Estimating Volumes of Simulated Lung Cancer Nodules*

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Abstract

Lung cancer is a disease of uncontrolled cell growth in tissues of the lung. Computed tomography (CT) shows promise in detecting lung cancers at earlier, more operable stages, when survival is better. CT scans generate multiple 2-D slice images of the lung and digital image processing software is used to combine these images into a 3-D representation of the lung and, in particular, an identified cancer lesion. Various CT scanners use, often different and usually proprietary, software to develop these 3-D images and generate parameters such as lesion volume. Tracking lesion volume is considered a good diagnostic tool for evaluating the results of patient treatment. The Food and Drug Administration (FDA) is conducting research on developing reference cancer lesions, called phantoms, to test CT scanners and their proprietary software. FDA loaned two semi-spherical phantoms to NIST, called Green and Pink, and asked to have the phantoms measured by a coordinate measuring machine (CMM) and the volumes estimated. This report describes in detail both the experimental and computational methods used to estimate the phantoms' volumes as well as a bootstrap method for estimating the uncertainties of the computed volumes. Three sets of CMM measured data were produced. One set of data involved data density measurements of a known calibrated metal sphere. The other two sets were measurements of the two FDA phantoms at two densities, called the coarse set and the dense set. Two computational approaches were applied to the data. In the first approach spherical models were fit to the calibrated sphere data and to the phantom data. The second approach was to model the data points on the boundaries of the spheres with surface B-splines and then use the Divergence Theorem to estimate the volumes. The results for the coarse data set tended to predict the volumes as expected with low expanded uncertainties and the results did show that the Green phantom was very near spherical. This was confirmed by both computational methods. The spherical model did not fit the Pink phantom as well and the B-spline approach provided a better estimate of the volume in that case. The results for the dense data set did not provide as nice a prediction of volumes as the coarse data set and produced larger expanded uncertainties. A discussion of some possible reasons for these results will be given in the conclusion section. The report includes the MATLAB codes used in the study.

Keywords: B-splines, computed tomography, coordinate measuring machine, divergence theorem, lung cancer, lung cancer phantoms, nonlinear least squares.

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1 Introduction

Lung cancer is a disease of uncontrolled cell growth in tissues of the lung. It is the most common cause of cancer related deaths in men and second most in women and is responsible for an estimated 560,000 deaths in the US in 2007. It is classified as small cell (13%) and non-small cell (87%) [12]. The most common cause is long term exposure to tobacco smoke. Lung cancer lesions may be seen in chest X-ray and computed tomography (CT), although chest X-ray analysis has shown limited effectiveness in improving survival. Computed tomography has shown promising results in detecting lung cancers at earlier, more operable stages, when survival is better [1].

Tomography is a method of imaging a single slice of the body, in this case the lung. Modern CT is a medical imaging method that uses tomography but also employs digital image processing techniques to generate three dimensional images built from a large sequence of two-dimensional X-ray images made around a single axis. There are a number of current CT methods but the one that is of most interest in this study is called a helical or spiral CT in which the X-ray source is connected to a rotating gantry. The patient, on a table, moves smoothly through the scanner. The name, helical or spiral CT, comes from the helical path traced out by the X-ray beam. The helical trajectory of the data is converted into two-dimensional image slices of the patient's body. The advantage of the helical or spiral CT is speed, since the patients must remain perfectly still and hold their breath. A large portion of the body can be scanned in 20-60 seconds [11]. CT provides excellent diagnostics for detecting changes in the lungs, where two-dimensional X-rays do not clearly show the lung cancer changes. A diagnosis is usually confirmed by biopsy. Treatments include surgery and chemotherapy. With treatment, the one-year survival was 42% in 2002 but the five-year survival rate was only 16% [1].

Evaluating the rate of growth/shrinkage of lung cancer lesions is important for pharmaceutical research in order to compare drug effects, for clinical practice to determine time to use drugs or surgery, and for patient follow up to quantify the effects of drugs or surgery. A goal then is to enable CT scans to become a measurement device, providing quantitative rather than only qualitative information. The current guidelines for measuring the increase or decrease of tumors involve linear measurements of the greatest diameter of a lesion from the sequence of CT slices [19]. However, the volume change of lesions is found to show more significant relative change than a relative diameter change. As an example, for a sphere a 30% decrease in the relative diameter would produce a 65.7% decrease in relative volume whereas a 30% increase in the relative diameter would produce approximately a 120% increase in relative volume.

The Food and Drug Administration (FDA) is conducting research on developing reference cancer lesions, called phantoms, to test CTs and their proprietary software. Two samples were loaned to NIST to estimate volumes. The material of which the phantoms are composed simulates lung cancer material. They can be inserted into a simulated body torso for CT scans. The two phantoms are shown in Fig. 1. Although they seem spherical they are slightly non-spherical. The larger one on the right is referred to as the Green phantom and the one on the left is called the Pink phantom due to the material colors

One approach to estimating the volume of the phantoms is to use what is called an Archimedes test in which the phantoms would be immersed in a liquid bath in a well calibrated container with fine measurement gradations to determine liquid displacement. However, in the case of these phantoms, the material used to manufacture them is porous and the phantoms would have to be coated. This, of course, would affect the "ground truth" volume estimate. Furthermore, the potential expense involved with the process was found to be sufficiently significant that another approach was chosen.

The approach chosen for this study is based on a fundamental theorem in calculus, called the Divergence Theorem (see Taylor [17]), which is an analogue of Green's Theorem in two dimensional



Figure 1: Two simulated lung cancers, called phantoms.

space. In the Divergence Theorem a volume integral is shown to be equal to a surface integral. Therefore, we surmised that, if a model of the surfaces of the phantoms could be developed, we could use the Divergence Theorem to estimate their volumes. In order to develop a surface model we needed to obtain data about the surface. This was done using a coordinate measuring machine (CMM) in the Manufacturing Engineering Lab (MEL) at NIST. This machine produced a set of (x, y, z) points on the surface of each phantom. The data were then transformed to spherical coordinates and modeled using a set of basis functions, called B-splines. After fitting, the B-spline model was then used to generate a grid of values on the surface. These values were used to form surface triangles that were then used to compute the necessary surface integrals and finally the volume. The quality of the volume estimates depended on the surface grid sizes, as will be clear from the discussion below. The method is not new, in that it was suggested in the book by Dierckx [6]. A reader can also consult the Dierckx references [4] and [5].

The report is divided as follows. Section 2 introduces B-splines and tensor products of B-splines. Section 3 describes the surface point generation experiments using the CMM. Section 4 describes the methods and results from applying the spherical and B-spline models to the CMM data. Appendices include the MATLAB codes used in the study.

2 Introduction to B-splines

In this section we will define cubic B-splines and show the construction of bivariate or tensor products of B-splines on a two dimensional array. We will then give a discussion of how B-splines are computed. These specific computational algorithms will not be implemented in code, since they are available as functions in the MATLAB spline toolbox. The algorithmic discussions are given, though, as background to understanding the output from the MATLAB functions.

2.1 Cubic B-Splines in One Variable

Suppose that a function y = f(x) is known at the m points $(x_1, y_1), \dots, (x_m, y_m)$, where $a < x_1 < x_2 < \dots < x_m < b$, $y_k = f(x_k)$. It is known that a polynomial of degree m-1, P(x), can be constructed to pass through these m points. In the case of highly accurate data points this polynomial can be constructed to interpolate these points by, for example, Lagrange polynomial or Newton divided difference algorithms. But, in the case of large m, it is also well known that

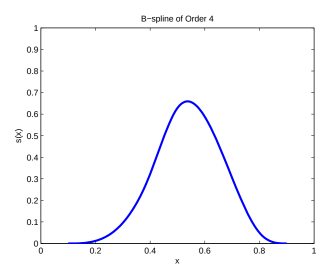


Figure 2: Plot of a B-spline of Order 4.

polynomials of high degree can produce unwanted oscillations between the interpolated points. It is crucial then to approximate sets of data with as low degree polynomials as possible. Of course these polynomials may or may not interpolate the data points but may be made to come as close to them as possible. This is a significant advantage of functions called splines.

Given a set of m real numbers, satisfying $a < x_1 < x_2 < \cdots < x_m < b$, a spline function, s(x), of order n (or degree n-1) with knots x_1, x_2, \cdots, x_m is a function that satisfies two properties: (1) in each of the intervals $x \le x_1$; $x_{j-1} \le x \le x_j$, $(j=2,3,\cdots,m)$; $x_m \le x$, s(x) is a polynomial of degree n-1 or less. (2) s(x) and its derivatives of orders $1,2,\cdots,n-2$ are continuous. This would mean, for example, a spline function of order four would be constructed from polynomials of degree three (cubic) or less on the intervals $x \le x_1$; $x_{j-1} \le x \le x_j$, $(j=2,3,\cdots,m)$; $x_m \le x$ with continuous derivatives of orders one and two. In this report only splines of order four will be used.

Instead of approximating data with one spline, another approach is to build an approximating function by forming a linear combination of functions. These functions are usually referred to as basis functions. This is a well known mathematical technique of constructing complex functions by forming linear combinations of simpler functions. In the current report data sets will be approximated by linear combinations of special spline functions, called B-splines, for Basis splines. Only B-splines of order four will be considered here. B-splines of order four are cubic splines that are non-zero only over four adjacent intervals between knots (see Fig. 2). The notation for a B-spline is $N_{i,k}(x)$, where $N_{i,k}(x)$ is zero everywhere except in the range $x_{i-k} < x < x_i$, where in this work k=4. As a note, sometimes B-splines are denoted in the literature by $B_{i,k}(x)$, but in this report we will use $N_{i,k}(x)$. To simplify notation let $N_i(x) = N_{i,4}(x)$. Then a B-spline of order four, or cubic B-spline, is a cubic spline with knots x_{i-4} , x_{i-3} , x_{i-2} , x_{i-1} , x_i that is zero everywhere except in the range $x_{i-4} < x < x_i$. $N_i(x)$ is defined uniquely except for a scaling factor and is conventionally taken to be positive throughout the range $x_{i-4} < x < x_i$ and has a single maximum value. Since $N_i(x)$ is a cubic spline it has continuous derivatives of order one and two at x_{i-4} and x_i . These derivatives are zero at the endpoint knots.

To define a complete set of B-splines on the set of points $a < x_1 < x_2 < \cdots < x_m < b$ it is necessary to introduce eight additional points at the boundaries given by x_{-3} , x_{-2} , x_{-1} , x_0 , x_{m+1} ,

 $x_{m+2}, x_{m+3}, x_{m+4}$. It is usual to have $x_0 = a, x_{m+1} = b$. With this augmented set of knots one can define m+4 fundamental cubic B-splines, $N_i(x), i=1,2,\cdots,m+4$. Then the general cubic B-spline has a unique representation in the range $a \le x \le b$ of the form

$$s(x) = \sum_{i=1}^{m+4} c_i N_i(x). \tag{1}$$

There are computational advantages of cubic B-splines. For any given x, all but four adjacent $N_i(x)$ are zero. In particular, if $x \in [x_{i-1}, x_i]$ the four non-zero cubic B-splines are $N_i(x)$, $N_{i+1}(x)$, $N_{i+2}(x)$, $N_{i+3}(x)$.

A least squares curve fitting problem to $a < u_1 < u_2 < \cdots < u_m < b, y_k = f(u_k)$ becomes one of determining the coefficients c_i as the least squares solution to the equations

$$\sum_{i=1}^{m+4} c_i N_i (u_r) = f_r, \quad r = 1, 2, \dots, m.$$
 (2)

These may be written in matrix notation as

$$Ac = f, (3)$$

where A is the $m \times (m+4)$ matrix whose element in column i of row r is $N_i(x_r)$ and c, f are column vectors with elements c_i , f_r respectively. If the data points are arranged in increasing order of x, then the matrix A becomes a banded matrix with bandwidth four. For a more thorough discussion of B-splines see de Boor [3].

2.2 Evaluation of B-splines

Let k be the order of the desired B-splines and let x_1, x_2, \dots, x_m be mesh points. In order to develop a complete set of B-splines we need to extend the mesh array by k points on each end as described in the previous section. On the left end, extend the mesh by setting

$$x_{-(k-1)} = x_{-(k-2)} = \dots = x_0 = a,$$
 (4)

and, on the right set

$$x_{n+1} = x_{n+2} = \dots = x_{n+k} = b. (5)$$

Now define a working array that incorporates all of the mesh and the extended mesh points as

$$xt_{1} = x_{-(k-1)}, xt_{2} = x_{-(k-2)}, \cdots, xt_{k} = x_{0},$$

$$xt_{k+1} = x_{1}, xt_{k+2} = x_{2}, \cdots, xt_{n+k} = x_{n},$$

$$xt_{n+k+1} = x_{n+1}, xt_{n+k+2} = x_{n+2}, \cdots, xt_{n+2k} = x_{n+k}.$$

$$(6)$$

In this section the double subscript for B-splines will be used since it is needed to define the computational algorithm. The B-splines, $N_{ik}(x)$, are constructed by a recursion formula given by

$$N_{1i}(x) = \begin{cases} 1 & \text{if } x_i \le x \le x_{i+1}, \\ 0 & \text{otherwise,} \end{cases}$$
 (7)

and

$$N_{ki}(x) = \frac{(x-x_i) N_{k-1,i}(x)}{x_{i+k-1} - x_i} + \frac{(x_{i+k} - x) N_{k-1,i+1}(x)}{x_{i+k} - x_{i+1}}.$$
 (8)

This formula is referred to in the literature as the Cox-deBoor formula (see Cox [2]).

To evaluate a B-spline of order k for a given $t \in [x_1, x_n]$ the first step is to find $i \in [k+1, n+k]$ so that $t \in [xt_i, xt_{i+1}]$. Note that if $t = x_n$ then i = n+k. Begin by initializing the array N(j, i) = 0 for $i = 1, \ldots, n+k, \ j = 1, \cdots, k$. By the Cox-deBoor formula set $N(1, i) = 1, \ N(1, j) = 0, j \neq i$ and

$$N(j,i) = \frac{(t-xt(i))N(j-1,i)}{xt(i+j-1)-xt(i)} + \frac{(xt(i+j)-t)N(j-1,i+1)}{xt(i+j)-xt(i+1)}.$$
 (9)

In any implementation of this formula care must be exerted due to the potential division by zero problem in the coefficients. We have dropped the function representation here and chosen an array form for the ease of algorithmic representation.

Once an interval index i has been identified with a given t, the evaluation of the final B-spline begins by setting N(1,i) = 1. It will happen that only k values N(k,i-k+1), N(k,i-k+2), ..., N(k,i) will be nonzero. This occurs since there is a triangular dependency of B-splines as

From the discussion here it will become clear why the computations for the relevant B-splines can be reduced to computing (10). In fact, the computation of the B-splines along the right leg of the triangle depends only on the B-splines just above on the right leg. The computation of the B-splines along the hypotenuse of the triangle depends only on the B-splines directly to the right and above along the hypotenuse. Finally, the B-splines within the triangle depend on the B-splines just above and to the right. This can easily be seen from the Cox-deBoor formula and (10). First of all N(1,i) = 1 initializes the recursion. By Cox-deBoor, N(2,i) depends on N(1,i) and N(1,i+1), but N(1, i+1) = 0, so N(2, i) depends only on N(1, i). Now N(2, i-1) depends on N(1, i-1) and N(1,i), but N(1,i-1)=0, so N(2,i-1) depends only on N(1,i-1). Now consider N(3,i). This depends on N(2,i) and N(2,i+1), but N(2,i+1) depends on N(1,i+1) and N(1,i+2), which are both zero. So N(3,i) depends only on N(2,i). N(3,i-1) depends, by Cox-deBoor, on N(2,i-1)and N(2,i), which are both nonzero. Finally, N(3,i-2) depends on N(2,i-2) and N(2,i-1), but N(2, i-2) = 0 since it depends on N(1, i-2) = 0 and N(1, i-1) = 0. This argument can easily be extended down the triangle, but the pattern is clear. Any implementation of the Cox-deBoor formula can then be done by proceeding down the right leg, then down the hypotenuse and then fill in the interior of the triangle.

There is a property of B-splines that can be used to check the calculation of the final k nonzero B-splines. It is called a "partition of unity" property in that

$$\sum_{j=i-k+1}^{i} N(k,j) = 1.$$
 (11)

2.3 Tensor Products of Cubic B-splines in Two Variables

In Section 2.1 the problem of least squares approximation of data by one dimensional B-splines was discussed. In this section the B-spline concept will be extended to two dimensions in order to fit two dimensionsl scattered data by a surface function. In this surface-fitting problem data points (u_r, v_r) , $r = 1, 2, \dots, k$ and values at these points, $z_r = f(u_r, v_r)$, $r = 1, 2, \dots, k$ are given. The surface model used to fit these data points will involve sums of products of B-splines.

To introduce this model, define a rectangle, say R, by $a \le x \le b$, $c \le y \le d$ in the (x,y) plane. The definition does not restrict the data points to the Euclidean plane. They could, for example be angular coordinates, as will be seen later in this report. The rectangle is subdivided by sets of knots x_i , $i=1,2,\cdots,m$ and y_j , $j=1,2,\cdots,n$, where $a < x_1 < x_2 < \cdots < x_m < b$ and $c < y_1 < y_2 < \cdots < y_n < b$. These knots divide the rectangle R into rectangular panels in the plane given by R_{ij} , $i=1,2,\cdots,m$, $j=1,2,\cdots,n$. Then, after extending the knots by eight in each dimension as done in the one dimensional case, a basis set of splines for this pairing of coordinates can be constructed as products of B-splines $N_i(x)N_j(y)$. In fact the surface spline model will be given by

$$s(x,y) = \sum_{i=1}^{m+4} \sum_{i=1}^{n+4} c_{ij} N_i(x) N_j(y),$$
(12)

called a tensor product of splines.

As in the one dimensional case these tensor product splines have a number of advantages. First of all, the basis functions $N_i(x)N_j(y)$ are each non-zero only over a rectangle composed of sixteen panels in a 4×4 arrangement. In particular $N_i(x)N_j(y)$ is non-zero only when $x_{i-4} \leq x \leq x_i$ and $y_{j-4} \leq y \leq y_j$. Next, if (u_r, v_r) , $r = 1, 2, \dots, m$ and values at these points, $z_r = f(u_r, v_r)$, $r = 1, 2, \dots, m$ are given, then the least squares fitting problem can be formulated as finding the least squares solution of

$$\sum_{i=1}^{m+4} \sum_{j=1}^{n+4} c_{ij} N_i(u_r) N_j(v_r) = z_r, \ r = 1, 2, \dots, k.$$
(13)

Again, this can be written in a matrix form as

$$Ac = f, (14)$$

where A is now a matrix with k rows and (m+4)(n+4) columns. If the indexing has been arranged so that $N_i(x)N_j(y)$ and the corresponding c_{ij} are arranged in the order for fixed j, with $i=1,2,\cdots,m+4$, where $j=1,2,\cdots,n+4$, then A has as its element $N_i(u_r)N_j(v_r)$ in row r of column (j-1)(m+4)+i. If the data points are also arranged according to the panels R_{ij} containing them in the order for fixed j, $i=1,2,\cdots,m+4$, where $j=1,2,\cdots,n+4$, then the matrix A takes the form

$$A = \begin{bmatrix} A_{11} & A_{12} & A_{13} & A_{14} \\ & A_{22} & A_{23} & A_{24} & A_{25} \\ & & A_{33} & A_{34} & A_{35} & A_{45} \\ & & \ddots & \ddots & \ddots & \ddots \\ & & & A_{n+1,n+1} & A_{n+1,n+2} & A_{n+1,n+3} & A_{n+1,n+4} \end{bmatrix}.$$
 (15)

Here each A_{rs} is a rectangular matrix of bandwidth four with m+4 columns and the number of rows is equal to the total number of data points in the panels $R_{1r}, R_{2r}, \dots, R_{n+1,r}$. For a full discussion of tensor products of B-splines see de Boor [3], Eberly [8], and Rogers and Adams [13].

2.4 Issues in Computing Tensor Product B-splines and the Relation to Least Squares

Unfortunately this block format for the matrix A is only achievable if the knots, defining the B-splines in the separate dimensions, can be ordered properly. If, for example, two dimensional points are dependent, as in a set of coordinate points (x_i, y_i) , $i = 1, \dots, n$ on a surface, it may not be

possible to order the variables as described above. In that case, the resulting tensor product matrix might be rank deficient and it will not be possible to use the traditional normal equations to solve a least squares problem. Another approach to solving the least squares problem in this case will be discussed below.

In this section, though, we will first discuss the MATLAB function tensor_prod_spl that is given in Appendix C.4. This function creates the tensor product matrix that will be called NM. The input to this function includes the set of knots for both the θ and ϕ angular dimensions as well as the vector arrays of θ and ϕ variables at which the tensor product of B-splines is to be evaluated.

The first thing done for each dimension is to augment the knots at each end. This is done by calls to the MATLAB spline toolbox function augknt, where the order of the B-splines has been set to four for cubic B-splines. This function is called for each of the two knot sets. The matrix for each B-spline, N for θ and M for ϕ , is then created by a call to the MATLAB spline toolbox function specility. This function can create the entire B-spline matrix for a vector of variables, provided the vector is ordered in ascending order. This is possible if the variables in both dimensions are independent. But in the case of coordinates, the variable arrays are not independent. In this case, although one dimension can be sorted in ascending order, the other one has to be ordered in such a way that the coordinates are always linked. Thus, although the θ could be sorted, the ϕ array would not necessarily be sorted properly. Therefore the approach taken in the program was to create the individual B-spline matrices one row at a time.

Once the N and M matrices are created for the θ and ϕ arrays, the tensor product matrix NM can be created. This is done by taking all possible products of the elements from the N and M matrices.

Once the tensor product B-spline matrix is computed, a least squares fit can be done to the scattered spherical coordinate data. Since the matrix NM could be rank deficient, with potentially zero rows or columns, we cannot rely on the standard normal equations for determining the coefficients. We will use the singular value decomposition (SVD) in this case. This is done in the MATLAB function lsq_2D_spline in Appendix C.5. The input to this function is the tensor product matrix NM and the array of radial values at the scattered spherical coordinate points, along with a desired data tolerance. The data tolerance is used to zero the small singular values. In this function the matrix NM, with the number of rows larger than the number of columns, is decomposed by the MATLAB function svd into the product of a column-orthogonal matrix U, a diagonal matrix S, and the transpose V' of a square orthogonal matrix, so that NM = USV'. In order to solve the problem NMc = r we compute c = $VS^+U'r$, where S^+ is the generalized inverse of S and U' is the transpose of U.

2.5 A Knot Selection Algorithm

The selection of B-spline knots for an optimum fit is a nontrivial task. One would be extremely lucky to manually select an initial set of knots and achieve an optimum fit. Therefore we approach knot selection in an iterative fashion that moves us toward an optimum selection. The overall strategy is to place knots in the vicinity of the fit where the local residuals are the largest. This process involves not only the placement algorithm issue but also the iterative stopping criteria.

First we will discuss the placement algorithm. It is based on a suggested strategy by Dierckx [6]. At the beginning of each iteration the assumption is that there exists a current set of knots, call them knts_theta and knts_phi. In the first iteration of the algorithm these would be an initial set chosen by the user. The tensor product of the B-splines are computed at the data points, a least squares fit is made to the data, and the current absolute residuals of the fit are computed at each data point. The current knots divide the θ , ϕ plane into rectangles. Some rectangles have data points and others

don't. For each i from 1 to the number of data points minus 1 the array indices of θ are found so that knts_theta(i) <= theta <= knts_theta(i+1). The sums of the residuals at these points are computed and the maximum of these sums is chosen. A similar calculation is done for ϕ and the maximum of the sums of the squares of the residuals is chosen. The maximum of these two maximums is chosen. If the maximum in the θ dimension is chosen then a knot is added to the knts_theta array, otherwise a knot is added to the knts_phi array. The algorithm steps are given below. For the sake of argument assume that there are currently g θ knots and h ϕ knots that form a grid defined by the rectangles [knts_theta(i), knts_theta(i+1)] × [knts_phi(j), knts_phi(j+1)], $i = 1, \dots, g-1, \ j = 1, \dots, h-1$. Here is the placement algorithm used.

$$R_{k} = r_{k} - S(\theta_{k}, \phi_{k}), \ k = 1, \cdots, n$$

$$\delta_{i}^{\theta} = \sum_{k} \left\{ R_{k}^{2} : \text{knts_theta}(i) \leq \theta(k) \leq \text{knts_theta}(i+1) \right\}$$

$$\delta_{j}^{\phi} = \sum_{k} \left\{ R_{k}^{2} : \text{knts_theta}(i) \leq \theta(k) \leq \text{knts_theta}(i+1) \right\}$$

$$\delta^{\theta} = \max_{1 \leq i \leq g-1} \delta_{i}^{\theta}$$

$$\delta^{\phi} = \max_{1 \leq j \leq h-1} \delta_{j}^{\phi}$$

$$\text{new_knt_theta} = \sum_{k} \left\{ R_{k}^{2} \theta(k) / \delta^{\theta} \right\}, \text{ if } \delta^{\theta} \geq \delta^{\phi}$$

$$\text{new_knt_phi} = \sum_{k} \left\{ R_{k}^{2} \phi(k) / \delta^{\phi} \right\}, \text{ if } \delta^{\phi} \geq \delta^{\theta}$$

The stopping criteria used in this algorithm is based on the use of the R^2 -statistic, called the coefficient of multiple determination (see Draper and Smith [7]). R^2 is the square of the correlation between the vector of observed data, r_k , $k=1,\cdots,n$, and the least squares predicted data, \hat{r}_k , $k=1,\cdots,n$. The statistic satisfies $0 \le R^2 \le 1$. This statistic is often used as a measure of how well the regression equation explains the variation in the data. It is known that building models based on adding terms in the regression equation, care must be taken in using this statistic. However, in the current algorithm the statistic is used in a somewhat non-conventional manner. We use it as a measure of the benefit of adding more knots to the tensor product spline model. The knot selection algorithm is terminated once $R^2 > 0.98$.

3 Surface Point Data Generation

In this section we discuss the experimental method by which boundary data was generated in Euclidean coordinates on each of the phantoms. Figure 3 shows the CMM system used to measure the phantoms and a calibrated sphere used to check the probe. The system is computer controlled and touches an object to be measured at programmend points in order to produce (x, y, z) values at the probed points.

The effective diameter of the CMM probe was measured by first measuring the high quality steel sphere, with a well-calibrated diameter. This sphere is separate from the calibrated one used to generate the sphere data in Section 4.2.1 below. The calibration sphere can be seen in the background of Fig. 4. The difference between the known diameter and the apparent diameter of the measured reference metal sphere gave the calibrated effective probe diameter. A similar pattern of probe touches was used for the measurement of the calibrated sphere and for all of the phantom sphere measurements.



Figure 3: Computer controlled CMM.

The FDA phantoms were created by a molding process and the mold marks on both spheres were visible. These mold marks were used to align the phantoms with the coordinate system of the CMM. The marks were laid out like lines of latitude, leading to the use of a natural nomenclature of latitude and longitude like those of the Earth. For each phantom, the north pole was chosen to be the one with darker, deeper, or more obvious mold marks.

The phantoms were held by a vacuum chuck (see Fig. 4) to minimize distortion and reduce the chance of damaging the spheres' surfaces. The chuck has a shallow cone, so contact is made around a circle of latitude at about -45° . The wall vacuum was strong enough to hold the spheres sufficiently that they did not move significantly, as shown by the repeatable results from run to run at the 1 μ m level.

Visually, the pink sphere was noticeably out of round, in the shape of an oblate spheroid. A dial caliper gave diameter measurements given in Table 1.



Figure 4: Lung nodule phantom on the vacuum chuck.

	Equatorial	Polar
Pink	$20.0 \mathrm{\ mm}$	18.4 mm
Green	$20.0 \; \mathrm{mm}$	$20.1 \mathrm{mm}$

Table 1: Dial Caliper Measurements of Phantoms' Diameters

The uncertainty of caliper measurements on hard steel surfaces is about 0.1 mm, and is estimated to be about 0.3 mm on the sample spheres due to the potential that the contact force would distort the soft surfaces of the spheres (all estimated uncertainties were k=2 expanded uncertainties according to the guidelines of Taylor and Kuyatt [18]). Since the CMM is primarily used to measure metallic artifacts produced by machining operations, we will see in this report some of the possible consequences of using the CMM to measure the non-metallic phantoms.

Two probing experiments were performed on each of the phantoms and the calibrated sphere. They created what we will call a coarse data set and a dense data set. In the next two sections we will discuss the methods used in the two experiments since they were slightly different.

3.1 Experiment 1: Coarse Data Set

In this experiment the plan was to measure each phantom on the CMM three times in each position, with 61 coordinate points per hemisphere. For the green sphere, each measurement set consisted of three separate sets of points: North pole up, South pole up, and prime meridian/equator intersection up. The pink sphere could not be held sideways in the first experiment, as the out-of-roundness prevented an effective vacuum seal. Therefore, a measurement data set for this sphere had only the North pole up and South pole up data.

Although 183 points were measured on the green sphere, 121 were selected in order to make the data sets for the green and pink phantoms consistent in terms of probed points. All 121 points from both hemispheres were merged together, and the plots in Fig. 5 and 6 show the radial deviation from a best-fit sphere for the full data sets. The figures show the radial residuals obtained by fitting sphere models to the Green and Pink data with an algorithm ordinarily used during sphere calibration work. In particular, they represent the residual errors between the distance from the fitted sphere center to the probed points and the fitted radius of the sphere model. The residuals, in the case of the Green phantom, range from approximately -0.1 mm to +0.1 mm, whereas the residuals, in the case of the Pink phantom, range from approximately -0.57 mm to +0.23 mm. This indicates the non-spherical nature of the Pink phantom. A separate nonlinear algorithm for sphere fitting will be described in Section 4. It produces consistent results. Figure 7 shows the typical distribution of the probe points on a sphere. The plot is a transparency so that probe points on the opposite side of the sphere are visible. The calibated sphere on the shaft, with a diameter of 19.05 mm, was also measured with 121 points, repeated five times.

3.2 Experiment 2: Dense Data Set

In this experiment 181 points were taken on the phantoms and the calibrated sphere, with five repeats in each position. The positions were taken the same as those in Experiment 1. That is, the alignment was selected with North pole up (position 1), South pole up (position 2), and prime meridian/equator intersection up (position 3). In this case it was possible to hold the Pink phantom in the sidways position 3. The calibrated sphere was measured in only one position, with five repeats, since it was permanently mounted on a support shaft.

Green Sphere Residuals

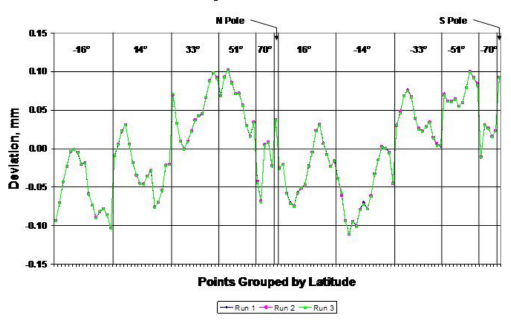


Figure 5: Residual Measure CMM Errors for the Green Phantom Sphere: Experiment 1.

Pink Sphere Residuals

Figure 6: Residual Measure CMM Errors for the Pink Phantom Sphere: Experiment 1.

Points Grouped by Latitude

--- Run 1 --- Run 2 --- Run 3

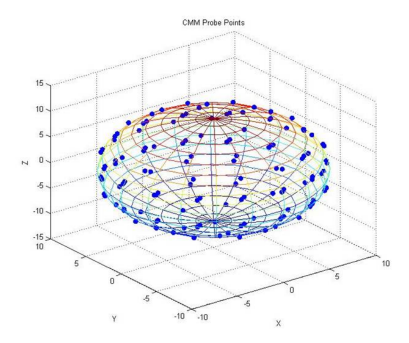


Figure 7: Distribution of the CMM Probe Points on a Sphere: Experiment 1.

4 Data Modeling and Volume Estimation

In this section two forms of data modeling will be discussed. Since the phantoms seemed to be nearly spherical the natural tendency was to first consider fitting a spherical model to each phantom and estimating the volume of the fitted spheres. Second, in order to develop a potentially more accurate volume estimation model the surface data was fit using tensor products of B-splines and the volumes estimated by the Divergence Theorem.

4.1 A Spherical Model

Since the phantoms were nearly spherical it was natural, as noted above, to consider how close the data could be modeled by first assuming spherical models for the data. The calibrated sphere, of course, could clearly be modeled with a spherical model.

In particular, let $c = (c_1, c_2, c_3, c_4)$ and set

$$f(x, y, z, c) = (x - c_1)^2 + (y - c_2)^2 + (z - c_3)^2 - c_4^2.$$
(17)

The unknown parameters c_1 , c_2 , c_3 represent the center of the sphere and c_4 is the radius. All of the data were measured in millimeters so that the parameters naturally have millimeter units. Define

$$S(c) = \sum_{i=1}^{n} \left\{ (x_i - c_1)^2 + (y_i - c_2)^2 + (z_i - c_3)^2 - c_4^2 \right\}.$$
 (18)

Since the function S is a nonlinear implicit function of the parameters we needed to use a nonlinear

mimimization algorithm to solve the problem

$$\min_{c} S(c). \tag{19}$$

There are a number of algorithms for fitting least squares models to data on geometric shapes. The reader can consult Shakarji [16]. There are also various algorithms for minimizing general nonlinear functions, such as (18). All of the algorithms involve iterative minimization of some form. Many require computing derivatives of the objective function in order to generate minimization search directions. Others do not involve derivatives but may be somewhat slower in the minimization search. The algorithm selected here, because of the relatively few parameters involved, and the fact that derivatives are not required, is a form of polygon search method called the Nelder-Mead method (see Sauer [14]). It should be noted here that the Newton's method, requiring derivatives, was initially used to estimate the parameters, but the Nelder-Mead tended to produce the smallest value to (18).

The Nelder-Mead method begins with an initial vector guess for the minimum in \mathbb{R}^4 , although the method works for \mathbb{R}^n , but due to the "curse of dimension", becomes prohibitive beyond \mathbb{R}^{10} . From the initial vector guess a polygon of five vertices is built. The function evaluations at the vertices, $w_i = S(v_i)$, of the polygon are tested and put into ascending order, $w_1 < w_2 < \cdots < w_5 = w_h$. The polygon vertex, v_h , with the largest function value, w_h , is replaced as follows. The centroid, \overline{v} , of the face of the polygon that does not contain v_h is chosen. The function (18) is evaluated at, v_r , the reflection point about the centroid, given by $v_r = 2\overline{v} - v_h$, is selected. If the value $w_r = S(v_r)$ lies in the range $w_1 < w_r < w_4$, then v_r replaces v_h , the function values of the vertices are again sorted, and the iterative step is repeated. If $w_r < w_1$, extrapolation is continued in the direction of w_r by setting $v_e = 3\overline{v} - 2v_h$. The better of v_e or v_r is selected to replace v_h . Finally, if $w_r > w_n$ then a contraction of points is attempted. Two points are selected. The outer contraction point is taken as $v_{oc} = 1.5\overline{v} - 0.5v_r$ and the inner contraction is $v_{ic} = 0.5\overline{v} + 0.5v_h$. The function (18) is evaluated at both contraction points. If there is no reduction in the value of (18) then the polygon is shrunk by a factor of 2 in the direction of the current minimum, v_1 . The iterations of sorting and testing continue until the volume of the polygon becomes less than some prespecified tolerance. Given a set of (x, y, z) points, a MATLAB code to estimate the center and radius of the best fit sphere is given in Appendix A. The output of the program produces a vector that is the median value of the vertex values of the final polyhedron, whose volume satisfies the tolerance specification. The volume is computed as a determinant involving the polygon vertices.

The uncertainties of the estimated center and radius were computed using the methods proposed in Draper and Smith [7] for nonlinear regression. In particular, if $\hat{c} = (\hat{c}_1, \hat{c}_2, \hat{c}_3, \hat{c}_4)$, then define the matrix with elements

$$\hat{Z}_{ij} = \frac{\partial f}{\partial c_i}(x_i, y_i, z_i, \hat{c}), \quad i = 1, \dots, n, \quad j = 1, 2, 3, 4.$$
 (20)

The matrix \hat{Z} is an $n \times 4$ matrix. The *i*-th row is given by

$$\hat{Z}_{i} = \begin{bmatrix} -2(x_{i} - \hat{c}_{1}) & -2(y_{i} - \hat{c}_{2}) & -2(z_{i} - \hat{c}_{3}) & -2\hat{c}_{4} \end{bmatrix}.$$
(21)

If we then form $\hat{Z}'\hat{Z}$ the standard error of \hat{c}_i , s.e. (\hat{c}_i) , is given by the *i*-th diagonal element of $(\hat{Z}'\hat{Z})^{-1}s^2$ where $s=S\left(\hat{c}\right)/(n-4)$. The expanded uncertainty is given by 2s.e. (\hat{c}_i) as defined in Taylor and Kuyatt [18]. The uncertainty limits of \hat{c}_i are $\hat{c}_i \pm 2$ s.e. (\hat{c}_i) . All units are in millimeters except volume, which is in cubic millimeters.

4.2 Computational Results for the Spherical Model

Tables 2, 3, and 4 present the results obtained by fitting spherical models to the data from the calibrated sphere and the two phantoms. In Experiment 1 there were three data sets for each of the Green and Pink phantoms, designated Green 4, 5, and 6 and Pink 5, 6, and 7. Since the output of the Nelder-Mead algorithm was a final polytope surrounding the minimum with the polytope vertices representing the final parameter esimates, the final reported parameters were selected as the median values of the vertex values. These are the first four entries in the table: Center x, Center y, Center z, and Radius. The units are millimeters. The fifth table entry is the spherical volume based on the median radius value in cubic millimeters. The radius residuals were computed as

$$\operatorname{resid}_{i} = \sqrt{(x_{i} - \hat{c}_{1})^{2} + (y_{i} - \hat{c}_{2})^{2} + (z_{i} - \hat{c}_{3})^{2}} - \hat{c}_{4}. \tag{22}$$

The sixth entry in the tables gives the mean value of these residuals. The unit is millimeters. The seventh and eighth table entries give the mean and standard deviation of the absolute values of the residuals in millimeters. The ninth through the twelfth entries are the expanded uncertainties for the estimated Center x, Center y, Center z, and radius.

The results in these sections on spherical model fitting involve strictly fitting the non-linear model (22) to the sparse data sets generated by the CMM. This process does not involve the B-spline algorithm used later in this report. Therefore any differences between coarse and dense results are most likely the consequence of the point selection in the metrology of the artifacts.

4.2.1 Results for the Spherical Model Fit to the Calibrated Sphere with Coarse and Dense Data

Table 2 gives the results of the fit of the sphere model to both the coarse (121 points) and the dense (181 points) data sets for the calibrated sphere. In both cases the average of the five repeat data sets was used for the fitting process. As can be seen, the two volume estimates differ by approximately 0.04%. The radii estimates differ by about 0.01%. It would seem that these differences are within the operating regime of the CMM. Although the estimated centers are slightly different, all other values are of the same order of magnitude. Based on these results we can accept that the CMM is producing accurate position data for spherical metallic artifacts. Tables 3 through 6, however, begin to show the consequences involved with attempting to measure slightly non-spherical and non-metallic artifacts with the CMM.

4.2.2 Results for the Spherical Model Fit to the Phantoms with Coarse Data

From Tables 3 and 4 it is clear that the Green phantom is more spherical than the Pink phantom as indicated by the residuals and the expanded uncertainties. This simply confirms the fact that the Pink phantom was more difficult to measure using the vacuum chuck due to its lack of sphericity. For example, the Mean Radial Residual for the Pink data is two orders of magnitude larger than for the Green data. The Standard Deviations for the Pink data are an order of magnitude greater, whereas all of the Expanded Uncertainties are two orders of magnitude greater. It is not clear how much the phantom material affected the results since it was difficult to set up a specific probe test for metallic versus phantom material. The artifacts would have to have been exactly the same size, positioned at exactly the right location, and probed at exactly the same coordinates to separate those factors from the material factor difference. There were no such comparable artifacts. We can make a guess, though, that there might be some affect due to probe force against the non-metalic material if we look at Table 2 and Table 3, the most spherical of the two phantoms. The Expanded Uncertainties

Spherical Fit Results for Calibrated Sphere			
Properties	Coarse	Dense	
Center x	0.2803×10^{-4}	-0.1331×10^{-2}	
Center y	-0.4881×10^{-3}	-0.2034×10^{-2}	
Center z	-0.6642×10^{-2}	-0.2564×10^{-2}	
Radius	9.5326	9.5314	
Est. Volume	3628.4	3627.1	
Mean Rad. Residual	0.3232×10^{-4}	0.8720×10^{-4}	
Mean Abs. Rad. Residual	0.1646×10^{-2}	0.1700×10^{-2}	
Stand. Dev. Rad. Residual	0.2156×10^{-2}	0.22302×10^{-2}	
Stand. Dev. Abs. Rad. Residual	0.1384×10^{-2}	0.1441×10^{-2}	
Expanded Uncert. Center x	0.3676×10^{-9}	0.2779×10^{-9}	
Expanded Uncert. Center y	0.3675×10^{-9}	0.2777×10^{-9}	
Expanded Uncert. Center z	0.9313×10^{-9}	0.7059×10^{-9}	
Expanded Uncert. Radius	0.2394×10^{-9}	0.1822×10^{-9}	

Table 2: Results of a Spherical Fit to Coarse and Dense Data for Calibrated Sphere

for the sphere fit to the calibrated metalic sphere and the Expanded Uncertainties for the sphere fit to the Green phantom data differ by five to six orders of magnitude. Since the repeatability of the CMM measurements is at the 1 μ m level, it is likely then that material difference had some significant affect on the difference in the uncertainties. It is a guess that this and the non-spherical shape of the Pink phantom account for a large part of the differences between the Pink phantom Expanded Uncertainties in Table 4, the Green phantom Expanded Uncertainties in Table 3, and the calibrated sphere Expanded Uncertainties in Table 2.

4.2.3 Results for the Spherical Model Fit to the Phantoms with Dense Data

We want to make some observations about the resulting measurements and the spherical fits to the dense data. If we first look at the residuals for the sphere fits to the dense data sets we see an interesting phenomenon occuring in the fits for both the Green and Pink phantoms in position 3 (prime meridian/equator intersection up). There appears to be a definite periodic oscillation in the residual data in Figures 10 and 13. It is not clear what in the measurement process produced these residual oscillations, although the points selected would have been chosen along longitudes as the CMM moved from top to bottom of the positioned phantom. These periodicities are possible considering the North to South symmetry across the equator in the sphere model fit. Next we examined the possible affects of these residual oscillations on the sphere volume estimates.

If we compare the volume results for the coarse Green and dense Green data we get the following. From Table 3 and Table 5 there appears to be only a change of about 1% from Green 4 volume to Dense Green 1 and similarly for Green 5 to Dense Green 2. Green 6 and Dense Green 3 volumes differ by about 0.08%. Therefore the oscillations in the residuals for the Dense Green 3 data do not seem to have affected the volume estimate. The residuals themselves in Fig. 10 oscillate between -0.15 mm and 0.1 mm. However, when we compare the Pink phantom volume estimates we get a vastly different result. Whereas the residuals in Figures 11 and 12 fall within the range of -0.4 to 0.15 mm, the residuals as shown in Fig. 13 oscillate between -0.6 mm and 0.6 mm, about six times the residual oscillations of the Dense Green 3 data. The damping of the oscillations may have been due to the slight non-spherical shape of the Pink phantom, although this is a conjecture. The volume

Spherical Fit Results				
Properties	Green 4	Green 5	Green 6	
Center x	0.2150×10^{-2}	0.2128×10^{-2}	0.2579×10^{-2}	
Center y	-0.4694×10^{-2}	-0.4813×10^{-2}	-0.4611×10^{-2}	
Center z	-0.5482×10^{-1}	-0.5489×10^{-1}	-0.5442×10^{-1}	
Radius	10.1124	10.1121	10.1118	
Est. Volume	4331.7	4331.3	4330.8	
Mean Rad. Residual	-1.7530×10^{-4}	-1.764×10^{-4}	-1.4684×10^{-4}	
Mean Abs. Rad. Residual	0.4592×10^{-1}	0.4607×10^{-1}	0.4586×10^{-1}	
Stand. Dev. Rad. Residual	0.5521×10^{-1}	0.5538×10^{-1}	0.5518×10^{-1}	
Stand. Dev. Abs. Rad. Residual	0.3038×10^{-1}	0.3043×10^{-1}	0.3041×10^{-1}	
Expanded Uncert. Center x	0.1849×10^{-3}	0.9352×10^{-4}	0.1844×10^{-3}	
Expanded Uncert. Center y	0.1849×10^{-3}	0.9351×10^{-4}	0.1844×10^{-3}	
Expanded Uncert. Center z	0.2249×10^{-3}	0.1138×10^{-3}	0.2243×10^{-3}	
Expanded Uncert. Radius	0.6551×10^{-4}	0.3314×10^{-4}	0.6535×10^{-4}	

Table 3: Results of a Spherical Fit to Coarse Data for the Green Phantom

Spherical Fit Results				
Properties	Pink 5	Pink 6	Pink 7	
Center x	-0.8612×10^{-2}	-0.8235×10^{-2}	-0.8373×10^{-2}	
Center y	0.1572×10^{-1}	0.1547×10^{-1}	0.1555×10^{-1}	
Center z	0.2315×10^{-1}	0.2365×10^{-1}	0.2359×10^{-1}	
Radius	9.7337	9.7334	9.7329	
Est. Volume	3863.02	3862.61	3862.03	
Mean Rad. Residual	-0.2222×10^{-2}	-0.2721×10^{-2}	-0.2356×10^{-2}	
Mean Abs. Rad. Residual	0.1748	0.1746	0.1746	
Stand. Dev. Rad. Residual	0.2128	0.2127	0.2126	
Stand. Dev. Abs. Rad. Residual	0.1203	0.1203	0.1203	
Expanded Uncert. Center x	0.3617×10^{-1}	0.3607×10^{-1}	0.3602×10^{-1}	
Expanded Uncert. Center y	0.3635×10^{-1}	0.3626×10^{-1}	0.3620×10^{-1}	
Expanded Uncert. Center z	0.4428×10^{-1}	0.4415×10^{-1}	0.4407×10^{-1}	
Expanded Uncert. Radius	0.1287×10^{-1}	0.1283×10^{-1}	0.1281×10^{-1}	

Table 4: Results of a Spherical Fit to Coarse Data for the Pink Phantom

Spherical Fit Results				
Properties	Dense Green 1	Dense Green 2	Dense Green 3	
Center x	-0.1073×10^{-1}	0.2112×10^{-1}	-0.1321×10^{-1}	
Center y	0.5112×10^{-1}	0.8661×10^{-2}	-0.4700×10^{-1}	
Center z	0.2550	0.4723×10^{-1}	0.2139	
Radius	10.0706	10.0710	10.1146	
Est. Volume	4278.1	4278.7	4334.5	
Mean Rad. Residual	-0.9344×10^{-4}	-0.7116×10^{-4}	-0.8881×10^{-4}	
Mean Abs. Rad. Residual	0.3502×10^{-1}	0.3267×10^{-1}	0.3980×10^{-1}	
Stand. Dev. Rad. Residual	0.4566×10^{-1}	0.3935×10^{-1}	0.4904×10^{-1}	
Stand. Dev. Abs. Rad. Residual	0.2918×10^{-1}	0.2181×10^{-1}	0.2850×10^{-1}	
Expanded Uncert. Center x	0.5320×10^{-4}	0.2980×10^{-4}	0.7197×10^{-4}	
Expanded Uncert. Center y	0.5324×10^{-4}	0.2980×10^{-4}	0.7158×10^{-4}	
Expanded Uncert. Center z	0.1233×10^{-3}	0.7006×10^{-4}	0.1670×10^{-3}	
Expanded Uncert. Radius	0.3058×10^{-4}	0.1778×10^{-4}	0.4158×10^{-4}	

Table 5: Results of a Spherical Fit to Dense Data for the Green Phantom

differences between Pink 5 in Table 4 and Dense Pink 1 in Table 6 as well as Pink 6 and Dense Pink 2 in the same tables are approximately 6%. The volume estimates between the Pink 7 results in Table 4 and the Dense Pink 3 results in Table 6, however, show about a 7% decrease. This last comparison may not be a fair one in that the Pink 7 data did not have data for the Pink phantom in position 3. Table 1 shows that the diameter of the Pink phantom falls approximately between 20 mm and 18.5 mm. This would imply that the sphere model should have a volume between 4188.8 mm³ and 3315.2 mm³. Table 4 for the Pink phantom shows that for the coarse data the volume estimate falls from 3862 mm³ to about 3863 mm³. These volumes are within the expected range. Whereas the volume estimates for Dense Pink 1, Dense Pink 2, and Dense Pink 3 from Table 6 are also within the expected range. The Pink phantom in position 3 may have shown difficulties in the data acquisition phase for the CMM. This could explain the lower volume estimate. In particular, an approximate 4% difference in radius estimate between Dense Pink 3 and Dense Pink 2 causes an approximate 13% decrease in volume estimate. This points how seemeingly small differences in linear parameter estiamtes can make significant volumetric differences.

4.3 Data Modeling by B-Splines

In the following sections we plan to introduce a more general approach to estimate the phantom volumes by modeling the surfaces with B-splines and using the Divergence Theorem to estimate the volume, although in the case of the Green phantom the sphere model and the B-spline model will confirm its near spherical nature.

4.3.1 Euclidean to Spherical Coordinate Transformation

The current study involves the modeling of the sets of CMM data points. These unstructured sets of (x, y, z) data points are called point clouds. Therefore, in this section we assume that we are given a set of n points, (x_i, y_i, z_i) , $i = 1, 2, \dots, n$, for some n, on a surface. As measured, these points are given relative to the CMM origin. The first step is to develop an origin for the data by using the

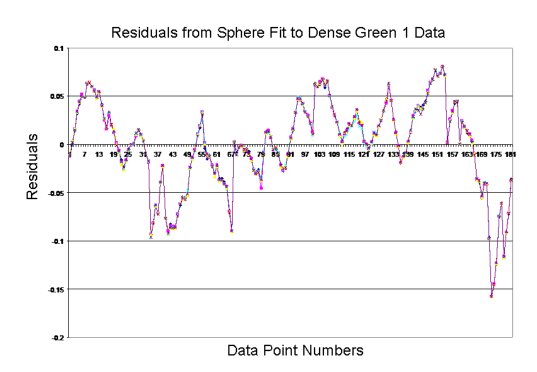


Figure 8: Residuals for the Sphere Fit to the Dense Green Data Set 1.

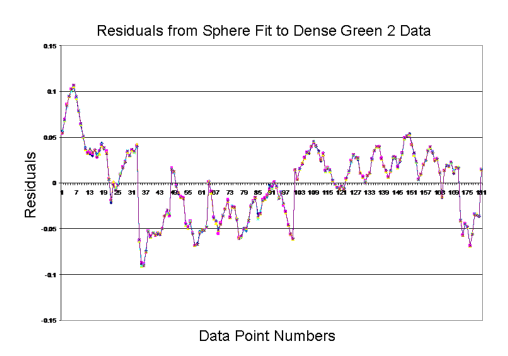


Figure 9: Residuals for the Sphere Fit to the Dense Green Data Set 2.

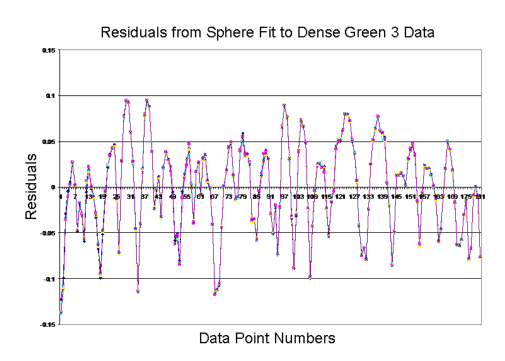


Figure 10: Residuals for the Sphere Fit to the Dense Green Data Set 3.

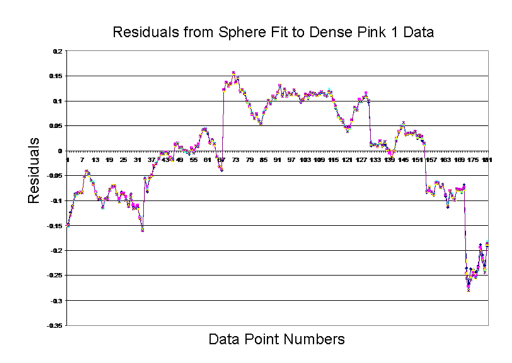


Figure 11: Residuals for the Sphere Fit to the Dense Pink Data Set 1.

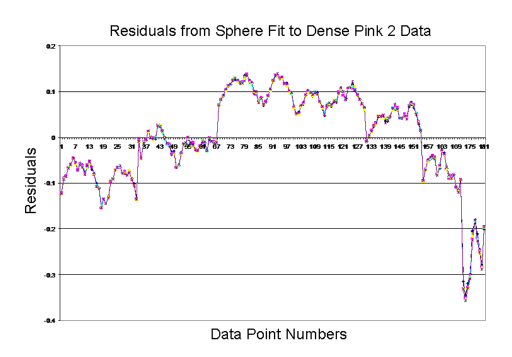


Figure 12: Residuals for the Sphere Fit to the Dense Pink Data Set 2.

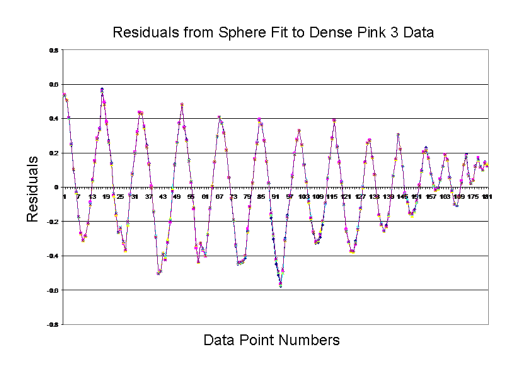


Figure 13: Residuals for the Sphere Fit to the Dense Pink Data Set 3.

Spherical Fit Results				
Properties	Dense Pink 1	Dense Pink 2	Dense Pink 3	
Center x	0.1472×10^{-1}	0.4573×10^{-1}	0.1496×10^{-1}	
Center y	0.1448×10^{-1}	0.1557×10^{-1}	0.6815×10^{-1}	
Center z	0.6848	0.7473	-0.3443	
Radius	9.9234	9.9370	9.5045	
Est. Volume	4093.2	4110.1	3596.5	
Mean Rad. Residual	-0.4393×10^{-3}	-0.4859×10^{-3}	-0.3282×10^{-2}	
Mean Abs. Rad. Residual	0.7993×10^{-1}	0.7989×10^{-1}	0.2149	
Stand. Dev. Rad. Residual	0.9764×10^{-1}	0.1001	0.2567	
Stand. Dev. Abs. Rad. Residual	0.5576×10^{-1}	0.5996×10^{-1}	0.1397	
Expanded Uncert. Center x	0.1060×10^{-2}	0.1162×10^{-2}	0.4686×10^{-1}	
Expanded Uncert. Center y	0.1057×10^{-2}	0.1157×10^{-2}	0.5073×10^{-1}	
Expanded Uncert. Center z	0.2398×10^{-2}	0.2610×10^{-2}	0.1178	
Expanded Uncert. Radius	0.5662×10^{-3}	0.6129×10^{-3}	0.3137×10^{-1}	

Table 6: Results of a Spherical Fit to Dense Data for the Pink Phantom

center-of-data mass of the point cloud. This point is given by

$$\overline{x} = \frac{1}{n} \sum_{i=1}^{n} x_i$$

$$\overline{y} = \frac{1}{n} \sum_{i=1}^{n} y_i$$

$$\overline{z} = \frac{1}{n} \sum_{i=1}^{n} z_i$$
(23)

We note here, implicitly, that we have assumed that the point cloud can be enclosed in a spherical bounded region. In order to use spherical coordinates to represent points on the boundary of a surface we need to restrict our analysis to surfaces that are called star-shaped. These are surfaces in which a ray drawn from the center-of-data mass intersects the boundary in a unique point.

To each point in the point cloud there is a vector from $(\overline{x}, \overline{y}, \overline{z})$ to (x_i, y_i, z_i) given by $V_i = (x_i, y_i, z_i) - (\overline{x}, \overline{y}, \overline{z})$. Furthermore, any point within the bounding sphere can be identified by spherical coordinates of the form $Q(\theta, \phi) = (r \sin \theta \cos \phi, r \sin \theta \sin \phi, r \cos \theta)$, where $x = r \sin \theta \cos \phi, y = r \sin \theta \sin \phi, z = r \cos \theta$, for $0 \le \theta \le \pi, 0 \le \phi \le 2\pi$. θ is referred to as the colatitude and ϕ is referred to as the azimuth. Table 7 associates the three dimensional octants with their spherical coordinates, where we have suppressed r. The conversion to spherical coordinates was necessary since tensor product B-splines need to be defined on rectangular regions. Therefore, the Euclidean coordinates were converted to θ , ϕ angles on a rectangle $[0, \pi] \times [0, 2\pi]$. The height at each θ , ϕ was taken as the estimated radius from the center-of-data mass of all of the Euclidean coordinates.

Points, not necessarily on a spherical surface of radius r, will in general be a function of θ and ϕ . At this point we will start with an (x, y, z) point from the point cloud and compute the value of $(r(\theta, \phi), \theta, \phi)$ relative to the center-of-data mass. There are eight cases to consider. In all cases we assume that $r(\theta, \phi) \geq 0$. r will always be taken so that $r^2 = x^2 + y^2 + z^2$.

Before considering individual cases there are some general relations that apply to multiple cases. For example, in cases one through four in Table 7 we have $z \ge 0$ so that $0 \le z \le r$ and we will

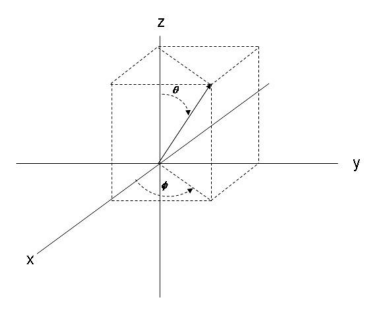


Figure 14: Spherical Coordinate angles. θ is the colatitude and ϕ is the azimuth.

3-D Octants to Spherical Coordinates				
Octant Number	Cartesian Coordinate	$Spherical\ Coordinate$		
1	$x \ge 0, y \ge 0, z \ge 0$	$0 \le \theta \le \pi/2, 0 \le \phi \le \pi/2$		
2	$x < 0, y \ge 0, z \ge 0$	$0 \le \theta \le \pi/2, \pi/2 \le \phi \le \pi$		
3	$x < 0, y < 0, z \ge 0$	$0 \le \theta \le \pi/2, \pi \le \phi \le 3\pi/2$		
4	$x \ge 0, y < 0, z \ge 0$	$0 \le \theta \le \pi/2, 3\pi/2 \le \phi \le 2\pi$		
5	$x \ge 0, y \ge 0, z < 0$	$\pi/2 \le \theta \le \pi/2, 0 \le \phi \le \pi/2$		
6	$x < 0, y \ge 0, z < 0$	$\pi/2 \le \theta \le \pi, \pi/2 \le \phi \le \pi$		
7	x < 0, y < 0, z < 0	$\pi/2 \le \theta \le \pi, \pi \le \phi \le 3\pi/2$		
8	$x \ge 0, y < 0, z < 0$	$\pi/2 \le \theta \le \pi, 3\pi/2 \le \phi \le 2\pi$		

Table 7: Octant Equivalence between Euclidean and Spherical Coordinates

always have $0 \le z/r \le 1$. For MATLAB this implies that

$$0 \le \theta = \arccos\left(\frac{z}{r}\right) \le \frac{\pi}{2}.\tag{24}$$

In cases five through eight in Table 7 z < 0 so that $-1 \le z/r < 0$. Then

$$\frac{\pi}{2} \le \theta = \arccos\left(\frac{z}{r}\right) \le \pi.$$
 (25)

There are also some special cases. In particular, if x = 0, y = 0, z = r, then we set $\theta = 0$, and, if x = 0, y = 0, z = -r, then we set $\theta = \pi$.

In Octant 1, $x \ge 0$, $y \ge 0$, $z \ge 0$. We assume (0,0,0) will not be considered. We estimate $r = \sqrt{x^2 + y^2 + z^2}$. We have θ from (24), so now we want to compute the associated ϕ . For $\theta \in (0, \pi/2)$ we have $0 < x/(r \sin \theta) < 1$. Then

$$0 < \phi = \arccos\left(\frac{x}{r\sin\theta}\right) < \frac{\pi}{2}.\tag{26}$$

In Octant 2, $x < 0, y \ge 0, z \ge 0$. Set $r = \sqrt{x^2 + y^2 + z^2}$ and θ from (24). Since $0 \le \theta < \pi/2$, then $-1 < x/(r\sin\theta) < 0$. Therefore

$$\frac{\pi}{2} < \phi = \arccos\left(\frac{x}{r\sin\theta}\right) < \pi,$$
 (27)

the desired interval.

In Octant 3, x < 0, y < 0, $z \ge 0$. Set $r = \sqrt{x^2 + y^2 + z^2}$ and θ from (24). We want $\pi \le \phi \le 3\pi/2$. Since $-1 < x/(r\sin\theta) < 0$ then $\pi/2 < \phi_1 = \arccos(x/(r\sin\theta)) < \pi$ and

$$\pi < \phi = \phi_1 + \frac{\pi}{2} < \frac{3\pi}{2}.\tag{28}$$

In Octant 4, $x \ge 0$, y < 0, $z \ge 0$. Set $r = \sqrt{x^2 + y^2 + z^2}$ and θ from (24). We want $3\pi/2 \le \phi \le 2\pi$. Since $0 < x/(r\sin\theta) < 1$, then $0 < \phi_1 = \arccos(x/(r\sin\theta)) < \pi/2$

$$\frac{3\pi}{2} \le \phi = \phi_1 + \frac{3\pi}{2} \le 2\pi. \tag{29}$$

In Octant 5, $x \ge 0, y \ge 0, z < 0$. Set $r = \sqrt{x^2 + y^2 + z^2}$ and solve for θ from (24). Since $0 < x/(r\sin\theta) < 1$,

$$0 < \phi = \arccos\left(\frac{x}{r\sin\theta}\right) \le \frac{\pi}{2}.\tag{30}$$

In Octant 6, $x < 0, y \ge 0, z < 0$. Set $r = \sqrt{x^2 + y^2 + z^2}$ and solve for θ from (24). We want $\pi/2 \le \phi \le \pi$. Since $-1 < x/(r\sin\theta) < 0$

$$\frac{\pi}{2} < \phi = \arccos\left(\frac{x}{r\sin\theta}\right) \le \pi.$$
 (31)

In Octant 7, x < 0, y < 0, z < 0. Set $r = \sqrt{x^2 + y^2 + z^2}$ and solve for θ from (24). We want $\pi \le \phi \le 3\pi/2$, but $\pi/2 \le \phi_1 = \arccos(x/(r\sin\theta)) \le \pi$ so we set

$$\pi \le \phi = \phi_1 + \frac{\pi}{2} \le \pi. \tag{32}$$

In Octant 8, $x \ge 0, y < 0, z < 0$. Set $r = \sqrt{x^2 + y^2 + z^2}$ and solve for θ from (24). We want $3\pi/2 \le \phi \le 2\pi$, but $0 \le \phi_1 = \arccos(x/(r\sin\theta)) \le \pi/2$ so we set

$$\frac{3pi}{2} < \phi = \phi_1 + \frac{3\pi}{2} < 2\pi. \tag{33}$$

Transformation Algorithm

The algorithm is relatively straightforward.

- 1. Load the measured points (x_i, y_i, z_i) into three arrays.
- 2. Compute length of the arrays and set = n. This assumes the arrays are of equal length.
- 3. Compute the center of data mass

$$\overline{x} = \frac{1}{n} \sum_{i=1}^{n} x_i,$$

$$\overline{y} = \frac{1}{n} \sum_{i=1}^{n} y_i,$$

$$\overline{z} = \frac{1}{n} \sum_{i=1}^{n} z_i.$$
(34)

4. Reset the (x_i, y_i, z_i) points as

$$x_{i} = x_{i} - \overline{x},$$

$$y_{i} = y_{i} - \overline{y},$$

$$z_{i} = z_{i} - \overline{z}.$$

$$(35)$$

- 5. Begin loop from i = 1 to n 5.1. Compute $r_i = \sqrt{x_i^2 + y_i^2 + z_i^2}$.
 - 5.2. Test for the special cases.
 - 5.2.1. If $x_i = 0, y_i = 0, z_i = r_i$, set $\theta_i = 0, \phi_i = 0$.
 - 5.2.1. If $x_i = 0, y_i = 0, z_i = -r_i$, set $\theta_i = \pi, \phi_i = 0$.
 - 5.3. Test for the Octants
 - 5.3.1. If $x_i \ge 0, y_i \ge 0, z_i \ge 0$

$$\theta_{i} = \arccos\left(\frac{z_{i}}{r_{i}}\right),$$

$$\phi_{i} = \arccos\left(\frac{x_{i}}{r_{i}\sin\theta_{i}}\right).$$
(36)

5.3.2. If $x_i < 0, y_i \ge 0, z_i \ge 0$

$$\theta_{i} = \arccos\left(\frac{z_{i}}{r_{i}}\right),$$

$$\phi_{i} = \arccos\left(\frac{x_{i}}{r_{i}\sin\theta_{i}}\right).$$
(37)

5.3.3. If $x_i < 0, y_i < 0, z_i \ge 0$

$$\theta_{i} = \arccos\left(\frac{z_{i}}{r_{i}}\right),$$

$$\phi_{i} = \frac{\pi}{2} + \arccos\left(\frac{x_{i}}{r_{i}\sin\theta_{i}}\right).$$
(38)

5.3.4. If $x_i \ge 0, y_i < 0, z_i \ge 0$

$$\theta_{i} = \arccos\left(\frac{z_{i}}{r_{i}}\right),$$

$$\phi_{i} = \frac{3\pi}{2} + \arccos\left(\frac{x_{i}}{r_{i}\sin\theta_{i}}\right).$$
(39)

5.3.5. If $x_i \ge 0, y_i \ge 0, z_i < 0$

$$\theta_{i} = \arccos\left(\frac{z_{i}}{r_{i}}\right),$$

$$\phi_{i} = \arccos\left(\frac{x_{i}}{r_{i}\sin\theta_{i}}\right).$$
(40)

5.3.6. If $x_i < 0, y_i \ge 0, z_i < 0$

$$\theta_{i} = \arccos\left(\frac{z_{i}}{r_{i}}\right),$$

$$\phi_{i} = \arccos\left(\frac{x_{i}}{r_{i}\sin\theta_{i}}\right).$$
(41)

5.3.7. If $x_i < 0, y_i < 0, z_i < 0$

$$\theta_{i} = \arccos\left(\frac{z_{i}}{r_{i}}\right),$$

$$\phi_{i} = \frac{\pi}{2} + \arccos\left(\frac{x_{i}}{r_{i}\sin\theta_{i}}\right).$$
(42)

5.3.8. If $x_i \ge 0, y_i < 0, z_i < 0$

$$\theta_{i} = \arccos\left(\frac{z_{i}}{r_{i}}\right),$$

$$\phi_{i} = \frac{3\pi}{2} + \arccos\left(\frac{x_{i}}{r_{i}\sin\theta_{i}}\right).$$
(43)

6. End loop on i.

This algorithm is implemented in the MATLAB function program in Appendix C.2. The results of the conversion of the probe points for the Green phantom from Euclidean coordinates to θ , ϕ coordinates are shown in Fig. 15.

4.4 Volume Estimation by the Divergence Theorem

In this section we will state the Divergence Theorem and indicate how it can be used to estimate the volume of a polyhedron.

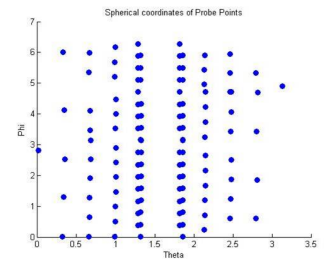


Figure 15: Plot of the θ , ϕ Coordinates of the Probe Points for the Coarse Data.

4.4.1 Divergence Theorem in 3-D and Volume Computation

Let R be a simply connected region in space and S the surface boundary. Then

$$\int \int_{R} \int \vec{\nabla} \cdot F \, dx dy dz = \int_{S} \int F \cdot \hat{n} \, d\sigma, \tag{44}$$

where $F(x, y, z) = (F_1(x, y, z), F_2(x, y, z), F_3(x, y, z))$ is a differentiable vector field,

$$\vec{\nabla} \cdot F = \frac{\partial F_1}{\partial x} + \frac{\partial F_2}{\partial y} + \frac{\partial F_3}{\partial z},\tag{45}$$

is the divergence of the vector field, \hat{n} is an outward-pointing unit normal vector to S, and $d\sigma$ is an infinitesimal element of the surface.

If we select F(x, y, z) = (1/3)(x, y, z) then $\vec{\nabla} \cdot F = 1$ and we can write

$$Vol(R) = \int \int_{R} \int dx dy dz = \frac{1}{3} \int_{S} \int \hat{n} \cdot (x, y, z) d\sigma.$$
 (46)

Now, if S is approximated by disjoint polyhedron (surface patches), then

$$S = \bigcup_{i=1}^{N} S_i, \tag{47}$$

and

$$\frac{1}{3} \int_{S} \int \hat{n} \cdot (x, y, z) d\sigma = \frac{1}{3} \sum_{i=1}^{N} \int_{S_{i}} \int \hat{n}_{i} \cdot (x, y, z) d\sigma, \tag{48}$$

where \hat{n}_i is the unit outer normal to S_i . Here we will model the surface patches by planar facets and, in particular, triangular facets. The plane S_i is given by $\hat{n}_i \cdot (x, y, z) = c_i$ for a constant c_i

associated with each triangular facet. Since this constant applies for all points on the triangular facet, it applies for any vertex point P_i of the triangular facet. The sum of the integrals over the facets reduces to

$$\frac{1}{3} \sum_{i=1}^{N} \int_{S_i} \int \hat{n_i} \cdot (x, y, z) d\sigma = \frac{1}{3} \sum_{i=1}^{N} \int_{S_i} \int c_i d\sigma = \frac{1}{3} \sum_{i=1}^{N} c_i \text{Area}(S_i).$$
 (49)

This method of computing the volume of an object from a surface integral can be found in Schneider and Eberly [15].

4.4.2 Area of a Planar Polyhedron in 3-D

We will describe the process of computing the area of a planar polyhedron in space by use of Stokes' Theorem and then we will particularize it to a planar triangle in space. Stokes' Theorem assumes that if C is a piecewise smooth boundary curve, oriented positively, of a surface S, and if F is a differentiable vector field defined on S and \hat{n} is a unit normal satisfying the right-hand rule relative to the boundary orientation, then

$$\int_{S} \int (\vec{\nabla} \times F) \cdot \hat{n} d\sigma = \int_{C} F \cdot d\vec{R}, \tag{50}$$

where the curl of $F(x, y, z) = (F_1(x, y, z), F_2(x, y, z), F_3(x, y, z))$ is

$$\vec{\nabla} \times F = \left(\frac{\partial F_3}{\partial y} - \frac{\partial F_2}{\partial z}, \frac{\partial F_1}{\partial z} - \frac{\partial F_3}{\partial x}, \frac{\partial F_2}{\partial x} - \frac{\partial F_1}{\partial y}\right),\tag{51}$$

and $d\vec{R} = (dx, dy, dz)$. If we set $F = (1/2)\hat{n} \times (x, y, z)$ then $\vec{\nabla} \times F = \hat{n}$ and $\hat{n} \cdot \hat{n} = 1$ so that

Area(S) =
$$\int_{S} \int d\sigma = \frac{1}{2} \oint_{C} (\hat{n} \times (x, y, z)) \cdot (dx, dy, dz),$$

= $\frac{1}{2} \oint_{C} \hat{n} \cdot ((x, y, z) \times (dx, dy, dz)),$
= $\frac{1}{2} \oint_{C} \hat{n} \cdot ((x(t), y(t), z(t)) \times (x'(t), y'(t), z'(t))) dt,$ (52)
= $\frac{1}{2} \oint_{C} \hat{n} \cdot (y(t)z'(t) - z(t)y'(t), z(t)x'(t) - x(t)z'(t), x(t)y'(t) - y(t)x'(t)) dt.$

We will now specialize the Stokes formula to find the area of a spatial triangle in terms of its three vertices oriented positively. Let the three vertices, in positive orientation, be specified by $v_1=(x_1,y_1,z_1),v_2=(x_2,y_2,z_2),v_3=(x_3,y_3,z_3)$. Parameterize the boundary of the triangle as follows. On [0,1] let $v=v_1+t(v_2-v_3)=(x_1+t(x_2-x_1),y_1+t(y_2-y_1),z_1+t(z_2-z_1))$. On [1,2] let $v=v_2+(t-1)(v_3-v_2)=(x_2+(t-1)(x_3-x_2),y_2+(t-1)(y_3-y_2),z_1+(t-1)(z_3-z_2))$. Finally on [2,3] let $v=v_3+(t-2)(v_1-v_3)=(x_3+(t-2)(x_1-x_3),y_3+(t-2)(y_1-y_3),z_3+(t-2)(z_1-z_3))$. The area of the spatial triangle, in terms of the parameterized boundary, can be written as

Area(S) =
$$\frac{1}{2} \oint_C \hat{n} \cdot (y(t)z'(t) - z(t)y'(t), z(t)x'(t) - x(t)z'(t), x(t)y'(t) - y(t)x'(t)) dt$$
,

$$= \frac{1}{2}\hat{n} \cdot \left(\int_0^3 (y(t)z'(t) - z(t)y'(t))dt, \int_0^3 (z(t)x'(t) - x(t)z'(t))dt, \right.$$

$$\left. \int_0^3 (x(t)y'(t) - y(t)x'(t))dt \right).$$
(53)

By straightforward integration over each of the parameterized segments it is easy to show that

Area(S) =
$$\frac{1}{2}\hat{n} \cdot ((y_1z_2 - y_2z_1) + (y_2z_3 - y_3z_2) + (y_3z_1 - y_1z_3),$$

$$(z_1x_2 - z_2x_1) + (z_2x_3 - z_3x_2) + (z_3x_1 - z_2x_3),$$

$$(x_1y_2 - x_2y_1) + x_2y_3 - x_3y_2) + (x_3y_1 - x_1y_3))$$
(54)

The normal vector to the oriented triangle can be computed as follows. Let $\vec{v}=(x_2-x_1,y_2-y_1,z_2-z_1)$ and $\vec{w}=(x_3-x_1,y_3-y_1,z_3-z_1)$. Then $\vec{n}=\vec{v}\times\vec{w}=((y_2-y_1)(z_3-z_1)-(z_2-z_1)(y_3-y_1),(z_2-z_1)(x_3-x_1)-(x_2-x_1)(z_3-z_1),(x_2-x_1)(y_3-y_1)-(y_2-y_1)(x_3-x_1))$. If we set $n_1=(y_2-y_1)(z_3-z_1)-(z_2-z_1)(y_3-y_1),\; n_2=(z_2-z_1)(x_3-x_1)-(x_2-x_1)(z_3-z_1),n_3=(x_2-x_1)(y_3-y_1)-(y_2-y_1)(x_3-x_1)$ then $\|\vec{n}\|=\sqrt{n_1^2+n_2^2+n_3^2}$ and $\hat{n}=\vec{n}/\|\vec{n}\|$.

4.5 Volume Estimation for the Phantom Grid Data

We divided the phantom surfaces into triangular surface patches as follows. First of all we assumed that the phantom surfaces had been partitioned into nh+1 latitude grid points and nv+1 longitude grid points, where the latitude grid points were set as $\phi_1 = 0 < \phi_2 < \cdots < \phi_{nh} < \phi_{nh+1} = 2\pi$ and longitude grid points given by $\theta_1 = 0 < \theta_2 < \cdots < \theta_{nv} < \theta_{nv+1} = \pi$. At every grid point there was a radius, $r(\phi, \theta)$, calculated from the surface B-spline model. At the north and south poles the radii was taken as the median value of the grid point radial values for the cases $\theta = 0$ and $\theta = \pi$.

The patches at the north and south poles were automatically triangular patches. In fact, at the north pole, where we started, there were nh triangular patches. The Euclidean points at the triangle corners for each patch were computed by

$$x = r \sin(\theta) \cos(\phi) \quad 0 \le \theta \le \pi,$$

$$y = r \sin(\theta) \sin(\phi) \quad 0 \le \phi \le 2\pi,$$

$$z = r \cos(\theta).$$
(55)

For all of the triangle facets at the north pole we designated the pole as $(x_1, y_1, z_1) = (0, 0, r_m)$, where r_m was the median value of the grid generated data for $\theta_1 = 0$. We looped through the grid points at θ_2 and $\phi_1, \phi_2, \dots, \phi_{nh+1} = \phi_1$. At each of these angle pairs there was a value $r_k = r(\phi_k, \theta_2)$, $k = 1, 2, \dots, nh+1$. We generated the volume from the Green's Theorem result by adding up the contributions of each facet to the volume total. We did this by initializing a variable for the volume, called vol. We then started to generate the contributions from the first layer of facets at the north pole. We set, as noted above, the north pole as (x_1, y_1, z_1) and then set

$$x_{2} = r(\phi_{1}, \theta_{2}) \sin(\theta_{2}) \cos(\phi_{1}),$$

$$y_{2} = r(\phi_{1}, \theta_{2}) \sin(\theta_{2}) \sin(\phi_{1}),$$

$$z_{2} = r(\phi_{1}, \theta_{2}) \cos(\theta_{2}),$$

and

$$x_3 = r(\phi_2, \theta_2) \sin(\theta_2) \cos(\phi_2),$$

$$y_3 = r(\phi_2, \theta_2) \sin(\theta_2) \sin(\phi_2),$$

$$z_3 = r(\phi_2, \theta_2) \cos(\theta_2).$$

These were set in a positive orientation. Next we computed the outward normal to the triangle facet, formed the vectors $v_1 = (x_2, y_2, z_2) - (x_1, y_1, z_1)$, $v_2 = (x_3, y_3, z_3) - (x_1, y_1, z_1)$, and then formed the normalized cross product

$$\hat{n} = \frac{v_1 \times v_2}{\|v_1 \times v_2\|} \tag{56}$$

We then computed the contribution that this facet made to the volume as

$$(\hat{n} \cdot (x_1, y_1, z_1)) \left\{ \hat{n} \cdot \left(\sum_{j=1}^{2} (y_j z_{j+1} - y_{j+1} z_j), (z_j x_{j+1} - z_{j+1} x_j), (x_j y_{j+1} - x_{j+1} y_j) \right) \right\}.$$

We proceeded to the next facet in the first layer. Again (x_1, y_1, z_1) was the north pole and we then used the previous computation to get $x_2 = x_3$, $y_2 = y_3$, $z_2 = z_3$ and set

$$x_3 = r(\phi_3, \theta_2)\sin(\theta_2)\cos(\phi_3),$$

$$y_3 = r(\phi_3, \theta_2)\sin(\theta_2)\sin(\phi_3),$$

$$z_3 = r(\phi_3, \theta_2)\cos(\theta_2).$$

We computed the normalized cross product as for the first facet and then computed the contribution of the second facet to the volume using equation (57). We continued this process for θ_2 , ϕ_j , $j = 1, \dots, nh$.

We next computed the contributions of the next layers in a two step process. We looped through the θ layers from 2 to nv-1. The last layer at the south pole was handled separately. At each layer there were nh four-vertex patches. Each patch was divided into two triangles. The four vertices of the patches were identified counterclockwise as $(\theta_i, \phi_j, r(i, j)), (\theta_{i+1}, \phi_j, r(i+1, j)), (\theta_{i+1}, \phi_{j+1}, r(i+1, j+1)), (\theta_i, \phi_{j+1}, r(i, j+1))$. For the first triangle in the patch we set

$$x_1 = r(i,j)\sin(\theta_i)\cos(\phi_j),$$

$$y_1 = r(i,j)\sin(\theta_i)\sin(\phi_j),$$

$$z_1 = r(i,j)\cos(\theta_i).$$

$$x_2 = r(i+1,j)\sin(\theta_{i+1})\cos(\phi_j),$$

$$y_2 = r(i+1,j)\sin(\theta_{i+1})\sin(\phi_j),$$

$$z_2 = r(i+1,j)\cos(\theta_{i+1}).$$

$$x_3 = r(i+1,j+1)\sin(\theta_{i+1})\cos(\phi_{j+1}),$$

$$y_3 = r(i+1,j+1)\sin(\theta_{i+1})\sin(\phi_{j+1}),$$

$$z_3 = r(i+1,j+1)\cos(\theta_{i+1}).$$

We then proceeded as above to compute the contribution of this triangle to the volume. For the second triangle of the patch we maintained the same x_1 , y_1 , z_1 and set $x_2 = x_3$, $y_2 = y_3$, $z_2 = z_3$ and then set

$$x_3 = r(i, j+1) \sin(\theta_i) \cos(\phi_{j+1}),$$

$$y_3 = r(i, j+1) \sin(\theta_i) \sin(\phi_{j+1}),$$

$$z_3 = r(i, j+1) \cos(\theta_i).$$

Again, the volume increment due to this triangle was computed as it was above.

Finally, at the south pole there were nh triangles to compute. Their vertices were chosen as follows.

```
x_{1} = r(nv, j) \sin(\theta_{n}v) \cos(\phi_{j}),
y_{1} = r(nv, j) \sin(\theta_{n}v) \sin(\phi_{j}),
z_{1} = r(nv, j) \cos(\theta_{n}v).
x_{2} = 0,
y_{2} = 0,
z_{2} = -r(nv + 1, 1).
x_{3} = r(nv, j + 1) \sin(\theta_{nv}) \cos(\phi_{j+1}),
y_{3} = r(nv, j + 1) \sin(\theta_{nv}) \sin(\phi_{j+1}),
z_{3} = r(nv, j + 1) \cos(\theta_{nv}).
```

The contribution of these triangles to the volume was computed as above. After all of the triangle contributions to the volume were computed the final volume was then computed as vol = (1/6)vol.

4.6 Computational Results for the B-spline Model

In this section we will describe the method used to estimate volume uncertainties and discuss the results of using a B-spline surface model and the Divergence Theorem to estimate the phantom volumes. A nonparametric method to estimate the volume uncertainties was chosen since there did not exist any "ground truth" values for the Green and Pink FDA phantom volumes.

4.6.1 Estimating Volume Uncertainties

The nonparametric "bootstrap" method is a computer intensive technique for estimating uncertainties. It involves repeated Monte Carlo resampling from the original spherical coordinates with radii values modified by the fitting residuals, refitting the model to estimate new volumes, and finally computing an uncertainty for the process from the set of computed volumes. For a full discussion of the bootstrap see Efron and Tibshirani [9]. We will give a brief description of the bootstrp method in Section 4.6.2 in the discussion of the coarse data results, but will use the same method for estimating the volume uncertainties in the case of the dense data.

4.6.2 Volume Estimates and Uncertainties for the Coarse Data

In this section we state the results of computing the volumes for the two phantoms using the six coarse data sets. Sixteen surface grid cases were used and the results are shown in Tables 8 through 13. Figures 16 through 26 show the dramatic asymptotic trend of the volume estimates as the grid sizes increase. It seems clear that the volume of the Green phantom may lie somewhere in the range of $4337 \ mm^3$ and $4340 \ mm^3$ while the volume of the Pink phantom volume seems to lie somewhere in the range $3910 \ mm^3$ and $3913 \ mm^3$. In terms of percentages these volume differences are in the range of about 0.07%. Figures 16 to 20 show the clear increase of the volume estimate to a stable value as the surface grid became finer. The volume estimates for the Green phantom differ from the volume estimates for the sphere fit in Table 3 by approximately 0.1% to 0.2%. This is indicative

of the fact that the Green phantom is near spherical. The fit to the Pink phantom in Tables 11 to 13 are all consistent and indicate the Pink phantom likely has a larger volume than that indicated by the sphere fit in Table 4. The differences of the stabilized Pink volume estimates differ from the spherical fit volume estimates by approximately 2%. Again this likely indicates the non-spherical nature of the Pink phantom.

The object of the uncertainty estimation procedure (bootstrap) was to develop volume uncertainty as a function of grid size. The process began with the conversion of the Euclidean data points to spherical coordinates. An automatic selection of knots was done. These steps were done once for a given data set. It was assumed that the modifications of the data made during the bootstrap would be small and not affect the knot selection. During the first pass of the bootstrap algorithm the radii of the spherical coordinates were fit by a tensor product of B-splines. The predicted values and residuals from this initial fit were called the master predicted values and master residuals respectively. The computed volume was then put in a list of volumes that would be added to in subsequent passes of the algorithm. The algorithm then iterated two hundred times as follows. In the first iteration the master residuals were sampled randomly uniformly with replacement. The resampled residuals were then added to the master predicted radii and a new fit was performed. The new computed volume was added to the volume list and the master residuals sampled randomly uniformly with replacement for the next iteration. The process continued for all two hundred iterations. The standard deviation of the volumes in the volume list was then used as an estimate of the volume uncertainty and the average volume was taken as the reference volume for the chosen grid size.

From Tables 8 to 10 the uncertainties range from $7.4~mm^3$ on the coarse surface grid to $3.3~mm^3$ on the finer grid for the Green phantom. These uncertainty values represent an approximate range of 0.2% to 0.07% of the estimated Green phantom volumes from the tables. Figures 17 to 21 graphically show the rapid descent to stable values for the uncertainties. Tables 11 to 13 show uncertainties for the Pink phantom that are about twice those for the Green phantom. This is not surprising, considering the non-spherical shape of the Pink phantom. The uncertainties for the Pink phantom have an approximate range of 0.5% to 0.2% of the estimated volumes. Figures 23 to 27 show a similar descent of the uncertainties as a function of grid refinement. In a similar manner to the volumetric behavior of the Green phantom as the grid refinement increases Figures 22 to 26 show the rapid increase to a stable value for the Pink phantom volume estimates.

Green 4 Data Set				
Volumes and Expanded Uncertainties vs Grid Size				
Grid Case	θ	ϕ	Volume (mm^3)	Uncertainty (mm^3)
1	10	20	4071.5	7.1
2	20	40	4278.6	5.1
3	30	60	4313.6	4.0
4	40	80	4325.5	4.0
5	50	100	4331.2	3.7
6	60	120	4334.4	3.6
7	70	140	4336.1	3.6
8	80	160	4337.5	3.7
9	90	180	4337.9	3.7
10	100	200	4338.8	3.7
11	110	220	4339.1	3.2
12	120	240	4339.4	3.7
13	130	260	4339.6	3.8
14	140	280	4339.9	3.9
15	150	300	4339.9	3.5
16	160	320	4340.0	3.3

Table 8: Estimated Volumes and Uncertainties for Green 4 Data

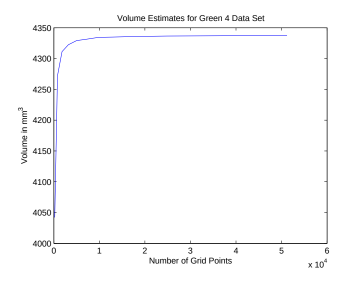


Figure 16: Volume Estimates for Green 4 Data Set vs. Grid Size.

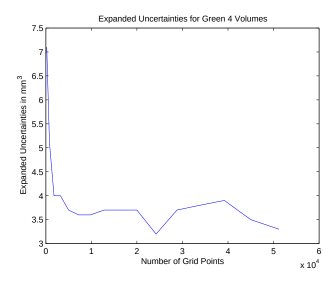


Figure 17: Uncertainty Estimates for Green 4 Data Set vs. Grid Size.

	Green 5 Data Set						
Volur	Volumes and Expanded Uncertainties vs Grid Size						
Grid Case	θ	ϕ	Volume (mm^3)	Uncertainty (mm^3)			
1	10	20	4068.5	7.4			
2	20	40	4275.4	4.8			
3	30	60	4311.4	4.2			
4	40	80	4323.1	3.8			
5	50	100	4328.5	3.7			
6	60	120	4331.1	3.7			
7	70	140	4333.2	3.5			
8	80	160	4334.5	3.6			
9	90	180	4335.1	3.4			
10	100	200	4335.6	3.4			
11	110	220	4336.1	3.6			
12	120	240	4336.5	3.5			
13	130	260	4336.8	3.1			
14	140	280	4336.9	3.3			
15	150	300	4337.0	3.4			
16	160	320	4337.1	3.5			

Table 9: Estimated Volumes and Uncertainties for Green 5 Data

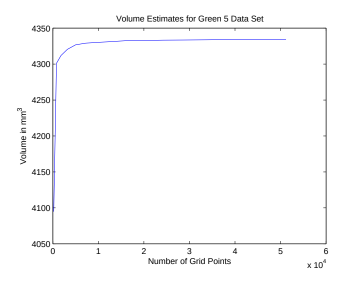


Figure 18: Volume Estimates for Green 5 Data Set vs. Grid Size.

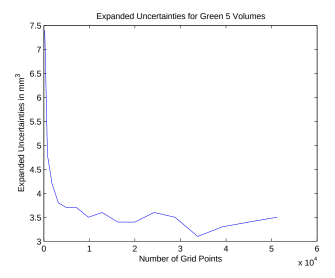


Figure 19: Uncertainty Estimates for Green 5 Data Set vs. Grid Size.

	Green 6 Data Set						
Volui	Volumes and Expanded Uncertainties vs Grid Size						
Grid Case	θ	ϕ	Volume (mm^3)	Uncertainty (mm^3)			
1	10	20	4071.3	7.4			
2	20	40	4277.6	4.8			
3	30	60	4312.9	3.8			
4	40	80	4325.0	4.1			
5	50	100	4330.6	3.4			
6	60	120	4333.7	3.3			
7	70	140	4335.6	3.4			
8	80	160	4336.9	3.4			
9	90	180	4337.4	3.9			
10	100	200	4338.0	3.4			
11	110	220	4338.5	3.8			
12	120	240	4338.9	3.6			
13	130	260	4339.1	3.4			
14	140	280	4339.6	3.5			
15	150	300	4339.5	3.4			
16	160	320	4339.8	3.4			

Table 10: Estimated Volumes and Uncertainties for Green 6 Data

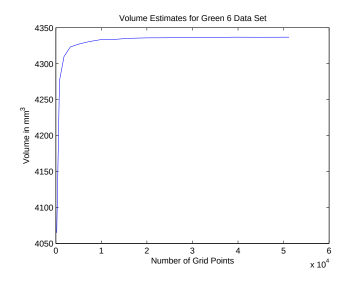


Figure 20: Volume Estimates for Green 6 Data Set vs. Grid Size.

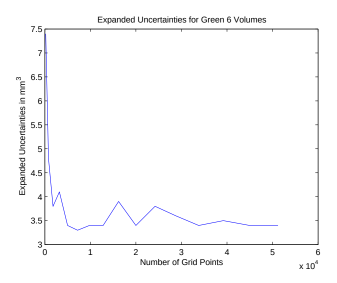


Figure 21: Uncertainty Estimates for Green 6 Data Set vs. Grid Size.

	Pink 5 Data Set						
Volu	Volumes and Expanded Uncertainties vs Grid Size						
Grid Case	θ	ϕ	Volume (mm^3)	Uncertainty (mm^3)			
1	10	20	3697.8	18.7			
2	20	40	3866.1	9.8			
3	30	60	3899.2	8.7			
4	40	80	3908.8	7.4			
5	50	100	3914.9	7.4			
6	60	120	3917.1	6.8			
7	70	140	3919.4	6.9			
8	80	160	3920.7	6.3			
9	90	180	3921.2	6.8			
10	100	200	3921.8	6.2			
11	110	220	3922.3	6.7			
12	120	240	3922.6	7.0			
13	130	260	3922.8	6.7			
14	140	280	3922.9	6.9			
15	150	300	3922.9	5.8			
16	160	320	3923.2	6.3			

Table 11: Estimated Volumes and Uncertainties for Pink 5 Data

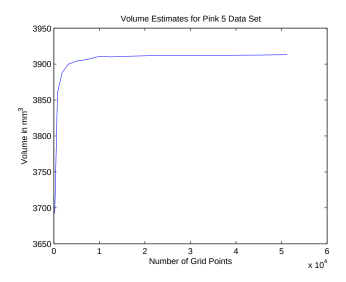


Figure 22: Volume Estimates for Pink 5 Data Set vs. Grid Size.

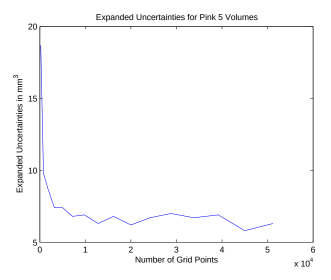


Figure 23: Uncertainty Estimates for Pink 5 Data Set vs. Grid Size.

Pink 6 Data Set							
Volur	Volumes and Expanded Uncertainties vs Grid Size						
Grid Case	θ	ϕ	Volume (mm^3)	Uncertainty (mm^3)			
1	10	20	3698.3	17.8			
2	20	40	3864.3	11.2			
3	30	60	3898.2	8.0			
4	40	80	3907.5	7.7			
5	50	100	3913.5	7.2			
6	60	120	3916.2	7.0			
7	70	140	3917.6	6.5			
8	80	160	3919.3	6.3			
9	90	180	3919.8	6.6			
10	100	200	3920.8	6.8			
11	110	220	3921.0	6.3			
12	120	240	3921.6	6.4			
13	130	260	3921.2	6.3			
14	140	280	3922.0	5.9			
15	150	300	3921.4	5.8			
16	160	320	3922.1	6.6			

Table 12: Estimated Volumes and Uncertainties for Pink 6 Data

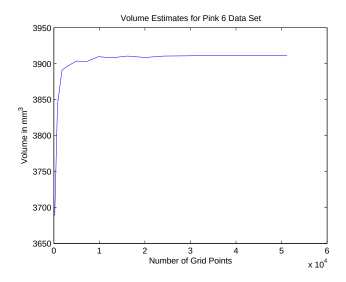


Figure 24: Volume Estimates for Pink 6 Data Set vs. Grid Size.

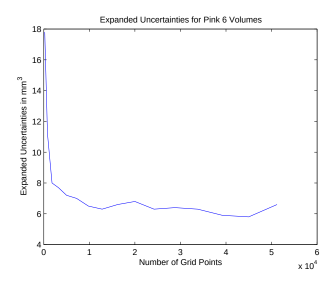


Figure 25: Uncertainty Estimates for Pink 6 Data Set vs. Grid Size.

	Pink 7 Data Set						
Volui	Volumes and Expanded Uncertainties vs Grid Size						
Grid Case	θ	ϕ	Volume (mm^3)	Uncertainty (mm^3)			
1	10	20	3697.2	19.2			
2	20	40	3865.3	10.8			
3	30	60	3897.5	9.0			
4	40	80	3907.6	7.6			
5	50	100	3913.6	6.8			
6	60	120	3915.9	6.6			
7	70	140	3917.3	6.2			
8	80	160	3919.2	6.5			
9	90	180	3919.9	6.2			
11	110	220	3920.8	6.4			
10	100	200	3920.7	6.6			
12	120	240	3921.1	6.4			
13	130	260	3921.2	6.6			
14	140	280	3921.7	6.4			
15	150	300	3921.7	6.3			
16	160	320	3921.9	6.8			

Table 13: Estimated Volumes and Uncertainties for Pink 6 Data

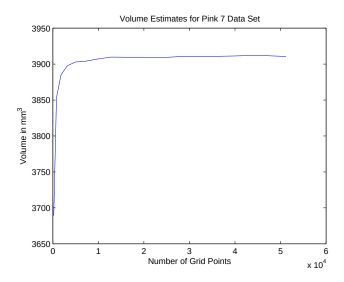


Figure 26: Volume Estimates for Pink 7 Data Set vs. Grid Size.

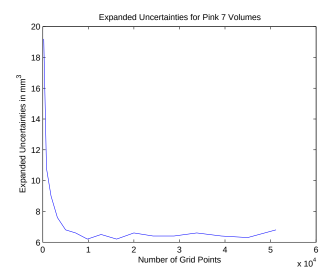


Figure 27: Uncertainty Estimates for Pink 7 Data Set vs. Grid Size.

	Green 1 Dense Data Set						
Volui	Volumes and Expanded Uncertainties vs Grid Size						
Grid Case	θ	ϕ	Volume (mm^3)	Uncertainty (mm^3)			
1	10	20	3816.4	210.6			
2	20	40	4014.0	135.8			
3	30	60	4052.5	111.6			
4	40	80	4065.4	105.0			
5	50	100	4067.5	93.6			
6	60	120	4070.6	94.0			
7	70	140	4071.3	91.4			
8	80	160	4073.0	91.4			
9	90	180	4073.7	91.4			
10	100	200	4073.5	89.6			
11	110	220	4074.8	89.8			
12	120	240	4074.5	88.4			
13	130	260	4075.4	89.2			
14	140	280	4075.0	87.2			
15	150	300	4075.3	88.0			
16	160	320	4075.2	87.4			

Table 14: Estimated Volumes and Uncertainties for Green 1 Dense Data

4.6.3 Volume Estimates and Uncertainties for the Dense Data

In this section we report the results of volume estimates and the extended uncertainties for the dense data sets of 181 probed points on the phantom surfaces. The results do not seem to be what one would have expected. The volumes and the uncertainties have not been plotted since in general they follow the pattern established by the volumes and uncertainties in the coarse data. We will just consider the differences. The first thing of interest is that the stabilized volume estimates for the Green dense data differ from the volume estimates for the Green coarse data by about 6%. This is in the negative direction in that the volume estimates for the Green dense data were smaller than for the Green coarse data. In the case of the volume estimates for the Pink dense data, the differences with the volume estimates for the Pink coarse data are approximately 4% larger. These estimates are clear from Tables 14 to 19. Another thing to notice is that the uncertainties are significantly higher as shown in Tables 14 through 19 compared to those estimated from the coarse data. In fact they appear to be an order of magnitude larger. It is not clear why the results for the dense data differ from the results for the coarse data, but we are reporting it nonetheless. The complete analysis of these differences is beyond the scope of this report but will be pursued in a coming study related to a redesign of the fitting algorithm.

5 Summary

The B-spline surface model joined with the Divergence Theorem seems to be a viable approach for estimating volumes of near spherical star-shaped artifacts, but the results seem to depend strongly on the distribution of the data points on the surface. In the case of sparse data the problem of extrapolating a surface fitted to the data to a grid on the surface leads potentially to unwanted

C 0 D D-+- C-+							
Green 2 Dense Data Set							
Volur	Volumes and Expanded Uncertainties vs Grid Size						
Grid Case	θ	ϕ	Volume (mm^3)	Uncertainty (mm^3)			
1	10	20	3788.2	225.6			
2	20	40	3989.5	137.4			
3	30	60	4027.8	123.4			
4	40	80	4034.4	112.2			
5	50	100	4040.2	103.6			
6	60	120	4041.2	99.8			
7	70	140	4042.7	102.2			
8	80	160	4043.7	100.4			
9	90	180	4045.4	98.4			
10	100	200	4044.9	98.6			
11	110	220	4044.7	97.2			
12	120	240	4044.9	96.8			
13	130	260	4045.4	96.8			
14	140	280	4045.3	95.8			
15	150	300	4045.7	95.6			
16	160	320	4046.3	95.2			

Table 15: Estimated Volumes and Uncertainties for Green 2 Dense Data

	Green 3 Dense Data Set						
Volui	Volumes and Expanded Uncertainties vs Grid Size						
Grid Case	θ	ϕ	Volume (mm^3)	Uncertainty (mm^3)			
1	10	20	3859.8	225.0			
2	20	40	4064.0	138.0			
3	30	60	4097.9	130.4			
4	40	80	4102.7	115.4			
5	50	100	4110.2	105.6			
6	60	120	4113.2	105.6			
7	70	140	4114.8	105.4			
8	80	160	4116.3	100.2			
9	90	180	4119.4	100.6			
10	100	200	4119.1	101.6			
11	110	220	4118.6	99.4			
12	120	240	4118.6	99.8			
13	130	260	4119.7	99.6			
14	140	280	4118.7	99.2			
15	150	300	4120.0	98.0			
16	160	320	4119.3	99.6			

Table 16: Estimated Volumes and Uncertainties for Green 3 Dense Data

Pink 1 Dense Data Set							
Volur	Volumes and Expanded Uncertainties vs Grid Size						
Grid Case	θ	ϕ	Volume (mm^3)	Uncertainty (mm^3)			
1	10	20	3663.7	193.0			
2	20	40	3847.9	105.6			
3	30	60	3878.9	88.0			
4	40	80	3891.9	83.8			
5	50	100	3896.0	75.2			
6	60	120	3898.5	73.2			
7	70	140	3902.3	73.2			
8	80	160	3903.0	72.0			
9	90	180	3903.2	72.0			
10	100	200	3903.8	71.8			
11	110	220	3905.0	71.0			
12	120	240	3905.0	70.8			
13	130	260	3905.2	69.2			
14	140	280	3904.7	69.6			
15	150	300	3905.5	69.2			
16	160	320	3905.3	69.0			

Table 17: Estimated Volumes and Uncertainties for Pink 1 Dense Data

	Pink 2 Dense Data Set						
Volui	Volumes and Expanded Uncertainties vs Grid Size						
Grid Case	θ	ϕ	Volume (mm^3)	Uncertainty (mm^3)			
1	10	20	3690.2	191.8			
2	20	40	3881.2	110.6			
3	30	60	3916.1	91.4			
4	40	80	3923.5	83.6			
5	50	100	3926.6	73.6			
6	60	120	3932.6	74.4			
7	70	140	3934.1	74.8			
8	80	160	3934.8	73.6			
9	90	180	3935.8	71.8			
10	100	200	3934.2	71.2			
11	110	220	3935.6	71.0			
12	120	240	3935.8	70.4			
13	130	260	3936.3	70.0			
14	140	280	3936.1	69.0			
15	150	300	3936.1	70.4			
16	160	320	3936.1	70.0			

Table 18: Estimated Volumes and Uncertainties for Pink 2 Dense Data

Pink 3 Dense Data Set							
Volur	Volumes and Expanded Uncertainties vs Grid Size						
Grid Case	θ	ϕ	Volume (mm^3)	Uncertainty (mm^3)			
1	10	20	3274.6	236.8			
2	20	40	3462.9	153.6			
3	30	60	3488.5	130.8			
4	40	80	3496.4	118.6			
5	50	100	3498.3	110.8			
6	60	120	3500.4	110.4			
7	70	140	3502.8	108.0			
8	80	160	3503.2	106.8			
9	90	180	3505.9	105.2			
10	100	200	3504.5	105.4			
11	110	220	3506.2	104.0			
12	120	240	3506.2	103.0			
13	130	260	3507.5	103.2			
14	140	280	3508.2	104.2			
15	150	300	3508.5	103.4			
16	160	320	3507.5	104.0			

Table 19: Estimated Volumes and Uncertainties for Pink 3 Dense Data

oscillations in the neighborhood of some of the sparse data points. In the current algorithm this problem was dealt with for convenience by a chopping operation, but this may not be a good procedure and may have affected the differences between the coarse data results and the dense data results, although this needs to be investigated further.

The results obtained from the coarse data distribution appear to be consistent with the results from the spherical model fits. As shown in Tables 8, 9, and 10 in Section 4.6.2 the volume estimates for the Green sphere are very close to the estimates obtained by the B-spline model suggesting that the Green sphere is essentially spherical in shape. As expected the volume estimates in Table 4 for the Pink sphere are significantly lower than the B-spline model estimates in Tables 11, 12, and 11, in this case suggesting that the Pink sphere has a significant non-spherical shape.

The results from the dense data distribution were not what would have been expected. The uncertainties were significantly higher than those for the coarse data. Tables 14 to 16 show volumes smaller than those from 8, 9, and 10. Tables 17 and 18 show similar volume estimates but Table 19 shows significantly lower volume estimates. This, of course, could have been brought about by any difficulty in holding the Pink phantom in position 3.

In summary we conclude that the combination of using the CMM to obtain surface data and the B-spline/Divergence Theorem volume estimation method can produce useful results but that there are some limitations. First, the CMM is primarily used for probing manufactured metalic artifacts and its use in probing non-metalic artifacts such as the FDA phantoms can lead to some large uncertainties in volume estimation. Second, the distribution of probed points can lead to unexpected results. So there is a problem of consistency here. Third, the methods used may not provide useful volume estimation for more complex artifacts that surely would arise in developing simulated lung cancers. The current artifacts were near-spherical and even these did not provide fully expected results based on data distribution. As a result of these limitations we think other means of estimating "ground truth" volumes for the current and any future phantoms be investigated

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7 Disclaimer

Certain commercial software products are identified in this paper in order to adequately specify the computational procedures. Such identification does not imply recommendation or endoresement by the National Institute of Standards and Technology nor does it imply that the software products identified are necessarily the best available for the purpose.

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A Fitting a Sphere to Probe Data

```
function sphere_fit_to_phantoms_simplex
%This functions uses a nonlinear optimization procedure to fit a set of
\%(x,y,z) data points determined by a CMM on the surface of two phantom lung
%cancer lesions supplied to NIST by the FDA.
global x y z
format long;
%Open the CMM data file
%select the file desired by removing the % sign
%fid= fopen('Tumor_Green_and_Pink_Results_Green_4.txt','r');
%fid= fopen('Tumor_Green_and_Pink_Results_Green_5.txt','r');
%fid= fopen('Tumor_Green_and_Pink_Results_Green_6.txt','r');
%fid= fopen('Tumor_Green_and_Pink_Results_Pink_5.txt','r');
%fid= fopen('Tumor_Green_and_Pink_Results_Pink_6.txt','r');
%fid= fopen('Tumor_Green_and_Pink_Results_Pink_7.txt','r');
%fid= fopen('Green_Results_dense_data_1.txt','r');
%fid= fopen('Green_Results_dense_data_2.txt','r');
%fid= fopen('Green_Results_dense_data_3.txt','r');
%fid= fopen('Pink_Results_dense_data_1.txt','r');
%fid= fopen('Pink_Results_dense_data_2.txt','r');
fid= fopen('Pink_Results_dense_data_3.txt','r');
%fid= fopen('Cal_121_sphere.txt','r');
%fid= fopen('Cal_181_sphere.txt','r');
%reads file into a 3-by-inf matrix and takes the transpose. This is
%required due to the way Matlab reads an array.
```

```
A=fscanf(fid, '%f %f %f', [3 inf])';
[n,m] = size(A);
x=A(:,1);
y=A(:,2);
z=A(:,3);
fclose(fid);
mnx = mean(x);
mny = mean(y);
mnz = mean(z);
%Set initial center and radius
c = [mnx mny mnz 10];
[c_final,val_final,iter]=neldermead_mod(@f2,c,1,1.0e-14,1000)
%Choose the median values of the tetrahedron vertex coordinates
c_final_median = median(c_final')'
Estimated_volume = (4/3)*pi*(c_final_median(4))^3
%Choose the median of the sum of squares of the function values at the
%vertices.
val_final_median = median(val_final')'
%Compute the residual radial value between the data points to the computed
%center and the estimated sphere radius.
%First get the radial values to the data points
radius = sqrt((x-c_final_median(1)).^2 + (y-c_final_median(2)).^2 + (z-c_final_median(3)).^2);
Residual_radius = radius - c_final_median(4);
for i = 1:length(x)
         disp(sprintf('%f
                                                     %f
                                                                   %f
                                                                                %f
                                                                                             %f ',x(i),y(i),z(i), radius(i), Residual_radius(i)));
end
Mean_residual_radius = mean(Residual_radius)
Mean_abs_residual_radius = mean(abs(Residual_radius))
Std_residual_radius = std(Residual_radius)
Std_abs_residual_radius = std(abs(Residual_radius))
%Estimating coefficient uncertainties
Z_{\text{hat}} = \begin{bmatrix} -2*(x(:)-c_{\text{final_median}}(1)) & -2*(y(:)-c_{\text{final_median}}(2)) & -2*(z(:)-c_{\text{final_median}}(3)) & -2*(z(:)-c_{\text{final_median}}(2)) &
s = f2(c_final_median)/(length(x) - 4);
SE = inv(Z_hat'*Z_hat)*s^2;
for i = 1:4
         se_c(i) = SE(i,i);
         expand_uncert_c(i) = 2*se_c(i);
end
se c
expand_uncert_c
%This function computes the sum of ((x(i) - c(1))^2 + (y(i) - c(2))^2
%+ (z(i) - c(3))^2 - c(4)^2)^2  for i = 1:n
function g = f2(c)
```

```
%F2 - Forms the sum of squares of the sphere model equations to be solved
%Input:
% c - Parameters defining a sphere. (c(1),c(2),c(3)) the center, c(4) the
     radius
global x y z
n = length(x);
g = zeros(n,1);
for i = 1:n
  p = x(i) - c(1);
   q = y(i) - c(2);
  r = z(i) - c(3);
   p2 = p^2;
  q2 = q^2;
  r2 = r^2;
   s2 = c(4)^2;
   pqrs = p2 + q2 + r2 - s2;
   g(i,1) = pqrs^2;
end
g = sum(g);
%Nelder-Mead search algorithm for minimizing a nonlinear function without
%derivatives.
function [x,y,iter]=neldermead_mod(f,xbar,rad,tol,maxit)
% Modified Nelder-Mead Search (modified with comments and convergence test)
% Input: function handle f, best guess xbar (column vector),
%
   initial search radius rad , tolerance on iterative polyhedron volume,
   and maximum iterations allowed.
% Output: matrix x whose columns are vertices of simplex,
%
   function values y of those vertices, and iterations used.
%Author: T. Sauer, George Mason University
%Modified: D. Gilsinn, National Institute of Standards and Technology
n=length(xbar);
x(:,1)=xbar;
               % each column of x is a simplex vertex
x(:,2:n+1)=xbar*ones(1,n)+rad*eye(n,n);
%determinant related to the volume of a polyhedron
v = zeros(n+1,n+1);
v(1,:) = 1;
```

```
for j = 1:n+1
    v(2:n+1,j) = x(:,j);
d = det(v);
for j=1:n+1
 y(j)=f(x(:,j));
                   % evaluate obj function f at each vertex
%The output array r is the ascending order of the values in the original y
[y,r]=sort(y);
                      % sort the function values in ascending order
x=x(:,r);
                           and rank the vertices the same way
iter = 1;
while ((abs(d) > tol)& (iter <= maxit))
    iter = iter + 1;
    %The mean function of a matrix produces a row vector of the column
    %means. Since we want row means across vertices we have to
    %transpose the matrix of vertices. We then transpose the row of
    %means to form a column vector.
  xbar=mean(x(:,1:n))'; % xbar is the centroid of the face
  xh=x(:,n+1);
                          % omitting the worst vertex xh
  %First step in each iteration is to reflect through the centroid
  %of the opposite face to the worst point and compute the function
  %value at the reflected point
  xr = 2*xbar - xh; yr = f(xr);
  %There are two major cases to test: yr < y(n), y(n) \le yr
  if yr < y(n)
     %If yr is better than the current best value then this inicates
     %that we might get better values by continuing in the same direction
     %by an expansion
    if yr < y(1)
                      % try expansion xe, i.e. xe = xbar + 2*(xbar-xh)
     xe = 3*xbar - 2*xh; ye = f(xe);
     %If the expansion leads to a better value accept the point
     %go to the end, resort the points and continue to the next iteration
                     % accept expansion
      if ve < vr
        x(:,n+1) = xe; y(n+1) = f(xe); %resort and go to the next iteration
                     % accept reflection point as the next best if yr<=ye
       x(:,n+1) = xr; y(n+1) = f(xr);
      end
                      % xr is middle of pack, accept reflection
    else
     x(:,n+1) = xr; y(n+1) = f(xr);
    end
  else
                      %Now test y(n) <= yr, use contractions
    if yr < y(n+1)
                     % try outside contraction xoc
     xoc = 1.5*xbar - 0.5*xh; yoc = f(xoc);
                     % accept outside contraction if the new point is better
      if yoc < yr
       x(:,n+1) = xoc; y(n+1) = f(xoc);
      else
                      % otherwise shrink polytope toward best point
       for j=2:n+1
          x(:,j) = 0.5*x(:,1)+0.5*x(:,j); y(j) = f(x(:,j));
```

```
end
      end
    else
                      %here y(n+1) <= yr and xr is even worse than the previous worst</pre>
     xic = 0.5*xbar+0.5*xh; yic = f(xic); %contract towards the previous worst point
      if yic < y(n+1) %if things are better accept inside contraction
        x(:,n+1) = xic; y(n+1) = f(xic);
                      % if y(n+1) \le yic then shrink polytope toward best point
      else
        for j=2:n+1
          x(:,j) = 0.5*x(:,1)+0.5*x(:,j); y(j) = f(x(:,j));
      end
    end
 end
  [y,r] = sort(y);
                        % resort the new obj function values
 x=x(:,r);
                             and rank the vertices the same way
 %determinant related to the volume of n-hedron
 for j = 1:n+1
      v(2:n+1,j) = x(:,j);
 end
 d = det(v);
end
                        %go back for the next iteration.
```

B Main Volume Estimation Code

```
function Main_Phantom_Volume
%MAIN_PHANTOM_VOLUME - This function is an interactive function that reads
%
                   a data file of Cartesian coordinates, obtained from
%
                   a CMM, converts them to spherical coordinates,
%
                   adaptively selects a near optimum set of knots to
%
                   fit a tensor product of B-splines to the data,
%
                   generates a grid of points in theta and phi and
%
                   extrapolates the B-spline function values to the
%
                   grid and finally computes the volume of a phantom
%
                   for that grid. A statistical bootstrap method is
%
                   used to estimate volumetric uncertainties.
%
%Author:
   David E. Gilsinn
   Mathematical and Computational Sciences Division
   Information Technology Laboratory
   National Institute of Standards and Technology
   Gaithersburg, MD 20899-8910
```

```
%Data initialization section, knot selection, and least squares fit.
%Executed once.
%Get the Cartesian data. Choose a file name included in the function
[x,y,z] = Get_CMM_data;
N = length(x);
%Convert to spherical coordinates. The term base is used to indicate
%results not affected by any scaling of the radial data, otherwise the term
%shift will be used.
[theta_base,phi_base,rad_base] = Get_Spherical_Coordinates(x,y,z);
%Take the mean out of the radial data. It will be added back later.
mean_rad_base = mean(rad_base);
max_rad_base = max(rad_base);
min_rad_base = min(rad_base);
%rad_shift = rad_base - mean_rad_base;
rad_shift = rad_base;
%Adaptively select an optimum set of knots for theta and phi
"The final fit is made to the shifted radial data."
[rad_pred_shift,c_shift,NM_shift,resid_shift,knts_theta,knts_phi,iter]...
  = adaptive_knot_selection(theta_base,phi_base,rad_shift);
disp('Initial Fitting Statistics');
disp(sprintf('Maximum fitting residual, shifted data = %g ', max(resid_shift)));
disp(sprintf('Minimum of fitting residual, shifted data = %g ', min(resid_shift)));
disp(sprintf('Mean of fitting residual, shifted data = %g ',mean(resid_shift)));
disp(sprintf('Standard deviation of fitting residual, shifted data = %g ',std(resid_shift)));
%End data initialization, knot selection, and initial least squares fit
%Begin interactive volume determination for selected grid sizes
disp('Interactive Grid Selection and Volume Estimate');
flag = 1;
```

```
while (flag > 0)
   flag = input('To continue with a new grid type 1, else type -1 > ');
   if (flag <= 0)
       break;
   end
   %Set up grid parameters for generating a longitudinal and latitudinal grid
   nt_grid = input('Enter desired number of theta grid points > ');
   np_grid = input('Enter desired number of phi grid points > ');
   %This function adjusts the extrapolation due to the shifted
   %coefficients by rescaling the predicted radial values on the grid.
    [t_grid,p_grid,rad1_base, N1M1]...
        = Generate_grid_data(c_shift,nt_grid,np_grid, knts_theta,knts_phi,rad_base);
   %Compute the volume
   vol(1) = volume(t_grid,p_grid,rad1_base);
   disp(sprintf('Initial volume in cubic mm = %g',vol(1)));
   Boot = 1;
   Boot = input('Type -1 to skip the bootstrap process > ');
   new_resid = zeros(N,1);
   data_tol = 1e-14;
   if(Boot > 0)
       "Bootstrap process to estimate volume uncertainty and ref volume
        for i = 2:200
            disp(sprintf('Bootstrap iteration %d out of 200.',i));
           for j = 1:N
                u = rand;
                ii = fix(u*N) + 1;
                new_resid(j) = resid_shift(ii);
            end
           new_pred_shift = rad_pred_shift + new_resid;
           %pred values are the extrapolated values to the grid based on
           %the new coefficients c
            c1 = lsq_2D_spline(NM_shift,new_pred_shift,data_tol);
           pred = N1M1*c1 + mean_rad_base;
            delta_plus = max_rad_base - mean_rad_base;
            delta_minus = mean_rad_base - min_rad_base;
            outlyer = find( (pred < mean_rad_base - delta_minus ) | (pred > mean_rad_base + delta_pl
           rnd = rand(length(outlyer),1);
           for ii =1:length(outlyer)
                j = outlyer(ii);
                if (rnd(ii,1) \le 0.5)
                   pred(j) = mean_rad_base - rnd(ii,1)*delta_minus;
                   pred(j) = mean_rad_base + rnd(ii,1)*delta_plus;
                end
            end
            vol(i) = volume(t_grid,p_grid,pred);
            %disp(sprintf('Volume(%d) = %f',i,vol(i)));
        end
```

```
vol_uncert = std(vol);
vol_ref = mean(vol);
disp(sprintf('Volume uncertainty = %f cubic mm',vol_uncert));
disp(sprintf('Reference Volume = %f cubic mm',vol_ref));
end
%clear variables for next pass
clear t_grid p_grid rad1_base
end
```

C Main_Phantom_Volume Support Programs

C.1 Function to Read CMM Data

```
function [x,y,z] = Get_CMM_data
%GET_CMM_DATA - This function opens a file of (x,y,z) data from the CMM
               and recenters the data
%
%Output:
  x,y,z - Re-centered CMM data
%
%Author:
  David E. Gilsinn
%
  Mathematical and Computational Sciences Division
  Information Technology Laboratory
%
   National Institute of Standards and Technology
   Gaithersburg, MD 20899-8910
%Open the CMM data file
%fid= fopen('Tumor_Green_and_Pink_Results_Green_4.txt','r');
%fid= fopen('Tumor_Green_and_Pink_Results_Green_5.txt','r');
%fid= fopen('Tumor_Green_and_Pink_Results_Green_6.txt','r');
%fid= fopen('Tumor_Green_and_Pink_Results_Pink_5.txt','r');
%fid= fopen('Tumor_Green_and_Pink_Results_Pink_6.txt','r');
%fid= fopen('Tumor_Green_and_Pink_Results_Pink_7.txt','r');
%fid= fopen('Green_Results_dense_data_1.txt','r');
%fid= fopen('Green_Results_dense_data_2.txt','r');
%fid= fopen('Green_Results_dense_data_3.txt','r');
%fid= fopen('Pink_Results_dense_data_1.txt','r');
%fid= fopen('Pink_Results_dense_data_2.txt','r');
%fid= fopen('Pink_Results_dense_data_3.txt','r');
%fid= fopen('Cal_121_sphere.txt','r');
fid= fopen('Cal_181_sphere.txt','r');
%fid= fopen('pointcloud.txt','r');
%reads file into a 3-by-inf matrix and takes the transpose. This is
%required due to the way Matlab reads an array.
A=fscanf(fid, '%f %f %f', [3 inf])';
%Now center the data
```

```
n=length(A(:,1));
meanx = sum(A(:,1))/n;
meany = sum(A(:,2))/n;
meanz = sum(A(:,3))/n;
x = zeros(n,1);
y = zeros(n,1);
z = zeros(n,1);
for i=1:n
    x(i,1)=A(i,1)-meanx;
    y(i,1)=A(i,2)-meany;
    z(i,1)=A(i,3)-meanz;
end
fclose(fid);
```

C.2 Function to Euclidean to Spherical Coordinates

```
function [theta,phi,rad] = Get_Spherical_Coordinates(x,y,z)
%GET_SPHERICAL_COORDINATES - Convert Cartesian to Spherical Coord and
%
                      returns the spherical coordinates of data
%Output:
  theta (colatitude from +z axis)
  phi (azimuth from + x axis)
  rad (radial data)
%Author:
% Amelia Tebbe
% Department of Mathematics
\% St. Mary's College of Maryland
% St. Mary's City, MD 20686-3001
B = sph\_coord(x,y,z);
%Read the columns of B into separate vectors.
theta = B(:,1);
phi = B(:,2);
rad = B(:,3);
function D=sph_coord(a,b,c)
%SPH_COORD - Converts Cartesian coordinates to spherical coordinates
% a - Vector of x Cartesian coordinates of points
% b - Vector of y Cartesian coordinates of points
  c - Vector of x Cartesian coordinates of points
%
%OUTPUT:
  D - Matrix of dimension length of a x 3 of Spherical coordinates
```

```
%
        (r,theta,phi) where r is the radius,phi is the azimuth from the
%
        positive x-axis, 0 \le phi \le 2pi, and theta is the colatitude
%
        from the positive z-axis, 0 \le \text{theta} \le \text{pi}. D(:,1) = \text{theta}, D(:,2) = \text{phi},
%
        D(:,3) = r.
n=length(a);
for i=1:n
   x=a(i);
   y=b(i);
   z=c(i);
   r = sqrt(x^2+y^2+z^2);
   \% If the point lies on the positive z-axis
   if (abs(x) \le eps \& abs(y) \le 0 \& abs(z-r) \le eps)
       theta=0;
       phi=0;
   % If the point lies on the negative z-axis
   elseif (abs(x)<=eps & abs(y)<=eps & abs(z+r)<=eps)
       theta=pi;
       phi=0;
   % Cases for each octant
       % 1st octant
       if x>=0 & y>=0 & z>=0
           theta= acos(z/r);
           phi=acos(x/(r*sin(theta)));
       % 2nd octant
       elseif x<0 & y>=0 & z>=0
           theta=acos(z/r);
           phi=acos(x/(r*sin(theta)));
       % 3rd octant
       elseif x<0 & y<0 & z>=0
           theta=acos(z/r);
           phi=pi/2+acos(x/(r*sin(theta)));
       % 4th octant
       elseif x>=0 & y<0 & z>=0
           theta=acos(z/r);
           phi=3*pi/2+ acos(x/(r*sin(theta)));
       % 5th octant
       elseif x>=0 \& y>=0 \& z<0
           theta=acos(z/r);
           phi=acos(x/(r*sin(theta)));
       % 6th octant
       elseif x<0 & y>=0 & z<0
           theta=acos(z/r);
           phi=acos(x/(r*sin(theta)));
```

C.3 MATLAB Function to Adaptively Select θ and ϕ knots

```
function [rad_hat, c,NM, resid,knts_theta,knts_phi,iter]...
   = adaptive_knot_selection(theta,phi,rad_shift)
%ADAPTIVE_KNOT_SELECTION - Choose a good set of knots for the least squares
%
                         fit of the scattered data.
%
%The selection algorithm is based on a suggested algorithm by P. Dierckx,
%Curve and Surface Fitting with Splines, Clarendon Press, Oxford, 2006.
%The object of the algorithm is to place knots in intervals where the
%residual errors are greatest.
%Input:
   theta, phi - Vectors of spherical coordinates for the probe
%
               points. theta and phi are in radians for theta in [0,pi]
%
                and phi in [0,2pi]
%
  rad_shift - rad_shift is in mm and is the radial data shifted by the
%
               mean value
%
%Output:
%
%
  rad_hat - Best fit radial values in mm. This is predicted values of
%
             shifted data.
%
           - Best fit B-spline basis function coefficients
%
  resid - Best fit residuals between rad_hat and rad_shift
%
  knts_theta - Optimum theta knots
%
   knts_phi - Optimum phi knots
%
   iter

    iteration counter

%
%Author:
```

```
David E. Gilsinn
  Mathematical and Computational Sciences Division
  Information Technology Laboratory
%
  National Institute of Standards and Technology
%
  Gaithersburg, MD 20899-8910
%Set a tolerance for computing the pseudo inverse for diagonal matrix S in
%the Singular Value Decomposition (SVD) solution of the least squares
%problem.
data_tol = 1e-14;
%set initial knots with 4 evenly spaced in theta and 8 in phi
knts_theta =[0.0 1.3 1.7 2.0 3 pi]'; %Specified knots for theta
knts_phi = [0 1.7 1.9 2.7 3.5 4 4.5 5.5 2*pi]'; %Specified knots for phi
%compute the fit for the initial data set and define the R_squared
%statistic
c = lsq_2D_spline(NM,rad_shift,data_tol); %Do a least squares fit by SVD
%Get the initial residual
resid = abs(NM*c-rad_shift);
%Compute the initial predicted values
rad_hat = NM*c;
%Compute the initail R squared statistic (coefficient of determination)
rad_bar = mean(rad_shift);
R_square = (sum((rad_hat - rad_bar).^2))/(sum((rad_shift - rad_bar).^2))
%Compute the degrees of freedom
n_dat = length(rad_shift);
df = n_dat - length(c);
%Initialize iteration counter
iter = 1;
while (R_square < 0.98)\&(df >= 1)
   iter = iter + 1; %Update iteration counter
   %Begin the automatic knot adjustment algorithm
   %Compute the sum of squares of the residuals of the fit to all points in
   %the interval (knts_theta(i),knts_theta(i+1))
   lkt = length(knts_theta);
   for i = 1:lkt-1
       t_index = find((knts_theta(i) <= theta) & (theta <= knts_theta(i+1)));</pre>
       Res = resid(t_index);
       del_t(i) = (norm(Res))^2;
   end
   %del_t
   u = find((del_t >= max(del_t))); %Get the maximum of the sums for theta
   "Compute the sum of squares of the residuals of the fit to all points in
   %the interval (knts_phi(i),knts_phi(i+1))
   lkp = length(knts_phi);
   for i = 1:lkp-1
       p_index = find((knts_phi(i) <= phi) & (phi <= knts_phi(i+1)));</pre>
```

```
Res = resid(p_index);
       del_p(i) = (norm(Res))^2;
   end
   %del_p
   v = find(del_p >= max(del_p)); %Get the maximum of the sums for phi
   %Test whether the maximum residuals are with respect to theta or phi and
   %then form a weighted sum of the squares of the residuals. Update the knot
   %arrays.
   if del_t(u) >= del_p(v) %Enter here to select a new theta knot
        t_index = find((knts_theta(u) <= theta) & (theta <= knts_theta(u+1)));
       l_t_index = length(t_index);
       res_t_index = resid(t_index);
       theta_t_index = theta(t_index);
       del_u = del_t(u);
       new_t_knt = 0;
       for k = 1:l_t_index
           new_t_knt = new_t_knt + (res_t_index(k)^2)*...
(theta_t_index(k)/del_u);
       knts_theta(lkt+1,1) = new_t_knt;
       knts_theta = sort(knts_theta);
   else %Otherwise select a new phi knot
       p_index = find((knts_phi(v) <= phi) & (phi <= knts_phi(v+1)));</pre>
       l_p_index = length(p_index);
       res_p_index = resid(p_index);
       phi_p_index = phi(p_index);
       del_v = del_p(v);
       new_p_knt = 0;
       for k = 1:l_p_index
           new_p_knt = new_p_knt + (res_p_index(k)^2)*...
(phi_p_index(k)/del_v);
        end
       knts_phi(lkp+1) = new_p_knt;
       knts_phi = sort(knts_phi);
   end
   %Refit the original data with the new knots
   NM = tensor_prod_spl(theta,phi,knts_theta,knts_phi);
   c = lsq_2D_spline(NM,rad_shift,data_tol);
   resid = abs(NM*c-rad_shift); %Compute the current residual
   rad_hat = NM*c; %Compute the current predicted values
   %Compute the new R-squared statistic and degrees of freedom
   rad_bar = mean(rad_shift);
   R_square = (sum((rad_hat - rad_bar).^2))/(sum((rad_shift - rad_bar).^2))
   df = n_dat - length(c);
end
```

C.4 Function to Compute the Tensor Product of Two B-splines

```
function NM = tensor_prod_spl(theta,phi,knts_theta,knts_phi)
%TENSOR_PROD_SPL - This function creates the bicubic tensor product spline
                 matrix for the spherical coordinate data points in theta
%
                 and phi given specified knots in knts_theta and knts-phi
%
%Input:
   theta, phi - Spherical coordinate angles for CMM data
%
   knts_theta, knts_phi - theta and phi knots
%
%Output:
  NM - Tensor product B-spline matrix
%
%Author:
  David E. Gilsinn
% Mathematical and Computational Sciences Division
% Information Technology Laboratory
% National Institute of Standards and Technology
  Gaithersburg, MD 20899-8910
kstheta = augknt(knts_theta,4); %Augment the knots at both ends
%The next generates four nonzero values for the four cubic B-splines that
%overlap at the specified point. The output is a matrix with a row
%associated with the argument. This will be done one row at a time
lt = length(theta);
lp = length(phi);
for i = 1:lt
   t = theta(i,1);
   N(i,:) = spcol(kstheta,4,t);
end
[row_N,col_N] = size(N);
%Create the associated M array for each data point
ksphi = augknt(knts_phi,4); %Augment the knots at the beginning and end
for j = 1:lp
   p = phi(j,1);
   M(j,:) = spcol(ksphi,4,p);
[row_M,col_M] = size(M);
%Create the tensor product matrix
for r = 1:lt
   for i = 1:col_N
       for j = 1:col_M
          q = (i-1)*col_M + j;
```

```
 \label{eq:NM(r,q) = N(r,i)*M(r,j);} \\ \text{end} \\ \text{end} \\ \text{end} \\ \\ \text{end} \\
```

C.5 Least Squares by Singular Value Decomposition

```
function c = lsq_2D_spline(NM,rad_shift, data_tol)
LSQ_2D_SPLINE - Given the tensor product spline matrix NM and the right
                 hand side data rad, this function produces the least
%
                 square solution of NM*c = rad with minimum norm for c
%
                 using the singular value decomposition. data_tol is the
                 uncertainty in the scattered data being approximated.
%
%
%Input:
   NM - Tensor B-spline matrix
%
   rad_shift - radial data shifted by the mean
% Compute the singular value decomposition of NM, "economy" style,
% dropping unneeded columns in U and V.
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                        ****************
[U,S,V] = svd(NM, 'econ');
%NM = U*S*V' and S is diagonal with nonnegative values [m,n] = size(S);
%Expect that ordinarily m > n
% Compute the pseudoinverse of S. It is a diagonal matrix
% with entries stored in the vector sinv.
\mbox{\ensuremath{\mbox{\%}}} Elements of S less than data_tol are assumed to be zero.
[m,n] = size(S);
sdiag = diag(S);
sinv = zeros(n,1);
for i=1:n
     if (sdiag(i) < data_tol)</pre>
        break
```

```
end
sinv(i) = 1 / sdiag(i);
end
% Form the solution vector as the pseudoinverse of NM times rad_shift.
c = V*(sinv.*(U'*rad_shift));
```

C.6 Generating Data Points on a Surface Grid

```
function [t_grid, p_grid, rad1, N1M1]...
   = Generate_grid_data(c,nt_grid,np_grid,knts_theta,knts_phi,rad)
%GENERATE_GRID_DATA - This function generates a grid of theta/phi points
                   and extrapolates the radial values to these points
%
                   given the coefficient vector c, optimum knots, and
%
                   the spherical coordinate data radial values, rad
%
%Input:
   c - The B-spline coefficients from the least squares fit to the
%
       spherical coordinate data values
%
  nt_grid - The number of desired theta grid values
  np_grid - The number of desired phi grid values
%
%
  knts_theta - The optimum theta knots
%
  knts-phi - The optimum phi knots
%
  rad - Unshifted radial data from the spherical coordinates
%
%Output:
%
   t_grid - Theta values at the grid points along the theta coordinate
   p_grid - Phi values at the grid points along the phi coordinate
%
   rad1 - Extrapolated radial values at the (t_grid,p_grid) coordinates
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"Select an evenly spaced grid of (theta, phi) pairs
t_grid = linspace(0,pi,nt_grid)';
p_grid = linspace(0,2*pi,np_grid)';
"Compute the tensor product matrix for the selected grid
lt = length(t_grid);
```

```
lp = length(p_grid);
kstheta = augknt(knts_theta,4); %Augment the knots at both ends
ksphi = augknt(knts_phi,4); %Augment the knots at the beginning and end
%Generate the B-spline tensor product at the grid points. Do it point at a
%time to take care of non-sorted arrays
for i = 1:lt
    t = t_grid(i);
    N1(i,:) = spcol(kstheta,4,t);
end
[row_N1, col_N1] = size(N1);
for j = 1:lp
    p = p_grid(j);
    M1(j,:) = spcol(ksphi,4,p);
[row_M1,col_M1] = size(M1);
for r1 = 1:row_N1
    for r2 = 1:row_M1
        r = (r1-1)*row_M1 + r2;
        for i = 1:col_N1
            for j = 1:col_M1
                q = (i-1)*col_M1 + j;
                N1M1(r,q) = N1(r1,i)*M1(r2,j);
            end
        end
    end
end
[r_n1m1,c_n1m1] = size(N1M1);
%Some important parameters of the unshifted radial data
mean_rad = mean(rad);
max_rad = max(rad);
min_rad = min(rad);
%Compute the new radii by adding back the mean of the original radial data
%rad1 = N1M1*c + mean_rad;
rad1 = N1M1*c;
%Adjust the values of rad1 values that fall outside of the original data
delta_plus = max_rad - mean_rad;
delta_minus = mean_rad - min_rad;
outlyer = find( (rad1 < mean_rad - delta_minus ) | (rad1 > mean_rad + delta_plus));
rnd = rand(length(outlyer),1);
for i =1:length(outlyer)
    j = outlyer(i);
    if (rnd(i,1) \le 0.5)
        rad1(j) = mean_rad - rnd(i,1)*delta_minus;
    else
        rad1(j) = mean_rad + rnd(i,1)*delta_plus;
    end
```

C.7 Computing the Volume Based on Grid Values

```
function vol = volume(t_grid,p_grid,rad1)
%VOLUME - This function computes the volume of a phantom based on a
         specified grid size using the Divergence Theorem. We accumulate
%
         the facet contributions to the volume estimate of the phantom.
%
%Input:
  t_grid - theta values at grid points
  p_grid - phi values at grid points
\% rad1 - extrapolated radial values at grid points
%
%Output:
  vol - volume in cubic millimeters
%
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lt = length(t_grid);
lp = length(p_grid);
%Transfer the rad1 array of estimated radii on a specified grid to a two
%dimensional array
nv = lt-1;
nh = lp-1;
for i = 1:nv+1
   for j = 1:nh+1
       p = (i-1)*(nh+1) + j;
       r(i,j) = rad1(p);
   end
end
r(1,1) = median(r(1,:));
r(nv+1,1) = median(r(nv+1,:));
%Initialize the volume
vol = 0;
%Loop through the layers of theta. Begin with the North Pole layer.
%Set up facet triangle vertices in counterclockwise manner. The top point
%will always be the north pole.
v1(1) = 0;
v1(2) = 0;
v1(3) = r(1,1);
```

```
%loop through each facet in the first layer
for j = 1:nv
    v2(1) = r(2,j)*sin(t_grid(2))*cos(p_grid(j));
    v2(2) = r(2,j)*sin(t_grid(2))*sin(p_grid(j));
    v2(3) = r(2,j)*cos(t_grid(2));
    v3(1) = r(2,j+1)*sin(t_grid(2))*cos(p_grid(j+1));
    v3(2) = r(2,j+1)*sin(t_grid(2))*sin(p_grid(j+1));
    v3(3) = r(2,j+1)*cos(t_grid(2));
   v = v2 - v1;
    w = v3 - v1;
   n = normal_vec(v, w);
   term = facet_term(n,v1,v2,v3);
    vol = vol + term;
end
%Next loop through layers 2 through nv-1. There are nh 4-vertex patches at
%each level. Each patch will be divided into two triangles. The four
%vertices of the patches will be identified in counterclockwise order as
%(t_grid(i),p_grid(j),r(i,j)),
(t_{grid}(i+1), p_{grid}(j), r(i+1, j)), (t_{gris}(i+1), p_{grid}(j+1), r(i+1, j+1)),
%(t_grid(i),p_grid(j+1),r(i,j+1))
for i = 2:nv-1
    for j = 1:nh
        %Get contribution of left triangle
        v1(1) = r(i,j)*sin(t_grid(i))*cos(p_grid(j));
        v1(2) = r(i,j)*sin(t_grid(i))*sin(p_grid(j));
        v1(3) = r(i,j)*cos(t_grid(i));
        v2(1) = r(i+1,j)*sin(t_grid(i+1))*cos(p_grid(j));
        v2(2) = r(i+1,j)*sin(t_grid(i+1))*sin(p_grid(j));
        v2(3) = r(i+1,j)*cos(t_grid(i+1));
        v3(1) = r(i+1, j+1)*sin(t_grid(i+1))*cos(p_grid(j+1));
        v3(2) = r(i+1,j+1)*sin(t_grid(i+1))*sin(p_grid(j+1));
        v3(3) = r(i+1,j+1)*cos(t_grid(i+1));
        v = v2-v1;
        w = v3-v1;
        n = normal_vec(v, w);
        term = facet_term(n,v1,v2,v3);
        vol = vol + term:
        v2 = v3;
        v3(1) = r(i,j+1)*sin(t_grid(i))*cos(p_grid(j+1));
        v3(2) = r(i,j+1)*sin(t_grid(i))*sin(p_grid(j+1));
        v3(3) = r(i,j+1)*cos(t_grid(i));
        v = v2-v1;
        w = v3-v1;
        n = normal_vec(v, w);
        term = facet_term(n,v1,v2,v3);
        vol = vol + term;
```

```
end
end
%Loop through the final layer at the South Pole
for j = 1:nh
   v1(1) = r(nv,j)*sin(t_grid(nv))*cos(p_grid(j));
   v1(2) = r(nv,j)*sin(t_grid(nv))*sin(p_grid(j));
   v1(3) = r(nv,j)*cos(t_grid(nv));
   v2(1) = 0;
   v2(2) = 0;
   v2(3) = -r(nv+1,1);
   v3(1) = r(nv, j+1)*sin(t_grid(nv))*cos(p_grid(j+1));
   v3(2) = r(nv, j+1)*sin(t_grid(nv))*sin(p_grid(j+1));
   v3(3) = r(nv, j+1)*cos(t_grid(nv));
   v = v2-v1;
   w = v3-v1;
   n = normal_vec(v, w);
   term = facet_term(n,v1,v2,v3);
   vol = vol + term;
end
%Get the final volume
vol = 1/6*vol;
function n = normal_vec(v,w)
%NORMAL_VEC - This function computes the unit normal vector n to the
%
              plane determine by v and w in counterclockwise form. The
%
              function computes the cross product of v and w using the
%
              right hand rule between v and w.
n(1) = v(2)*w(3) - v(3)*w(2);
n(2) = v(3)*w(1) - v(1)*w(3);
n(3) = v(1)*w(2) - v(2)*w(1);
n = n/norm(n);
```