Surface-bounded growth modeling applied to human mandibles

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Abstract—From a set of longitudinal three-dimensional scans of the same anatomical structure, we have accurately modeled the temporal shape and size changes using a linear shape model. On a total of 31 computed tomography scans of the mandible from six patients, 14851 semilandmarks are found automatically using shape features and a new algorithm called geometry-constrained diffusion. The semilandmarks are mapped into Procrustes space. Principal component analysis extracts a one-dimensional subspace, which is used to construct a linear growth model. The worst case mean modeling error in a cross validation study is 3.7 mm.

Index Terms—Geometry-constrained diffusion, morphometrics, nonrigid shape-preserving registration, principal component analysis, Procrustes analysis, semilandmarks.

I. INTRODUCTION

PEDiatric craniofacial surgeons need insight into expected facial growth. This paper is concerned with the mandible, a particularly complex bony structure both in its shape and in its growth process, as two sets of teeth erupt asynchronously while the direction of the condylar process changes by a considerable angle.

Our data set here comprises 31 mandibular surfaces extracted from computed tomography (CT) scans of a total of six children diagnosed with Apert syndrome.

The analysis falls under two major headings. First is the representation of the set of mandibular surfaces by one vector of 14851 points that can be considered to be semilandmarks: that is, points that do not have names, but that correspond across all the cases of a data set under a reasonable model of deformation from their common mean [1]. Second is the summary of these 31 point sets in a linear subspace of surprisingly low dimension, affording surprisingly accurate growth predictions.

The organization of the remainder of this text is as follows. Section II sets the stage for our algorithm by reviewing the literature of nonrigid registration by deformable models. Section III introduces the patients to which we have access, their CT images, and the mechanism by which we produced the 31 mandibular surfaces of the data set. Section IV describes the initialization of our diffeomorphism by detection and matching of crest lines, and Section V shows how we proceed to a full surface-constrained diffusion. In Sections VI and VII, the (31 − 1)-dimensional sample subspace of the full space of semilandmark shape is subjected to certain conventional multivariate biometric analyses that yield powerful predictors of unobserved (future) form. We assess their accuracy using measures of surface-to-surface discrepancy. Section VIII is a retrospect over all these tactics, emphasizing the surprising power of the diffusion methodology to uncover commonalities in the six independent growth processes of this sample. We close with a plea for corresponding studies of normative samples.

II. RELATED WORK

The literature treating registration methods is very extensive (e.g., [2] or [3] for surveys). This section, therefore, mainly concentrates on the literature covering both registration and deformable models (for reviews, e.g., [4] or [5]) or morphometrics (e.g., [6] and [7]). We will emphasize the registration method of Feldmar and Ayache [8], as it most resembles the geometry-constrained diffusion method [9], [10]. Feldmar and Ayache [8] perform a surface registration based on a distance measure that relies on local geometrical properties of the surfaces. The surface registration is a collection of local affine registrations, spatially blurred so as to result in a smoothly varying global registration. Geometry-constrained diffusion (Section V) does not make a collection of local affine frames, but a global registration field. We do not exploit any metric properties of the surfaces, but look for a globally simple registration field. This also creates a tendency to match points of similar geometry since the field, otherwise, must be more complex.

Deformable models have been widely studied [4], [5]. When using landmarks to drive the correspondence between objects one main drawback is the need for their manual location in advance. A lot of researchers have worked on this. Bookstein has reported a method where the semilandmarks are placed on contours automatically [1]. Note, a landmark is a point that can be identified by verbal characterization on the single case [11].
Fig. 1. The raw dataset of patient 3, scan #1 (1-mo scanning). The mandibular bone and teeth have low density, therefore, cavities and holes are introduced when segmented.

Fig. 2. The crest lines (in red) and \( k \)-max lines (in blue) on the three Gaussian smoothed (kernel size, \( \sigma = 3 \) mm) mandibles at (left) 9 mo, (middle) 21 mo, and (right) 7-yr old. The mandibles are from patient 6, scan #1, #2, and #3, respectively. The surfaces are translucent.

Semilandmarks are points that do not have names, but correspond across all the cases as images of the same point of their average, so one can carry out statistics just as if they were landmark points [1].

Fleute and Lavallée [12] extrapolate a small number of range data to obtain a complete surface representation. Principal component analysis (PCA) is used to reduce the dimensionality. Data sets are registered together using an elastic registration method of Szeliski and Lavallée [13] based on octree-splines. The method is a least squares minimization of the distances between a sparse and unorganized set of points and a dense set used to build a three–dimensional (3-D) octree-spline distance map. The result is a smooth deformation field.

A registration technique based on thin-plate splines that takes landmark errors into account has been reported in [14]. The semilandmarks are located semiautomatically or manually.

The present work has been greatly inspired by the seminal work of Cootes and colleagues ([15] for an overview) and Dean et al. [16] (see below). Manually detected landmarks have been used for Cootes and colleagues’ analysis [15]. Principal components are calculated from the Procrustes analysis and an active shape model is made. The deformation of the active shape model is restricted by the principal components. For segmentation, gray-level information near the object boundaries is also modeled.

Kelemen et al. [17] have used the same method as in [15], but in order to automate the landmark generation, Fourier-descriptors [18], [19] were found very powerful. Restrictions on the topology of the surfaces are the main drawback when using Fourier descriptors.

In the present paper, the registration method of Subsol et al. [20] gives the initial object correspondence (Section IV). Crest lines [21] are registered together taking into account the constraints inferred by lines and a heuristic algorithm based on the iterative closest-point (ICP) algorithm [22].

Grenander and Miller [23] have formalized a model of anatomy in which anatomies are represented as deformable templates, collections of zero-, one-, two-, and three-dimensional manifolds. They have three principal themes in computational anatomy: large deformations maps [24], empirical probability laws that represent anatomical variation in populations, and inferences on populations, such as disease classification.

Lorenz and Krahnstöver [25] present a 3-D shape model based on surface points. The overall strategy is similar to that in the present paper. However, the initial correspondence is based on a manually defined set of landmarks on each object. From the sparse mapping defined by the landmarks, a dense mapping is calculated by thin-plate interpolation. If it happens that the deformed mesh is folded, it is unfolded using a mesh relaxation balancing bending force against surface mesh force by a mass-spring model.

Dean et al. [16] analyze pairs of frontal and lateral X-rays from 32 individuals (16 males and 16 females) aged from 3- to 18-yr old. The objective is to investigate which 3-D landmarks could be collected with high precision, to identify ontogenetic trends in landmark configuration shape change, and to detect patterns of sexual dimorphism. The 32 landmarks are...
transformed to points in two Procrustes shape spaces (one for each gender). Relative warps are then used to search for trends in ontogenetic shape change. It is interesting that the result of the analysis is almost identical to the present work even though the 32 landmarks are placed mostly off the mandible, and the subjects are normal children.

We are not aware of any other growth studies in which longitudinal 3-D acquisitions from humans have been used for growth modeling except in [26]–[28], where the growth was modeled for the subject who is patient 6 here. Subsol [26] modeled the craniofacial growth using a linear model between a set of the controlling points. Bro-Nielsen et al. [27] used a nonrigid registration method to model the growth of the mandible. The method was a surface interpolation that did not preserve the mandibular shape; e.g., the condyles disappeared for intermediate interpolated time instances. Andresen et al. [28] used the same object registration technique as in Section IV to register the mandibles. A second-order polynomial was used to interpolate the longitudinal displacement. Unfortunately, because the polynomial model there was not well-suited to mandibular growth modeling, the mandible was forecast to shrink in the course of growth!

### III. DATA MATERIAL

The data consisted of CT scans of six subjects with Apert syndrome, four males and two females. Patient 6 is Danish, the
TABLE II
PRINCIPAL COMPONENT ANALYSIS OF THE SEMILANDMARK DATA

<table>
<thead>
<tr>
<th>$\sigma = 3$, $k = 9087$</th>
<th>$\sigma = 1$, $k = 14851$</th>
</tr>
</thead>
<tbody>
<tr>
<td>no diffusion</td>
<td>diffusion</td>
</tr>
<tr>
<td>$\lambda_1 \cdot 10^5$</td>
<td>$\lambda_1 \cdot 10^5$</td>
</tr>
<tr>
<td>$\delta_{\text{var}}$</td>
<td>$\delta_{\text{var}}$</td>
</tr>
<tr>
<td>0.3134</td>
<td>0.3107</td>
</tr>
<tr>
<td>0.0496</td>
<td>0.0294</td>
</tr>
<tr>
<td>0.0415</td>
<td>0.0233</td>
</tr>
<tr>
<td>0.0373</td>
<td>0.0197</td>
</tr>
</tbody>
</table>

Fig. 9. Left and right images show the deformed (green) and the original (red) crest lines before and after applying the diffusion algorithm ($\sigma_{\text{NI}} = 2$ mm). In the initial registration, crest lines are registered with crest lines. Where the topology does not change and away from umbilic points (for which the curvature is the same in all directions, e.g., at the condyles), (almost) no movement of the green crest lines is seen. Because erupting teeth change the topology on the upper surface (cf. Fig. 6), the green crest lines have moved in that region.

Fig. 10. The deformation vectors are moved too far away from the surface (the value of $\sigma_{\text{NI}}$ is too high) resulting in a wrong projection back onto the surface.

Fig. 11. Scatterplots of the first and second principal component for the six cases (left) before and (right) after the geometric-constrained diffusion is applied. For visual clarity, each patient’s mean is restored. Numbers are ages in months. It is clear that the six growth trajectories span an angular range of nearly 90° between the earliest growth segments of patient 2 (dashed line) and 3 (dotted line) in this projection; after diffusion (right), the alignment is far tighter.

TABLE III
CROSS-VALIDATION OF SHAPE PC1

<table>
<thead>
<tr>
<th>Patient excluded</th>
<th>Percentage variance explained in model ($= 100% \cdot \lambda_1 / (\sum \lambda_i)$)</th>
<th>$\alpha$</th>
</tr>
</thead>
<tbody>
<tr>
<td>none</td>
<td>62.2</td>
<td>0.0</td>
</tr>
<tr>
<td>1</td>
<td>64.5</td>
<td>4.1</td>
</tr>
<tr>
<td>2</td>
<td>65.8</td>
<td>5.8</td>
</tr>
<tr>
<td>3</td>
<td>62.1</td>
<td>6.5</td>
</tr>
<tr>
<td>4</td>
<td>62.3</td>
<td>8.3</td>
</tr>
<tr>
<td>5</td>
<td>60.8</td>
<td>5.9</td>
</tr>
<tr>
<td>6</td>
<td>61.1</td>
<td>3.9</td>
</tr>
</tbody>
</table>

A previous two-dimensional roentgencephalometric study has shown that the mandible in Apert syndrome has relatively normal morphology except for some adaptive changes [29]. As these subjects are prepubescent and the sample is small, we have pooled the two sexes in the analysis to follow.

The mandibles were extracted from the CT scans by thresholding, with manual segmentation around the condyles. Holes inside the object were filled and the mandibles trilinearly resliced to 0.5-mm cubic voxels (Fig. 1). The reference mandible from which we propagated all the semilandmark points in this study is the 56-mo scan for patient 2; the semilandmarks were chosen to be the vertices of the triangulated mandibular surface.

IV. REGISTRATION: OBJECT CORRESPONDENCE

In order to establish object correspondence, we search for features that match areas with equivalent morphology. Since the topology is not changed dramatically for the mandible when growing, features reflecting the “stable” geometry are used. The local shape of a surface is totally characterized by the principal curvatures $k_1, k_2$ ($k_1 > k_2$) and their derivatives in the coordinate system defined by the principal directions $(t_1, t_2)$ [30].
We work with the extraction and matching of ridge lines [21], [28], [31]. Ridge lines come in four type types, corresponding to maximum or minimum Gaussian curvature \( k_1 \) and \( k_2 \). Here, we use the maximal \( k_1 \) type and the maximal \( k_2 \) type, called “crest lines” and “k2-max lines,” respectively. For three mandibles, these are shown in Fig. 2.

The overall framework follows the ideas originally proposed in [20] and already used in [28]. First, the crest lines and k2-max lines for each dataset are extracted at a high scale, in order to get the more global features (Fig. 2). The crest lines are registered (Fig. 3) to provide a sparse initial deformation field. A dense field is then calculated by adaptive Gaussian filtering [32] (Section IV-A). The k2-max lines are then deformed according to the initial dense deformation field and subsequently registered. From the two sets of matches (one from the crest lines, the other from the k2-max lines) a combined deformation field is calculated. This sparse field is also interpolated to a dense field by adaptive Gaussian filtering. The resulting sparse deformation field is used as an initial guess in the point matching algorithm in Section V. Another iteration is done at a fine scale, but using the dense high-scale deformation field as initial displacement for the crest lines at fine scale.

This extremal mesh is manipulated in the course of registration first by a second-order moment registration to the target form (Section IV-B); then by two first-order polynomial deformations and two second-order polynomial deformations (Section IV-C); finally, by an unconstrained nonrigid deformation for all points on all lines (Section IV-D). Details of these steps are as follows.

### A. Adaptive Gaussian Filtering

Gaussian filtering equals Gaussian smoothing for unevenly distributed points with a renormalization so that the total filter weight becomes unity. Adaptive Gaussian filtering [32] equals Gaussian filtering, but the scale parameter (the size of the Gaussian kernel) equals the square root of the distance to the nearest point.

### B. Second-Order Moment Registration

The description in this section is adopted from [33]. The dispersion matrix for the mesh is given by

\[
D = \frac{1}{L} \sum_{i=1}^{L} (\mathbf{q}_i - \mathbf{q}) (\mathbf{q}_i - \mathbf{q})^T \tag{1}
\]

where

- \( \mathbf{q}_i \) three–dimensional points in the mesh;
- \( \mathbf{q} \) center of mass for the points;
- \( L \) number of points;
- \( (\cdot)^T \) transpose.

\( D \) is factorized by determining the eigensolution of the matrix.

\[
D = \mathbf{P} \Lambda^{1/2} (\mathbf{P} \Lambda^{1/2})^T \tag{2}
\]

where \( \mathbf{P} \) is the orthonormal matrix of eigenvectors and \( \Lambda \) is a diagonal matrix containing the eigenvectors.

The affine transformation \( (\mathbf{q}_T = \mathbf{R} \mathbf{q}_T + \mathbf{t}, \mathbf{R} \) and \( \mathbf{t} \) are the rotation matrix and translation vector, \( \mathbf{q}_T \) is the transformed set) that gives set \( T \) the same first and second-order matrix as set \( S \) is determined by

\[
\mathbf{R} = \mathbf{P} \Sigma^{-1/2} \Lambda_T^{-1/2} \mathbf{P}_T^{-1}, \quad \mathbf{t} = \mathbf{q}_S - \mathbf{R} \mathbf{q}_T. \tag{3}
\]

### C. \( v \)-th Order Polynomial Deformation, \( v = 1, 2 \)

Having the point correspondence between the two sets \( S \) and \( T \) (Section IV-E), the \( v \)-th order polynomial deformation that optimally—in a least square sense—maps \( T \) to \( S \) is given by

\[
\langle \hat{\mathbf{q}} \rangle_T = \sum_{i} \sum_{j} \sum_{k} \alpha_{ij,k} \mathbf{x}_{i} \mathbf{y}_{j} \mathbf{z}_{k}, \quad i + j + k \leq v \tag{5}
\]

where \( \hat{\mathbf{q}} \) is the mapped dataset and \( \langle \cdot \rangle \) is one of the \( x, y, \) or \( z \) coordinates. The parameters \( \alpha_{ij,k} \) are determined from the correspondence between the two sets by least squares [34].

### D. Totally Nonrigid Deformation

All the points are freely moved to their corresponding match in the other set (Section IV-E).

### E. Making the Point Correspondence

The correspondence between meshes is determined by the ICP algorithm [22] modified as in [20] to take the line constraints into account.

### V. REGISTRATION: POINT CORRESPONDENCE

From the sparse deformation field (the mapping between the two extremal meshes), derived from Section IV, a dense field must be calculated as we wish to register all the points on the whole surface, not only the extremal mesh. The points are the vertices of the triangulated mandibular surface. These points will be assumed to make up the semilandmarks after the geometry–constrained diffusion algorithm has been applied.

Adaptive Gaussian filtering can fold the mesh registration. To regularize the dense field, including simplifying (unfolding) it, we adopt the geometry-constrained diffusion algorithm originally published in [9] and [10].

The algorithm (Fig. 4) alternates Gaussian smoothing of the Cartesian components of displacement independently, which pulls them away from the surface, with projection back to the closest point on the surface. The detailed flow is as in Fig. 5; a typical example is set out in Figs. 6–8. Fig. 9 shows the improved registration achieved by the diffusion algorithm. In all cases, the “initial displacement field” is the dense deformation field (interpolated from the extremal mesh onto the whole surface) constructed in Section IV.

In the diffusion step, for each Cartesian component of the displacement field, a Gaussian weighted average is constructed. The standard deviation of the Gaussian \( \sigma_D \) is the only parameter in the numerical scheme. In effect, it regulates the “time steps” between projections onto the surface. For simple surfaces, it may be large. For surfaces including regions of high
The final surface registration is not sensitive to the value of $\sigma_D$, as long as the surface is not torn apart (Fig. 10). The influence of $\sigma_D$ is investigated further in [10].

For matching, as in [35], we use a kD-tree [36] for finding the closest point on the target surface. As reference points on the triangulated target surface we use the center of mass (CM) points of its triangles. The closest CM point is located (by the kD-tree) and then the closest actual surface point is estimated as the intersection of the triangle’s plane with the line through the reference point parallel to the target normal. If this intersection is outside the target triangle, its neighbors are searched. The proposed algorithm has the advantage that the points can move continuously on the target surface.

The diffusion is stopped when

$$\sum_{p_i} ||D_n(p_i) - D_{n-1}(p_i)||^2 < \epsilon$$

where

- $p_i$: points on the source surface;
- $D_n$: displacement in the $n$th-iteration;
- $\epsilon$: user-chosen parameter.

Ten iterations are normally sufficient.

VI. STATISTICAL DESCRIPTION: GEOMETRIC MORPHOMETRIC ANALYSIS

In Section IV and Section V, our aim was to register the points on the 31 mandibles in such a way that we could assume that the points were semilandmarks. On the reference mandible the semilandmarks were chosen to be the vertices of the triangulated mandibular surface. The Gaussian-smoothed reference mandible has 14,851 and 9087 vertices for the kernel sizes $\sigma_D = 1$ and 3 mm, respectively. These meshes are deformed onto all other mandibles using the simplified dense deformation field set out in Section V. The semilandmarks that result are subjected...
Fig. 16. Modeled last scan (scan #5) based on the first scan (scan #1) for patient 1. The large errors are located at the cavities as described in the text. The left images being the observed scan seen from two different viewpoints. The right images are the modeled mandible seen from the same two viewpoints. The surface is rainbow color coded from blue (0 mm) to red (10 mm) with the error (Fig. 23), calculated as the Euclidean distance between the same semilandmarks in the observed and modeled scan. The mean error is 2.4 mm.

Fig. 17. Modeled last scan (scan #7) based on the first scan (scan #1) for patient 2. The large errors are located at the cavities introduced by the segmentation of the 1-mo-old scan. The mean error is 3.5 mm. The staircases on the observed scan #7 is due to errors in the original volume.

Fig. 18. Modeled last scan (scan #5) based on the first scan (scan #1) for patient 3. The mean error is 2.8 mm. By looking at the first scan (Fig. 1) it is not surprising to find the large errors at the cavities. The notch in the side of the modeled mandible is related to the hole found at the same place on the first scan.

Fig. 19. Modeled last scan (scan #7) based on the first scan (scan #1) for patient 4. The mean error is 3.7 mm. The quality of the model is surprising. None of the scans in the model is as old as scan #7 (12-yr old). The oldest scan in the model is 7-yr old (patient 6, scan #3). This indicates the stability of the growth. The red area shows a missing tooth, which has been extracted. The prominent chin is not modeled very well. Using older scans solves the problem (Figs. 24 and 25).

to conventional multivariate biometric analyses that yield powerful predictors of unobserved (future) form.

At the outset, the configurations of 14,851 semilandmarks, specimen by specimen, are aligned in a common coordinate framework by Procrustes analysis [7], [37], which is a least-squares superposition in which position, orientation, and scale are jointly standardized. We used the usual iterative approach to produce an average shape \( \bar{X} \) to which the specimens are all aligned. Let \( X_i \) be the \( k \times 3 \) matrix of 3-D semilandmarks for the \( i \)th specimen.

1) The center of mass (the centroid) for each shape is translated to the origin.
2) For each specimen, centroid size (CS) is computed, square root of the the summed squared distance of all semilandmarks from this centroid, and the form is rescaled so that this sum of squares becomes 1.0.
3) Make an initial guess at \( \bar{X} \), for instance, \( X_1 \).

4) Orient each shape to this tentative mean. To do so, let \( UDV^T \) be the singular value decomposition of \( X_i^T X_i \). As long as all elements of \( D \) are positive (which will be the case in all realistic applications), the rotation that superimposes \( X_i \) on \( \bar{X} \) is the \( 3 \times 3 \) matrix \( VU^T \).
5) Re-estimate the mean shape as the average of these rotated configurations.
6) Return to step 4) until convergence, which is invariably after two or three iterations.

Now that all the shapes are aligned, we calculate the patient mean shapes for each patient

\[
\bar{X}_i = \frac{1}{n_i} \sum_{j=1}^{n_i} X_{i,j}
\]

(7)
where \( n_i \) is the count of scans for the \( i \)th patient. Write \( \bar{X}_{pi,j} \) for the patient-specific deviation from his/her own average

\[
\bar{X}_{pi,j} = X_{pi,j} - \bar{X}_i.
\]

Rearrange each \( \bar{X}_{pi,j} \) into a column vector of length \( 3k \), where \( k \) is the count of semilandmarks (9087 or 14851, depending on the computation), and concatenate all of these column vectors into a new data matrix \( \bar{X}_{3k \times 31} \).

The covariance matrix of the patient-specific residual coordinates is \( S = \bar{X}\bar{X}^T/30, \ 3k \times 3k \). By the Eckart-Young’s theorem [38], we can extract its eigenvectors very simply from the corresponding eigenvectors of \( T = \bar{X}\bar{X}^T/30 \), which is \( 31 \times 31 \).

Let \( \bar{e}_p \) be the eigenvector of \( T \) with corresponding eigenvalue \( \lambda_p \), sorted in descending order. It follows from [38] that the 31 vectors

\[
e_p = \frac{1}{\sqrt{\lambda_p}}\bar{X}_{pi,}\bar{e}_p
\]

are all eigenvectors of \( S \) with corresponding eigenvalues \( \lambda_p \). The last \( 3k - 31 \) eigenvectors are all zero.

VII. SHAPE EVALUATION AND PREDICTION

Table II shows some results of these PCAs. There are two comparisons here: the effect of the surface-bounded diffusion algorithm (Section V), expressed in the difference of the two conditions for \( \sigma = 3 \), and the effect of reducing the spatial scale \( \sigma \) of the smoothing, observable in the relation of the diffusion with \( \sigma = 3 \) to that with \( \sigma = 1 \). For each set of 31 semilandmark configurations, the table shows the first four eigenvalues and the percentages of total Procrustes variation for which they account. The diffusion greatly strengthens the systematic aspects of the analysis; for instance, the shape variation along the first eigenvector increases by a factor of one and a half. Decreasing the size of the Gaussian smoothing radius \( \sigma \), conversely, results in local differential changes that are less likely to be aligned across these 31 mandibles and, thus, a decrease of the explanatory power of these same eigenvectors.

Any summary of an ostensibly homogeneous sample of biological material, in this case, observed growth in six children...
with the same diagnosis of synostosis, is persuasive to the extent that that categorization "explains" the quantifications: the extent to which the extracted measurements are homogeneous over the class. From Fig. 11, it is clear that the six growth trajectories span an angular range of nearly 90° between the earliest growth segments of patients 2 and 3 in this projection; after diffusion, the alignment is far tighter. As shown in Table II, there is far less variation around the common direction of these growth trajectories after diffusion than before (68% vs. 45%).

From the right image in Fig. 11, it appears that the three to seven forms of each case lie reasonably close to a line in Procrustes space. We can, thus, summarize each by its own first principal component [16], and then examine the resulting six vectors to see how they may be ordinated.

Fig. 12 indicates the distribution of these six vectors in the 6-D subspace they span. Each principal component is normalized for its actual Procrustes variance explained, and the presentation here preserves that metric. We see a strong central tendency in these six growth vectors, closest to the observed trend for patient 3 or patient 4. The variation of the tips of these six vectors seems to cover about the same diameter in the six different directions plotted here, without apparent correlation in the planes selected for plotting, and so it is reasonable to imagine it to be spherical, i.e., without a preferred directional structure within this space.

We demonstrate the power of this procedure by a series of growth predictions that, in all cases, predict the oldest form for each of the six patients by altering each of the earlier forms for that patient according to the regression on CS and the first principal component (PC1) given by the other five patients.

The model is built as described above except that one patient at a time, m=1, ... , 6, is excluded from the analysis. We will attend to the eigenvectors e1,m that arise as first eigenvectors of the data set excluding patient m’s scans. As seen in Table III the growth models prove highly stable: fractions of Procrustes variance explained by the six eigenvectors e1,m range from 60.8% to 65.8% only, and the angle that these eigenvectors make with the pooled e1 is limited to no more than 8° (a cosine of 0.99). No patient controls the variability of the pooled analysis. Other computations, not shown here, confirm the observation, first set out in [16], that removing variation of patients’ mean shape greatly enhances the reliability of the resulting PCA analysis of growth per se. Because patient m’s scans are already aligned with the grand mean shape, a new shape may be modeled by

$$X_{\text{future}} = X_\ell + t \cdot e_{1,m} , \quad t \in \mathcal{R}$$

(10)

where \(t\) is a growth index parameter and \(X_\ell\) is one of patient m’s earlier forms. If the new form is already available, the value of \(t\) can be found by projection of the form onto the line of slope \(e_{1,m}\) through the earlier form. In a context of prediction instead, we can estimate an appropriate value of \(t\) by regression using age or its transforms, CS, or mandibular length. These lead to equivalently accurate predictions (Figs. 13–15); we will use the one calibrated by CS, as that is the commonest in other applications of these Procrustes coordinates. Fig. 13 shows the clear relationship between CS and PC1 for the complete data set of all 31 mandibles. To evaluate this prediction, of course we need to use the regression coefficient \(\alpha\) based on the data set omitting the \(m\)th case, the one we are predicting. The final model is, therefore

$$X_{\text{future}} = X_\ell + (CS_{\text{future}} - CS_{\ell}) \cdot \alpha \cdot e_{1,m} .$$

(11)

From (11), it is seen that only one scan of the patient and the future CS or age (Fig. 15) is needed to make the prediction.

Table IV shows mean, standard deviation, and root-mean-squared errors of surface prediction when each patient’s last scan is predicted by each of his or her earlier scans at the appropriate CS, using predictions of size-related shape change based on the other five patients. In general, accuracy increases with decrease of age interval, but the prediction errors from earliest to latest scan seem gratifyingly homogeneous, in keeping with the evident parallelism of individual growth curves in Fig. 11.

Some of the predicted mandibles are shown in Figs. 16–21. The histograms of the errors for the figures are shown in Fig. 22. The left images in Figs. 16–21 represent the last observed scan for each patient. The right images show the modeled mandibles. The rainbow color coding shows the errors (Fig. 23). Error is the unsigned Euclidean distance between matched semilandmarks when the modeled surface is rescaled to match the observed scan in CS.

When segmenting the scans at 1–3 mo of age, large cavities are erroneously introduced because of the very low X-ray absorption (Fig. 1). As seen from Fig. 18, these errors are not removed by the growth model, as expected. For that reason, the largest errors (>7 mm) are seen at the cavities. Also, removal of a tooth is obviously not modeled, as seen in Fig. 19. Besides these very specific errors, we do not see errors larger than approximately 5 mm, except for patient 4, scan #7 (Fig. 19). This
Fig. 24. Predicted shape of the most recent scan of patient 4 using each of her earlier scans, from top left to bottom right. The problem of cavities occurs only for the prediction from the earliest surface, and the accuracy of the prediction increases as the prediction interval becomes shorter. Notice, the model itself—the first eigenvector in (11)—stays the same.

patient, unlike the others, has a prominent chin, about which the prediction has no information.

The ability to model patient 4, scan #7 accurately is a bit surprising. None of the other patients have been scanned at that age, the oldest scan (when patient 4 is excluded) being 7-yr old (patient 6, scan #3), but patient 4, scan #7 is 12-yr old. This also indicates that the growth is modeled correct.

Figs. 24–25 are included to show how the errors decrease as successively older scans are used. Notice, how small the errors are when patient 4, scan #6 (11-yr old) is used as the basis for the model. 12 715 of the 14 851 semilandmarks (85.6%) have an error less than 3 mm. Fig. 26 shows the histograms.

VIII. CONCLUSION

We have presented a linear 3-D growth model that relies on principal components of Procrustes shape coordinates. The model, which is formally independent of mandibular form per se [seen by subtracting $x_c$ from (11)], is able to predict mandibular shape changes to an acceptable accuracy. The great increase in percent of Procrustes variance explained by the first eigenvector after geometry-constrained diffusion (Table II) is surprising and deserves further investigation. Inclusion of zero-dimensional and one-dimensional constraints (landmark points and curves) would further improve this step in the processing.

Other studies, such as [40], show that the growth of the mandible is nonlinear, when using a “biological coordinate system.” The present study does not reject that hypothesis, but indicates that the growth is linear if modeled in Procrustes space. This also complies very well with the result of [16]. A combination of the two “frames” might be very fruitful. Procrustes space could be used for the growth modeling and
the “biological space” for the visualization. The ability to extrapolate the growth period by 5 yr strengthens our hypothesis about linear growth.

At present, we can only obtain the closely spaces CT scans that allow this kind of analysis from clinical cases with various types of craniofacial growth disturbances. As mentioned above, in Apert syndrome, the mandible is not affected by the primary anomaly [29]. Other craniofacial syndromes also show fairly normal mandibular development e.g., Crouzon syndrome and achondroplasia. It will be interesting to ascertain which of these groups are characterized by these same mean growth trajectories. If the predictive accuracy we have demonstrated here extends to other syndromes lacking a primary mandibular dysmorphology, one might then speculate that it characterizes the normal growth process as well.

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REFERENCES


