

# Volume-Preserving Free-Form Solids

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Abstract: Some important trends in geometric modeling are the reliance on solid models rather than surfacebased models and the enhancement of the expressive power of models, by using free-form objects in addition to the usual geometric primitives and by incorporating physical principles. An additional trend is the emphasis on interactive performance.

In this paper we integrate all of these requirements in a single geometric primitive by endowing the tri-variate tensor product free-form solid with several important physical properties, including volume and internal deformation energy. Volume preservation is of benefit in several application areas of geometric modeling, including computer animation, industrial design and mechanical engineering. However, previous physics-based methods, which usually have used some forms of 'energy', have neglected the issue of volume (or area) preservation.

We present a novel method for modeling an object composed of several tensor-product solids while preserving the desired volume of each primitive and ensuring high-order continuity constraints between the primitives. The method utilizes the Uzawa algorithm for non-linear optimization, with objective functions based on deformation energy or least squares.

We show how the algorithm can be used in an interactive environment by relaxing exactness requirements while the user interactively manipulates free-form solid primitives. On current workstations, the algorithm runs in real-time for tri-quadratic volumes and close to realtime for tri-cubic volumes.

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# 1 Introduction

Modern geometric modeling emphasizes solid models rather than surface-based models, usage of free-form objects in addition to the usual geometric primitives, incorporation of physical principles, and interactive performance. In this paper we integrate these four issues in a single setting by endowing the tri-variate tensor product Bézier free-form solid with physical properties.

#### 1.1 Background

The common approach to representing and manipulating free-form objects is by using a boundary representation (Brep), with parametric surfaces for the boundary. Adjacencies between neighboring surface patches are stored explicitly. Using a Brep, it is inherently difficult to model physical attributes associated with the object. Such attributes are easier to consider when using parametric free-form *solids* instead of surfaces. The difference between the two is the dimension of the parameter space (two for surfaces and three for solids.)

Some previous systems have used free-form solids (e.g. [Farouki85].) However, parametric volumes are usually not used in the way that surfaces are used, for direct object design, but rather for design of separate deformation entities used for modification of existing objects. This can be explained by the fact that if only the boundary of the object is of interest, there is no need to use free-form solids, which enable control over what happens 'inside' the object.

Free-form deformations (FFD) were introduced in [Sederberg86] as a technique for defining a smooth deformation on a space including the objects embedded within that space, regardless of their geometric representation. FFD utilizes a tri-variate tensor-product parametric Bézier solid defined by a lattice of control points. The defining parameter space is the unit cube. To deform an object point, its local coordinates inside the unit cube are computed. Then the image of the point under the deformation is computed using the Bézier control points and basis functions. Naturally, other basis functions (such as NURBS) could be used as well [Griessmair89].

Sederberg suggested a user interface based on control point manipulation, with which is it rather difficult and tedious to obtain a desired deformation. Direct manipulation of object points instead of control point manipulation was suggested in [Borrel91, Hsu92]. The user directly moves an object point, and the system automatically computes the control point configuration yielding the desired point displacement constraints. [Rappoport94] extends this method to approximate ('probabilistic') point constraints with a non-isotropic shape parameter. [Joy91] gave methods to manipulate a group of control points in a single operation. A more general type of extension to FFD was presented in [Coquillart91], who defined an arbitrary volume and used numerical routines to compute local coordinates within this volume. Neither of the above methods attaches any physical meaning of the deformation. Simple constrained deformations were described in [Borrel94].

Physics-based modeling is a successful research area in geometric modeling. Several papers [Terzopoulos94, Welch92, Kallay93, Moreton92, Celniker91, Greiner93] presented surface design schemes based on minimization of an energy functional subject to linear point constraints such as location and tangent vectors. We are not aware of any work using similar ideas for free-form solids. Other applications of physics-based modeling are in reconstruction and tracking [Fang92], motion control [Shapiro88], and modeling of flexible and rigid objects [Barzel88].

The only relevant reference we are aware of for volume preservation<sup>1</sup> is [Aumann92], which gives an algorithm that approximates a surface of revolution by a surface which is not a surface of revolution while trying to preserve the original volume. Free-form solids are not discussed, and it seems that the algorithm is not suited for them at all. Formulae for computing the area or volume enclosed by curves and surface patches were given in [Elber94, Liu87].

### 1.2 Proposed Approach

We use free-form solids as design primitives. In the context of solid model design in general and specifically of free-form solids, one of the most basic physical properties of a space cell is its volume size. A major drawback of current user interaction techniques when applied to free-form solid design is that the user has no way of controlling the contained volume size. Currently, solid design (as opposed to using volumes for free-form deformations) is not much more than design of the surfaces bounding the volume, each of them independently. We present a novel method for modeling an object composed of several tensor-product solids while preserving the desired volume of each primitive and ensuring high-order continuity constraints (and any linear constraints in the control points) between the primitives. The method utilizes the Uzawa algorithm for non-linear optimization, with an objective function based on deformation energy or least squares (LSQ).

The algorithm is very useful for several applications. For example, hierarchical FFDs were used by [Chadwick89] for computer animation of muscles. A similar effect could be achieved by a combination of point displacement constraints and smooth modification of desired volume size. The algorithm is useful in industrial design, where basic functional requirements are automatically obeyed without imposing limitation on the creativity of the designer. When the object material is known, volume preservation means weight preservation, hence is attractive for mechanical engineering applications when the engineer designs a part or an assembly. The preservation of volume of each element of the objects enables us to keep required proportions between volumes and weights of object parts. Obviously, simple scaling of the object in order to achieve a desired volume is not possible, due to the presence of point location and continuity constraints.

Our algorithm uses Bézier solids of arbitrary orders as the underlying mathematical definition of a free-form solid primitive. A Bézier solid of known orders is completely specified by its control points. The input to our algorithm consists of a desired object form (a set of primitives defined by their control points configurations), desired primitives volume sizes and a set of linear constraints on the control points implied by continuity requirements between the primitives or imposed directly by the user. The control points configurations can either be given directly by the user through control point manipulation, or computed from point displacement constraints specified by direct solid manipulation as in [Borrel91, Hsu92]. The algorithm computes a control point configuration closest to the given one (in a deformation energy minimization or least square sense) such that the deformed primitives contain volume of the given sizes and obeying the linear constraints. The algorithm does not automatically guarantee that the boundaries do not self-intersect.

Note that it is the global volume of a given free-form cell that is being preserved, not the volume of an object embedded within the cell or of local sub-cells. This approach was introduced in the finite element method for rubber type materials, but here we avoid the complexity of the penalty approach [Bercovier81] and use a duality argument to deal with the constraint, based on the Uzawa algorithm for non-linear programming [Arrow58. Ciarlet88].

Special measures were taken in order to endow the algorithm with real-time performance on current worksta-

<sup>&</sup>lt;sup>1</sup>[Sederberg86] refers to an unpublished report about volume preserving deformations, but such deformations cannot be everywhere locally satisfied with polynomial fields except for the simple case of pure shears.

tions. We utilize the fact that the volume size actually depends only on the boundary surfaces of the deformed primitive, hence volume size computation can be done with a subset of the control points. The inside points are of no interest to the user as well for the object's geometry, but are required for physical computations on the object, such as tear strength or deformation energy. The inside control points are computed from the outside points using a 3-D variant of the Coons surface formula when energy computation is required. This does not prevent them in general from crossing the parametric boundary, but intersection is not caused for most modeled objects.

In an interactive setting, the algorithm relaxes its accuracy requirements during object manipulation, computing an accurate solution only when real-time performance is no longer essential. This technique gives the user a feeling that volume is preserved during interaction.

Although in this work we limit the method description to Bézier solids, it can easily be adopted for most of the other common definitions of free-form solids, for example NURBS. The only restriction on the mathematical definition of the solid that we have is that it should be a defined as a linear combination of the control points.

The paper is organized as follows. Section 2 gives necessary mathematical notations. Section 3 formalizes the mathematical problem involved. Section 4 explains in detail how to compute the size of the volume enclosed by a tensor product Bézier solid and the partial derivatives of the volume size function. Section 5 explains how to represent continuity constraints. Section 6 explains how to compute the energy required for a change of a tensor product Bézier solid from one control point configuration to another, and the energy derivative. Section 7 presents the numerical algorithm used to solve the mathematical problem, and Section 8 describes our implementation and results.

### 2 Notations

We introduce here the formal mathematical notations used during the rest of the work. A tensor product Bézier solid is defined using a set of control points  $P_{ijk} \in \mathbb{R}^3$ . The image of a parametric point (u, v, w) in the unit cube is

$$F(u, v, w) = \sum_{i=0}^{n_u} \sum_{j=0}^{n_v} \sum_{k=0}^{n_w} B_i^{n_u}(u) B_j^{n_v}(v) B_k^{n_w}(w) P_{ijk}$$
(1)

where  $B_i^n(t)$  is the Bernstein polynomial defined by

$$B_i^n(t) = \binom{n}{i} (1-t)^{n-i} t^i$$

Denote the x, y, z coordinates of a control point by  $P_{ijk}^x, P_{ijk}^y, P_{ijk}^z$  respectively. Denote the volume of the

solid primitive defined by a set P of control points by Volume(P), and denote by  $\partial Volume(P)/\partial P$  the vector whose components are the partial derivatives

$$\frac{\partial Volume(P)}{\partial P_{abc}^{x}}, \frac{\partial Volume(P)}{\partial P_{abc}^{y}}, \frac{\partial Volume(P)}{\partial P_{abc}^{z}}$$

for every triplet  $abc \in [1 \dots n] \times [1 \dots m] \times [1 \dots l]$ .

Denote the energy of a transformation from a Bézier solid defined by a configuration Q of control points to one defined by configuration P of control points by Energy(P-Q), and denote by  $\partial Energy(P-Q)/\partial P$ the vector whose components are the partial derivatives

$$\frac{\partial Energy(P-Q)}{\partial P_{abc}^{x}}, \frac{\partial Energy(P-Q)}{\partial P_{abc}^{y}}, \frac{\partial Energy(P-Q)}{\partial P_{abc}^{z}}$$

Denote by  $\overline{P}$  the column vector of all the control points from all the Bézier solids in the system,  $(P_{ijk}^x, P_{ijk}^y, P_{ijk}^z, 1) = (P, 1)$ .

### **3** Problem Statement

The general problem we handle is finding a control point configuration that satisfies the constraints (linear and volume) and which results in an object as close as possible to the given one. The change of an object can be represented in two ways. The simpler is as the sum of squares of distances between the original control point positions and the new ones. The second is as the energy required to get from the original object to the new one. In this section we formalize this problem as a set of mathematical requirements that the target control points configuration should satisfy.

We denote by Dist(P,Q) the distance between two objects resulting from control points locations P and Q, which can stand for:

- Energy(P-Q), when using an energy approach,
- $\frac{1}{2}(P-Q)^T(P-Q)$ , when using a LSQ approach.

In case objects are modeled directly, the original objects are usually close to the desired final ones, in which case the the distance measure should be LSQ since we want the resulting object control points to be close to the original ones so that the shape of the object will incur a minimal change.

With physics-based modeling, we use as the original object the element in an initial state and we deform it by applying linear constraints and minimizing the energy. The resulting object then simulates the behavior of an elastic material with internal pressure. Initial control point configurations and the specification of constraints can be obtained by any method, including direct control point manipulation and direct manipulation of points and vectors inside or on the object.

The resulting constrained minimization problem (M) is: given a control point configuration  $Q_1, \ldots, Q_n$  (each

 $Q_i$  representing a single tri-variate primitive), a set of corresponding volume sizes  $V_1, \ldots, V_n$  and a matrix C representing linear constraints on the control points, find a new control point configuration  $P = P_1, \ldots, P_n$ such that the following holds:

- P is the solution of  $\min_{P'} \sum_{i=1}^{n} Dist(P'_i, Q_i)$ ,
- For each i,  $Volume(P_i) = V_i$ ,
- $C\bar{P}=0.$

The desired volumes  $V_i$  could be the initial volume sizes or any other number. For example, smooth variation of the desired volumes can be used for dilating the object during animations.

# 4 The Volume Function

Our volume preservation algorithm requires the computations of Volume(P) and of  $\frac{\partial Volume(P)}{\partial P}$ . Below we show how to analytically compute the exact volume size of a tensor product Bézier solid. We show that the computation of the volume size can be represented as a scalar product of two vectors: one whose components are the multiplication of the coordinates of the solid's control points, and a second one whose components are based on the Bézier basis functions and therefore can be computed off-line just once for each combination of orders of basis functions.

#### 4.1 Computing the Volume

The size of the volume specified by a three-dimensional function F(u, v, w) defined over the unit cube is

$$\int_0^1\int_0^1\int_0^1J_Fdudvdw$$

where  $J_F$  is the determinant of the Jacobian matrix of F:

$$J_F = \det\left(\frac{\partial F_i}{\partial x_j}\right), \ i = x, y, z \quad x_j = u, v, w.$$

In our case F is given by Equation 1. For example, the entry in the first row and column of the Jacobian matrix is

$$\frac{\partial F_x}{\partial u} = \sum_i \sum_j \sum_k \frac{d}{du} B_i^{n_u}(u) B_j^{n_v}(v) B_k^{n_w} P_{ijk}^x.$$

The derivative of a Bernstein polynomial of order n can be expressed by the scaled difference of two Bernstein polynomials of order n-1 [Farin92]:

$$\frac{d}{du}B_i^n(u) = n(B_{i-1}^{n-1}(u) - B_i^{n-1}(u))$$
(2)

with the convention that  $B_b^a(u) = 0$  for b < 0, a < 0, or b > a. Denote

$$\bar{u}_{ijk} = \frac{d}{du} B_i^n(u) B_j^{n_u}(v) B_k^{n_w}(w) =$$

$$_u (B_{i-1}^{n_u-1}(u) - B_i^{n_u-1}(u)) B_j^{n_u}(v) B_k^{n_w}(w),$$
(3)

and similarly  $\bar{v}_{ijk}$  and  $\bar{w}_{ijk}$ . The determinant  $J_F$  can be written as:

$$\det \left(\begin{array}{ccc} \sum_{ijk} \bar{u}_{ijk} P^x_{ijk} & \sum_{ijk} \bar{v}_{ijk} P^x_{ijk} & \sum_{ijk} \bar{w}_{ijk} P^x_{ijk} \\ \sum_{ijk} \bar{u}_{ijk} P^y_{ijk} & \sum_{ijk} \bar{v}_{ijk} P^y_{ijk} & \sum_{ijk} \bar{w}_{ijk} P^y_{ijk} \\ \sum_{ijk} \bar{u}_{ijk} P^z_{ijk} & \sum_{ijk} \bar{v}_{ijk} P^z_{ijk} & \sum_{ijk} \bar{w}_{ijk} P^z_{ijk} \end{array}\right)$$

expanding, we obtain:

,

n

$$J_{F} = \sum_{ijk} \bar{u}_{ijk} P_{ijk}^{x} \sum_{ijk} \bar{v}_{ijk} P_{ijk}^{y} \sum_{ijk} \bar{w}_{ijk} P_{ijk}^{z} \\ - \sum_{ijk} \bar{u}_{ijk} P_{ijk}^{x} \sum_{ijk} \bar{v}_{ijk} P_{ijk}^{z} \sum_{ijk} \bar{w}_{ijk} P_{ijk}^{y} \\ - \sum_{ijk} \bar{v}_{ijk} P_{ijk}^{x} \sum_{ijk} \bar{u}_{ijk} P_{ijk}^{y} \sum_{ijk} \bar{w}_{ijk} P_{ijk}^{z} \\ + \sum_{ijk} \bar{v}_{ijk} P_{ijk}^{x} \sum_{ijk} \bar{w}_{ijk} P_{ijk}^{y} \sum_{ijk} \bar{u}_{ijk} P_{ijk}^{z} \\ + \sum_{ijk} \bar{w}_{ijk} P_{ijk}^{x} \sum_{ijk} \bar{w}_{ijk} P_{ijk}^{y} \sum_{ijk} \bar{v}_{ijk} P_{ijk}^{z} \\ - \sum_{ijk} \bar{w}_{ijk} P_{ijk}^{x} \sum_{ijk} \bar{v}_{ijk} P_{ijk}^{y} \sum_{ijk} \bar{u}_{ijk} P_{ijk}^{z} .$$

Since the determinant is a multylinear operator and due to the structure of the summations, we can write:

$$J_F = \sum_{ijk} \sum_{lmn} \sum_{opq}$$
(4)

$$\det \begin{pmatrix} \bar{u}_{ijk} & \bar{u}_{lmn} & \bar{u}_{opq} \\ \bar{v}_{ijk} & \bar{v}_{lmn} & \bar{v}_{opq} \\ \bar{w}_{ijk} & \bar{w}_{lmn} & \bar{w}_{opq} \end{pmatrix} P^x_{ijk} P^y_{lmn} P^z_{opq}$$

Let I = ijklmnopq be a new index notation, in the range  $1 \dots (n_u n_v n_w)^3$ . Denote the determinant in Equation 4 by  $det_I(u, v, w)$ , and denote

$$c_I = \int_0^1 \int_0^1 \int_0^1 det_I(u, v, w) du dv dw$$

Since the integral is a linear operator, the volume can be written as:

$$Volume(P) = \sum_{ijk} \sum_{lmn} \sum_{opq} c_I P_{ijk}^x P_{lmn}^y P_{opq}^z.$$
(5)

Let **p** be a column vector indexed by *I* containing all terms of the form  $P_{ijk}^x P_{imn}^y P_{opq}^z$ , and let **c** be a column vector of the same size whose components are the  $c_I$ 's. Then Equation 5 can be expressed as the scalar product of **p** and **c**:

$$Volume(P) = \mathbf{c}^T \mathbf{p}$$

The vector c depends only on the orders of the Bézier basis functions, hence can be computed once and for all for every practical order combination (the number of all practically useful order combinations is small.) Computing the elements of c via symbolic integration is very complicated even for relatively small Bézier orders, therefore we compute them using Gauss numerical integration [Press88], which gives an exact result since the integrated functions are polynomials. A component  $c_I$  is computed as

$$c_I = \sum_r \sum_s \sum_t w_r w_s w_t det_I(x_r, x_s, x_t)$$

where  $w_r$ ,  $w_s$ ,  $w_t$  are the Gauss weights corresponding to the points  $x_r$ ,  $x_s$ ,  $x_t$  in the unit interval. The number of sample points on each dimension is determined according to the order of the basis function in that dimension.

The description above was simplified for ease of explanation. Actually, volume size depends only on the boundary surfaces (Stokes' formula [Gibson44].) In Bézier volumes the boundary surfaces are not influenced at all by the 'inner' control points, which can be completely neglected during the computation of the volume size. In fact, when computing the elements of the c vector we find that for any index ijklmnopq containing coordinates of an inside point the value  $c_{ijklnmopq}$  is zero. In practice, then, to accelerate the volume computation we let the index I run only on values of ijklmnopq which define 'outer' control points.

#### 4.2 Computing the Volume Derivative

The volume preservation algorithm requires the computation of the vector  $\partial Volume(P)/\partial P$  whose components are of the form  $\partial Volume(P)/\partial P_{abc}^{r}$  where r is x, y or z. For example,

$$\frac{\frac{\partial Volume(P)}{\partial P_{abc}^{r}}}{\frac{\partial \left(\sum_{ijk}\sum_{lmn}\sum_{opq} C_{ijk}lmnopq}P_{ijk}^{r}P_{lmn}^{y}P_{opq}^{z}\right)}{\partial P_{abc}^{r}}$$

Since for every  $ijk \neq abc$  the partial derivative vanishes, we get

$$\frac{\partial Volume(P)}{\partial P^{x}_{abc}} = \sum_{lmn} \sum_{opq} c_{abclmn\,opq} P^{y}_{lmn} P^{z}_{opq}.$$

### 5 The Constraints

In this section we explain the different linear constraints imposed on the control point configuration required in order to achieve desired geometric or physical results.

#### 5.1 Continuity Constraints

Continuity constraints between primitives in an object are essential for any object design. Continuity of order k ( $C^k$ ) between two adjacent volumes  $F_1(u, v, w)$  and  $F_2(u, v, w)$  defined on [0, 1] in the *u* direction is achieved when the following holds

$$\left(\frac{\partial^{k} F_{1}(u, v, w)}{\partial^{k} u}\right)(1, v', w') = \left(\frac{\partial^{k} F_{2}(u, v, w)}{\partial^{k} u}\right)(0, v', w').$$

for every  $(v', w') \in [0, 1] \times [0, 1]$ .

In our case for two adjacent primitives defined by control points configurations P and Q we will get

$$\left(\frac{\partial^{k}(\sum_{ijk}B_{i}^{n_{u}}(u)B_{j}^{n_{v}}(v)B_{k}^{n_{w}}(w)P_{ijk})}{\partial^{k}u}\right)(1,v',w') = \left(\frac{\partial^{k}(\sum_{ijk}B_{i}^{n_{u}}(u)B_{j}^{n_{v}}(v)B_{k}^{n_{w}}(w)Q_{ijk})}{\partial^{k}u}\right)(0,v',w').$$

Since the derivative is a linear operator we get

$$\left( \sum_{ijk} \left( \frac{\partial^k B_i^{n_u}(u)}{\partial^k u} \right) B_j^{n_v}(v) B_k^{n_w}(w) P_{ijk} \right) (1, v', w') = \\ \left( \sum_{ijk} \left( \frac{\partial^k B_i^{n_u}(u)}{\partial^k u} \right) B_j^{n_v}(v) B_k^{n_w}(w) Q_{ijk} \right) (0, v', w').$$

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and therefore

$$\sum_{jk} B_{j}^{n_{u}}(v') B_{k}^{n_{u}}(w') (\sum_{i} (\frac{\partial^{k} B_{i}^{n_{u}}(u)}{\partial^{k} u} (1) P_{ijk} - \frac{\partial^{k} B_{i}^{n_{u}}(u)}{\partial^{k} u} (0) Q_{ijk})) = 0.$$

For this to hold for each  $v', w' \in [0, 1] \times [0, 1]$ , a necessary and sufficient condition is that for  $j = 1 \dots n_v$ ,  $k = 1 \dots n_w$ 

$$\sum_{i} \left( \frac{\partial^{k} B_{i}^{n_{u}}(u)}{\partial^{k} u}(1) P_{ijk} - \frac{\partial^{k} B_{i}^{n_{u}}(u)}{\partial^{k} u}(0) Q_{ijk} \right) = 0$$

thus getting a set of  $n_v n_w$  linear equations in  $P_{ijk}$ ,  $Q_{ijk}$ .

The derivative of a Bernstein polynomial of order n was given in Equation 2. Since  $B_l^m(0) \neq 0$  for l = 0 only and  $B_l^m(1) \neq 0$  only for l = m then the number of *i*'s for which  $\frac{\partial^k B_l^{n_u}(u)}{\partial^k u}(1) \neq 0$  is k + 1 and the same holds for  $\frac{\partial^k B_l^{n_u}(u)}{\partial^k u}(1)$ . Therefore,  $C^k$  continuity conditions between adjacent Bézier volumes are expressed as a set of  $n_v n_w$  linear equations on k+1 layers of control points of each volume from the adjacent border.

For the most common cases the conditions are:

$$C^{0} = P_{n_{u},i,j} - Q_{1,i,j} = 0, \quad i = 1 \dots n_{v}, \quad j = 1 \dots n_{w}$$

$$C^{1} = n_{u}(P_{n_{u},i,j} - P_{n_{u}-1,i,j}) - n_{u}(Q_{2,i,j} - Q_{1,i,j}) = 0,$$

$$i = 1 \dots n_{v}, \quad j = 1 \dots n_{w}.$$

Another kind of continuity constraint between elements is geometric continuity, which is more general than parametric continuity. Geometric continuity yields non-linear constraints which are difficult to express and solve, and therefore we do not use them in this work. For more details on geometric constraints see [Bercovier93].

#### 5.2 Other Constraints

The following types of constraints can easily be handled in addition to continuity constraints:

- Fixing a point at a given location, resulting in equations such as  $P_i^r c_i^r = 0$ , r = x, y, z, where  $c_i^r$  is a constant.
- Attaching two points together, resulting in equations such as  $P_i^r - Q_i^r = 0$ , r = x, y, z.
- Preserving a given distance between points.

#### 5.3 Summary

A general linear equation on variables  $P_i$  is expressed as

$$\sum_{i} c_i P_i + c_{n+1} = 0$$

or in vector representation as

$$c^T(P,1)=0.$$

If we denote by C the matrix whose rows are the coefficients c of the linear equations and by  $\tilde{P}$  the column vector of all the control points, the constraints are achieved when  $C\tilde{P} = 0$ .

### 6 The Energy Function

Energy computation for a deformation of a Bézier primitive from one control point configuration to another is required by our algorithm. Here we show that it can be computed using a matrix whose elements depend only on the order of Bézier basis functions.

#### 6.1 Computing the Energy

The energy of a deformation of a unit cube specified by a 3-D vector function  $F(x_1, x_2, x_3)$  is usually [Terzopolus94] described as

$$\int_{0}^{1} \int_{0}^{1} \int_{0}^{1} \frac{1}{2} \left(\beta \sum_{i} \left(\frac{\partial F_{i}}{\partial x_{i}}\right)^{2} + \left(\beta \sum_{i \neq j} \left(\left(\frac{\partial F_{i}}{\partial x_{j}}\right) + \left(\frac{\partial F_{j}}{\partial x_{i}}\right)\right)^{2}\right) dx_{1} dx_{2} dx_{3}$$

$$(6)$$

with  $\alpha$  and  $\beta$  being material property constants. We can write

$$Energy(F) =$$
(7)  

$$\frac{1}{2}\beta \sum_{i} \int_{0}^{1} \int_{0}^{1} \int_{0}^{1} \left(\frac{\partial F_{i}}{\partial x_{i}}\right)^{2} du dv dw +$$

$$\frac{1}{2}\alpha \sum_{i \neq j} \left(\int_{0}^{1} \int_{0}^{1} \int_{0}^{1} \left(\frac{\partial F_{i}}{\partial x_{j}}\right)^{2} du dv dw +$$

$$+ \int_{0}^{1} \int_{0}^{1} \int_{0}^{1} \left(\frac{\partial F_{j}}{\partial x_{i}}\right)^{2} du dv dw +$$

$$2 \int_{0}^{1} \int_{0}^{1} \int_{0}^{1} \left(\frac{\partial F_{j}}{\partial x_{i}}\right) \left(\frac{\partial F_{i}}{\partial x_{j}}\right) du dv dw \right).$$

In our case the deformations is of a body defined by one Bézier point configuration to a body defined by another one. Hence the deformation is defined as a trivariate Bézier function with distances between the control points of the two configurations serving as its control point lattice. Using P for these new 'control points', we can write

$$\frac{\partial F_x}{\partial x_1} = \frac{\partial F_x}{\partial u} = \sum_{ijk} \bar{u}_{ijk} P^x_{ijk}$$

and consequently

$$\frac{\partial F_x}{\partial u}^2 = \sum_{ijk} \sum_{lmn} \bar{u}_{ijk} \bar{u}_{lmn} P^x_{ijk} P^x_{lmn}.$$

We have

$$\int_{0}^{1} \int_{0}^{1} \int_{0}^{1} \left(\frac{\partial F_{x}}{\partial u}\right)^{2} du dv dw =$$

$$\int_{0}^{1} \int_{0}^{1} \int_{0}^{1} \left(\sum_{ijk} \sum_{lmn} \bar{u}_{ijk} \bar{u}_{lmn} P_{ijk}^{x} P_{lmn}^{x}\right) du dv dw =$$

$$\sum_{ijk} \sum_{lmn} P_{ijk}^{x} P_{lmn}^{x} \left(\int_{0}^{1} \int_{0}^{1} \int_{0}^{1} \bar{u}_{ijk} \bar{u}_{lmn} du dv dw\right)$$

Let Du be a matrix indexed by ijk and lmn, defined by

$$Du_{ijk,lmn} = \int_0^1 \int_0^1 \int_0^1 \bar{u}_{ijk} \bar{u}_{lmn} du dv dw$$

and define Dv and Dw similarly. Let Px be a column vector with components  $P_{ijk}^x$ , and similarly define Py and Pz, then we have

$$\int_0^1 \int_0^1 \int_0^1 \left(\frac{\partial F_x}{\partial u}\right)^2 du dv dw = P^{xT} D u P^x.$$

Denote by Duv the matrix of the mixed derivatives given by

$$Duv_{ijk,lmn} = \int_0^1 \int_0^1 \int_0^1 \bar{u}_{ijk} \bar{v}_{lmn} du dv dw$$

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and define Duw and Dvw similarly ( $Dvu = Duv^T$ .) We have

$$\int_0^1 \int_0^1 \int_0^1 \left(\frac{\partial F_y}{\partial u}\right) \left(\frac{\partial F_x}{\partial v}\right) du dv dw = P^{yT} Du v P^x.$$

The elements of the three matrices above can also be computed numerically. Substituting the matrices into Equation 7,

$$Energy(F) = \frac{1}{2}(\beta(P^{rT}DuP^{r} + P^{yT}DvP^{y} + P^{zT}DwP^{z}) + \alpha((P^{rT}DvP^{r} + P^{yT}DuP^{y} + P^{rT}Duv^{T}P^{y} + P^{yT}DuvP^{r}) + (P^{rT}DwP^{r} + P^{zT}DuP^{z} + P^{rT}Duw^{T}P^{z} + P^{zT}Duw^{T}P^{z} + P^{zT}DuwP^{x}) + (P^{yT}DwP^{y}) + (P^{yT}DwP^{y}) + (P^{zT}DvP^{z} + P^{zT}DvP^{z} + P^{zT}DvP^{z}))).$$

Finally, let D be the matrix whose rows are

$$\frac{1}{2}(\beta Du + \alpha (Dv + Dw), \alpha Duv^{T}, \alpha Duw^{T}),$$
$$\frac{1}{2}(\alpha Duv, \beta Dv + \alpha (Du + Dw), \alpha Dvw^{T}),$$

and

$$\frac{1}{2}(\alpha Duw, \alpha Dvw, \beta Dw + \alpha (Du + Dv)).$$

Let **P** be a column vector concatenating (Px, Py, Pz). Then, we have

$$Energy(P) = \mathbf{P}^T D \mathbf{P}.$$

All the elements of D depend only on the orders of the Bézier basis functions, hence can be computed exactly once and for all for every practical order combination, using Gauss quadrature.

#### 6.2 Computing the Energy Derivative

To minimize the deformation energy the algorithm requires the computation of the vector  $\partial Energy(P)/\partial P$ whose components are of the form  $\partial Energy(P)/\partial P_{abc}^{r}$ where r is x, y or z. It is easy to see that, for example,

$$\frac{\partial Energy(P)}{\partial P_{abc}^{r}} = 2\sum_{r}\sum_{lmn} D_{xabc,rlmn} P_{lmn}^{r},$$

since for every  $rijk \neq xabc$  the partial derivative of Energy(P) according to  $P_{abc}^{x}$  vanishes.

# 7 The Uzawa-Based Volume Preservation Algorithm

In this section we explain in detail the algorithm we use for solving the problem as defined in Section 3.

#### 7.1 Lagrangian Multiplier Method

To convert the constrained minimization problem (M): minimize  $\sum_{i=1}^{n} Dist(P_i, Q_i)$  subject to the constraints  $C\bar{P} = 0$  and  $Volume(P_i) - V_i = 0$ ,  $i = 1 \dots n$ , into an unconstrained min-max problem, we define a new functional L called the Lagrangian associated with the problem (M) by

$$L(P, \lambda, \gamma) = \sum_{i} Dist(P_{i}, Q_{i}) + \sum_{i} \lambda_{i}(Volume(P_{i}) - V_{i}) + \gamma C\bar{P}$$

where  $\gamma$  is vector with size the number of linear constraints. The vector  $(\lambda_1 \dots \lambda_n, \gamma_1 \dots \gamma_m)$  is called the Lagrange multipliers vector.  $\lambda_i$  is called the Lagrange multiplier for the constraint  $Volum \epsilon(P_i) - V_i = 0$  and  $\gamma_j$  is called the Lagrange multiplier for the constraint  $C_j \bar{P} = 0$  ( $C_j$  stands for row j of C).

As explained in [Ciarlet88], the constrained minimization problem (M) can be reformulated as finding a solution to the unconstrained min-max problem (S) defined by

$$(S) = \max_{\lambda,\gamma} \min_{P} L(P,\lambda,\gamma)$$

A necessary condition for a triplet  $(P, \lambda, \gamma)$  to be a solution of (S) is the vanishing of the partial derivatives:

$$\frac{\partial L}{\partial P} = \frac{\partial L}{\partial \lambda} = \frac{\partial L}{\partial \gamma} = 0.$$

which means that for each  $i = 1...n, j = 1...n, n_w n_w n_w n_w n_w x = x, y, z$ :

$$\frac{\partial L}{\partial P_{i,j}^r} = \frac{\partial Dist(P_i, Q_i)}{\partial P_{i,j}^r} +$$

$$\lambda_i \frac{\partial Volume(P_i)}{\partial P_{i,j}^r} + \gamma C|_i = 0.$$
(8)

 $(C|_i$  denotes the columns of C that multiply the points of  $P_i$  in  $\overline{P}_i$ ) and

$$\frac{\partial L}{\partial \lambda_i} = Volume(P_i) - V_i = 0.$$
$$\frac{\partial L}{\partial \gamma_j} = C_j \bar{P} = 0, \quad j = 1...m.$$

#### 7.2 Solution Method

The volume derivative expression is non-linear, hence the usual direct methods (such as  $LDL^T$  and Gauss elimination) cannot be used to solve (S). We use a version of the Uzawa method tailored to our problem [Ciarlet88]. Uzawa's method is an iterative method allowing one to solve an inequality constrained minimization problem by replacing it with a sequence of unconstrained minimization problems. Since we do not have inequality constraints we can use a simpler version. Given the problem (M) the iteration starts with an arbitrary values for  $\lambda^0 \in R_+^n$ ,  $\gamma^0 \in R_+^m$  (we start with 0 for both), and with an initial value for  $P^0$  for which we use Q. These initial guesses are especially suitable in an interactive setting, where it is expected that Q will not change much after the constraints are satisfied. A sequence of triplets

$$(P^k, \lambda^k, \gamma^k) \in V \times R^n_+ \times R^m_+, \ k \ge 0$$

is defined by means of the following iterations:

$$P^{k}: solves \inf_{P' \in V} (\sum_{i} Dist(P'_{i}) + \sum_{i} \lambda_{i}(Volume(P'_{i}) - V_{i}) + \gamma C\tilde{P}'),$$
(9)

$$\lambda_i^{k+1} = max(\lambda_i^k + \rho_1(Volume(P_i) - V_i), 0) \qquad (10)$$
  
$$i = 1 \dots n,$$

$$\gamma_j^{k+1} = max(\gamma_j^k + \rho_2(C_j\bar{P}), 0)$$
(11)  

$$i = 1 \dots m.$$

The algorithm runs until the constraints are satisfied or the number of iterations exceeds a given limit.

Pseudo-code for the algorithm is shown on Figure 1. The initial values for  $P, \lambda$  and  $\gamma$  are set in lines 1 and 2. Line 3 computes the current volumes  $v_i$  and line 4 initializes the loop counter k. The main ('outer') loop of the algorithm is performed in lines 5-10. The loop iterates while the constraints are not satisfied, stopping after the limit on the number of iterations has been reached. In each iteration the system in Equation 8 is solved (line 6) and the current value of  $\lambda$  and  $\gamma$  is updated using the tuning parameters  $\rho_1$  and  $\rho_2$  respectively (lines 7-8). Line 11 returns P as the answer.

The choice of tuning parameters  $\rho_1$  and  $\rho_2$  as used in Equation 10 and Equation 11 is the most difficult practical issue when using Uzawa's method. Each type of problem has its own best range of values for  $\rho_i$ . In our case we found it best to use  $\rho_1 = \rho_2 = 0.15$  for an energy distance function and  $\rho_1 = \rho_2 = 0.5$  for a least squares distance function. In general, the larger the  $\rho$ 's the faster the convergence becomes, but the risk of nonconvergence due to overstepping the convergence point increases.

Pseudo-code for one way of solving the inner problem is shown in Figure 2. The inner problem is solving Equation 8 for P with the given  $\gamma$  and  $\lambda$ . It is a nonlinear problem; Figure 2 shows how to solve it using successive approximation on P. We iteratively compute new values for P based on Equation 8 until the distance between two successive iterations is small enough  $(|| P - P' ||^2 < \delta_{dist})$ . There are several other possible techniques for solving a set of non-linear equations which can be used here as well. Usually when solving physics-based problems by Lagrange multipliers methods the additional variables added as multipliers have physical meaning. In our case one can interpret  $\lambda$  as an inner hydrostatic pressure to keep the volume at a given value. We are looking for the value of that pressure: the Uzawa outer step can be seen as augmenting or diminishing the hydrostatic pressure until convergence. This tuning is done with the parameter  $\rho_1$ . This observation relates our method to so-called mixed finite element methods for the Stokes problem [Hughes87]. In our case we have constant pressure for each small volume element.

### 8 Results

The algorithm was implemented in C under Unix using SGI/GL for graphics and Motif for the user interface. The interface lets the user work with a number of Bézier primitives, the order of each selectable by the user. In the initial state the primitives are displayed as unit cubes (cubes whose volume is 1.) Control points on each primitive can be selected and manipulated in 3-D. We did not implement direct manipulation of boundary surface points since it is immaterial to the problem being tackled. The primitives as whole can be selected as well and manipulated.

Constraints are inserted via a Motif-based user interface where the type of the constraint is set and then through direct point manipulation the points or surfaces involved are chosen.

There are two methods for object design. In the first method, volume preservation can be turned off during interaction and performed only when arriving at a desired configuration. In the second method, it can be turned on during the whole interaction process. The first option is necessary since for high orders the performance is not fully interactive.

Due to the complexity of computations in the interactive stage we cannot satisfy volume and linear constraints simultaneously, so the user has to choose which one is preferred.

There are three sets of parameters to the algorithm: parameters that influence volume preservation during interaction while the user drags the mouse, parameters that are for solving volume constraint when leaving the mouse, and parameters for global computation when solving all the constraints. Typically, for the interaction mode the iteration limits are lower and the convergence tolerances are larger than for the final mode, for the global computation the tolerances usually do not increase but the iteration limits are larger and the  $\rho_i$  used are smaller.

Different sets of parameters do not cause divergence of the algorithm, since during interaction the current configuration is very close to a solution satisfying the volume constraint, and the algorithm needs fewer iterations to reach a solution. The parameter sets can be tuned using a dialog box.

The user can manipulate a scale widget that defines the desired volume for a chosen primitive. The volume preservation algorithm is performed repeatedly while the scale is dragged.

Tri-quadratic free-form volume design is fully interactive. For a typical movement of a single control point, to reach a final volume tolerance of  $10^{-4}$  and a final distance tolerance of  $10^{-3}$  requires about 15 outer iterations, each of them with 1-2 inner iterations. This takes about 3 seconds on Silicon Graphics workstations with a MIPS R-4000 processor. During interaction it is enough to set both tolerances to  $10^{-2}$ , in which case the solution is completed in real-time.

For a tri-cubic free-form volume, to reach the same tolerances requires about 25-30 outer iterations, each of them with one inner iteration. This takes about 15 seconds. When both tolerances are set to  $10^{-2}$  during interaction the solver takes about 3 seconds, hence tri-cubic interaction could be done in real-time using a faster processor.

The running times above are of course dependent on the number of linear constraints and on how far the current configuration is from their solution.

designed using the system. The amphora is modeled from a single primitive, and the phone was modeled from three tri-cubic primitives with  $C^1$  continuity conditions between them. Its parts were designed by volume modifications to create the right proportions between them while keeping the desired shape constraints and continuity.

# 9 Conclusion

We presented an approach for modeling with free-form solid primitives while preserving the volume contained within each primitive and satisfying continuity constraints between the primitives. Careful tuning allows our Uzawa-based non-linear optimization algorithm to be fully interactive for tri-quadratic volume elements and almost interactive for tri-cubic elements. The algorithm possesses several possible applications in computer animation, industrial design and mechanical engineering, broadening the scope of physics-based geometric modeling.

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Input: A set of control point configurations  $Q_1, \ldots, Q_n$ A set of desired volumes  $V_1, \ldots, V_n$ A matrix C of linear constraints Output: A set of new control point configurations  $Q_1, \ldots, Q_n$ that satisfies the Goal conditions. Parameters: Convergence tolerances  $\delta_{vol}, \delta_{cons}, \delta_{dist}$ Iteration limits  $L, L_{dist}$ Tuning parameters  $\rho_1, \rho_2$ Goal: Minimize  $\sum_{i} Dist(P_i, Q_i)$  subject to  $Volume(P_i) = V_i, C\bar{P} = 0$ Algorithm: 1 P = Q2  $\lambda = 0, \gamma = 0$ 3  $v_i = Volume(Q_i)$   $i = 1 \dots n$ 4 k = 05 while  $\sum_{i} |v_i - V_i| > \delta_{vol}$  and  $|| C\bar{P} || > \delta_{cons}$  and k < L6 find  $P_1 \dots P_n$  that solve  $\frac{\partial Dist(P_i)}{\partial P_i} + \lambda_i \frac{\partial Volume(P_i)}{\partial P_i} + \gamma C|_i = 0$ 7  $\lambda = \lambda + \rho_1(v - V)$ 8  $\gamma = \gamma + \rho_2 C\bar{P}$ 9  $v_i = Volume(P_i)$   $i = 1 \dots n$ 10 k = k + 1end 11 return P

Figure 1: The Uzawa-based volume preservation algorithm.

Solution of 
$$\frac{\partial D_{ist}(P_{1})}{\partial P_{i}} + \lambda_{i} \frac{\partial Volume(P_{1})}{\partial P_{i}} + \gamma C|_{i} = 0, i = 1 \dots n$$
:  
1 if dist function is energy  
2  $P_{ij}^{new} = Q_{ij} - \frac{\sum_{k \neq j} 2D_{jk}(P_{ik} - Q_{ik}) + \lambda_{i} \frac{\partial Volume(P_{i})}{\partial P_{ij}} + \gamma C|_{ij}}{2D_{jj}}$   
for each control point coordinate  $i, j$   
3 else  
4  $P_{ij}^{new} = Q_{ij} - \lambda_{i} \frac{\partial Volume(P_{i})}{\partial P_{ij}} - \gamma C|_{ij}$   
for each control point coordinate  $i, j$   
5 endif  
6  $l = 0$   
7 while  $|| P - P^{new} ||^{2} > \delta_{dist}$  and  $l < L_{dist}$   
8  $P = P^{new}$   
9 if dist function is energy  
10  $P_{ij}^{new} = Q_{ij} - \frac{\sum_{k \neq j} 2D_{jk}(P_{ik} - Q_{ik}) + \lambda_{i} \frac{\partial Volume(P_{i})}{\partial P_{ij}} + \gamma C|_{ij}}{2D_{jj}}$   
for each control point coordinate  $i, j$   
11 else  
12  $P_{ij}^{new} = Q_{ij} - \lambda_{i} \frac{\partial Volume(P_{i})}{\partial P_{ij}} - \gamma C|_{ij}$   
13 endif  
14  $l = l + 1$   
end  
15 return  $P^{new}$ 

Figure 2: The 'inner' P computation by successive approximation.



Figure 3: An amphora, modeled with a single tri-cubic free-form solid.



Figure 4: A phone, modeled with three tri-quadratic elements.