, $0.0 < t \leq 0.125$, constant pressure $0.125 < t \leq 0.75$, and deflation $0.75 < t \leq 1.0$. The final fractional area gain is 2.64, corresponding to 77.4 cm$^2$.
model previously proposed [55], we would like to point out that some limitations remain. First, motivated by experimental observations, which report normal cell differentiation upon tissue expansion [61], we have assumed that the material microstructure remains unaffected by the growth process, \( F^g = \sqrt{\nu^g} I + \left[ 1 - \sqrt{\nu^g} \right] n_0 \otimes n_0 \). Here, for the sake of simplicity, we have modeled this microstructure as isotropic and elastic. We have recently shown that it is straightforward combine our growth model with in-plane anisotropy, introduced through pronounced stiffness along Langer’s lines [9, 30]. It might also be interesting to elaborate out-of-plane anisotropy and model the different skin layers individually [39]. We have demonstrated how to model the growth process itself as anisotropic as well [16]. This could imply growth \( \vartheta || \) exclusively along specific microstructural directions such as Langer’s lines, \( F^g = I + [\vartheta || - 1] \nu^g_0 \otimes \nu^g_0 \), or major in-plane growth \( \vartheta \perp \) along Langer’s lines combined with minor in-plane growth \( \vartheta || \) orthogonal to Langer’s lines, \( F^g = \vartheta || \nu^g_0 \otimes \nu^g_0 + \vartheta \perp \nu^g_0 \otimes \nu^g_0 + n_0 \otimes n_0 \). Similarly, we could even introduce a progressive reorientation of the collagen network to allow for the material to align with the maximum principal strains [33, 42]. Ideally, the growth law would be tied to the underlying mechanobiology [11]. Comparative tissue histology of grown and ungrown tissue samples could help to identify the mechanisms that trigger skin growth to validate or, if necessary, refine our evolution equation (13) for the growth tensor.

Second, for the sake of simplicity, our finite element mesh consists of one single brick element with two integration points across the skin thickness. We have previously studied the sensitivity of growth with respect to thickness refinement using a higher resolution across the thickness [8, 9]. However, our results were rather insensitive to mesh refinement. This insensitivity might be explained by the fact that, upon expander inflation, the skin is almost in a pure membrane state. During deflation, however, we observe buckling associated with strain gradients across the skin thickness, which might play a critical role in the development of residual stresses. To explore these residual stresses further, we are currently refining our model utilizing a shell kinematics [47] with a higher resolution across the thickness direction. This will also allow us to simulate the individual skin layers [35, 54] and their interaction during the expansion process, which we believe to be a major source of residual stress in real tissue expansion cases [41, 62].

Third, for the sake of simplicity, we have modeled tissue expansion only implicitly through controlling the applied pressure. In real tissue expansion, the external control parameter is the expander volume [36]. This implies that our virtual tissue expansion displays creep under constant loading, while clinical tissue expansion might rather display relaxation under constant deformation [9], similar to our parameter study in Figure 4.

Fourth, here, we have assumed that the expander is connected tightly to the expanded tissue, neglecting effects of interface sliding and shear [55]. This seems to be a reasonable first assumption though, since most current expanders have well-designed textures to promote mild tissue in-growth, primarily to prevent expander migration [6]. To address these potential limitations, we are currently refining the elastic model, the growth model, and the boundary conditions, to render our future simulations more realistic.

Last, while our computational model seems well suited to provide qualitative guidelines and trends, at its present state, it is not recommended for quantitative statements. We will need to perform acute and chronic in vitro and in vivo experiments to truly calibrate the underlying material parameters, to potentially refine and fully validate our model, to eventually make it applicable for clinical practice. Nevertheless, we believe that using the equations on nonlinear continuum mechanics represents a significant advancement over the current gold standard to predict tissue growth exclusively in terms of kinematic quantities [53, 60].

Conclusion

We have presented a novel computational model to predict the chronic adaptation of thin biological membranes when stretched beyond their physiological limit. Here, to illustrate the features of this model, we have demonstrated its performance during tissue expansion in pediatric forehead reconstruction. We have quantified its capability to rationalize these parameters to create skin flaps ofstanding of the gradually evolving stress and strain fields in growing skin may help the surgeon to prevent tissue damage and optimize clinical process parameters such as expander geometry, expander size, expander placement, and inflation timing. Ultimately, through inverse modeling, computational tools like ours have the potential to rationalize these parameters to create skin flaps of desired size and shape. Overall, we believe that predictive computational modeling might open new avenues in reconstructive surgery and enhance treatment for patients with birth defects, burn injuries, or breast tumor removal.

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