# Fast Analytical Computation of Richards's Smooth Molecular Surface

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#### Abstract

An algorithm for rapid computation of Richards's smooth molecular surface is described. The entire surface is computed analytically, triangulated, and displayed at interactive rates. The faster speeds for our program have been achieved by algorithmic improvements, parallelizing the computations, and by taking advantage of the special geometrical properties of such surfaces. Our algorithm is easily parallelizable and it has a time complexity of  $O(k \log k)$  over n processors, where n is the number of atoms of the molecule and k is the average number of neighbors per atom.

# 1 Introduction

The smooth molecular surface of a molecule is defined as the surface which an exterior probe-sphere touches as it is rolled over the spherical atoms of that molecule. This definition of a molecular surface was first proposed by Richards [16]. This surface is useful in studying the structure and interactions of proteins, in particular for attacking the protein-substrate docking problem. An example of such a molecular surface for Felix with a probe-sphere radius of 1.4Å can be seen in Figure 1.

Present systems to compute the surfaces of molecules are batch-oriented. They take a few minutes to compute the surface for a couple of thousand atoms. Our goal has been to compute and display these surfaces at interactive rates, by taking advantage of results from the field of computational geometry, making further algorithmic improvements, and parallelizing the computations.

Interactive computation of molecular surfaces should benefit biochemists in three important ways. First, the ability to change the probe-radius interactively helps one study the surface. Second, it helps in visualizing the changing surface of a molecule as its atom positions are changed. These changes in atom positions could be due to user-defined forces as the user attempts to modify a molecular model on a computer. Third, it assists in incorporating the effects of the solvent into the overall potential energy computations during the interactive modifications of a molecule on a computer.

# 2 Previous and Related Work

The first molecular surface computation algorithms were numerical in nature (ie. they were computed by sampling) Connolly [2], Greer [12]. Connolly [2] computes the sampled surface (also known as the *dot-surface*) by placing a probe tangent to either one atom, or two atoms, or three atoms and



Figure 1: Smooth Molecular Surface for Felix

checking to see if it intersects any of the other neighboring atoms. If it does not and it is tangent to (i) one atom, then a dot is placed at the point of tangency between the probe and that atom, (ii) two atoms, then a concave arc of dots connecting the two points of tangency is created, (iii) three atoms, then a concave spherical triangle of dots is created between the three points of tangency. This generates a dot-representation of the entire surface.

The analytic computation of the molecular surface was also first done by Connolly [3], [4]. Here a molecular surface is represented by a collection of spherical and toroidal patches as follows:

- The molecular surface for the regions of a molecule where the probe is in contact with a single atom are modeled by convex spherical patches.
- The molecular surface for the regions of a molecule where the probe is in simultaneous contact with two atoms are modeled by saddle-shaped toroidal patches.
- The molecular surface for the regions where the probe is in simultaneous contact with three atoms are modeled by concave spherical triangular patches.

The issues of triangulation of such surfaces are discussed by Connolly in [5].

Only recently have the issues of algorithmic complexity of these algorithms begun to be addressed. Let n be the number of atoms in a molecule and let k be the average number of *neighboring* atoms for an atom in the molecule. By *neighboring* we mean the atoms that are near enough to affect probe placement on a particular atom. Yip and Elber [18] present an algorithm for computation of the list of neighboring atoms that is linear in n. It is based on spatial subdivision by a global grid. Perrot *et al.* [13], [14] present a O(kn) algorithm that generates an approximation to the solvent-accessible surface. In this approximation, every concave spherical triangular patch between three atoms is represented by a planar triangle with vertices at the centers of these three atoms. Saddle-shaped toroidal regions and convex spherical patches are ignored. In terms of sequential algorithmic complexity this is good, however some points remain unaddressed here. This algorithm is inherently sequential, as it always needs to start from some concave spherical triangular region of the molecule and from there it proceeds by adding an adjacent face at a time. Besides being hard to parallelize, it fails for the cases where the solvent-accessible surface folds back to intersect itself or where the molecule has two or more sub-parts connected by only two overlapping spheres. Also, it cannot generate the interior cavities of a molecule.

In computational geometry, the  $\alpha$ -hull has been defined as a generalization of the convex hull of point-sets by Edelsbrunner, Kirkpatrick, and Seidel [6], [8]. For  $\alpha > 0$ , the  $\alpha$ -hull of a set of points P in two-dimensions is defined to be the intersection of all closed complements of discs with radius  $\alpha$  that contain all points of P. If we generalize this notion of  $\alpha$ -hulls over point-sets to the corresponding hulls over spheres of unequal radii in three-dimensions, we would get the molecular surface (along with the surface defining the interior cavities of the molecule). It has been shown in [8] that it is possible to compute the  $\alpha$ -hulls from the Voronoi diagram of the points of P. For  $\alpha = \infty$  the  $\alpha$ -hull over the set of points P is the same as their convex hull. Richards [16] had also suggested computing the molecular surface by computing a 3D Voronoi diagram first and then using its faces to determine which nearby atoms to consider.

Edelsbrunner and Mücke [9] extend the definition of  $\alpha$ -hulls to points in three-dimensions. Here an  $\alpha$ -shape over a set of points P has been defined to be the polytope that approximates the  $\alpha$ -hull over P, by replacing circular arcs of the  $\alpha$ -hull by straight edges and spherical caps by triangles. An  $\alpha$ -shape of a set of points P is a subset of the Delaunay triangulation of P. Edelsbrunner in [7], extends the concept of  $\alpha$ -shapes to deal with weighted points (i.e. spheres with possibly unequal and non-zero radii) in three-dimensions. An  $\alpha$ -shape of a set of weighted points  $P_w$  is a subset of the regular triangulation of  $P_w$ . Since these methods involve computing the entire triangulation first and then culling away the parts that are not required, their complexity is  $O(n^2)$  in time. This is worst-case optimal, since an  $\alpha$ -shape in three-dimensions could have a complexity of  $O(n^2)$ . We next discuss a different approach that works better for environments where the maximum density of P in a given volume is some constant smaller than n. Molecules are a good example of such environments.

# 3 Our Approach

Our goal has been to formulate a parallel analytical molecular surface algorithm that has expected linear complexity with respect to the total number of atoms of a molecule. For achieving this goal, we have avoided computation of the complete three-dimensional regular triangulation over the entire set of atoms - a process that takes time  $O(n^2)$ , where n is the number of atoms in the molecule.

Molecules considered as a collection of weighted points in three-dimensions, where the coordinates of each point  $p_i$  correspond to the center of atom i and the weight  $r_i$  is the radius of atom i, have two interesting properties: (i) the minimum distance  $d_{ij}$  between any two points  $p_i$  and  $p_j$  is strictly positive,  $d_{ij} > 0$  and (ii) the set of all the weights can be bounded from above and below by non-zero values,  $r_{min} \leq r_i \leq r_{max}$ . We take advantage of the first property to arrive at better running times for our algorithm. Stated simply, the first property says that the number of neighboring atoms within a fixed distance from any atom *i*, is always bounded from above by a constant  $k_{max}$  that depends on the minimum spacing between any two atoms. If the average number of neighbors for an atom is *k*, then we can just compute an approximation to the power cell (discussed in Section 3.1), which we call a *feasible cell*, by considering only these neighbors. Each feasible cell can be computed in parallel in time  $O(k \log k)$ . For *n* atoms, this task requires *n* processors, each processor computing the feasible cell for one atom.

### 3.1 Formal Notation

Here we consider the underlying space to be three-dimensional Euclidean Space  $\Re^3$ , although these results can be generalized to higher dimensions.

Let  $M = \{S_1, \ldots, S_n\}$ , be a set of spheres, where each sphere,  $S_i$ , is expressed as a pair  $\langle c_i, r_i \rangle$ ,  $c_i$  being the center of the sphere and  $r_i$  being the radius of the sphere.

Let x, y be two points. Define d(x, y) to be the Euclidean distance between x and y. The power of a point x with respect to a sphere  $S_i$  is defined as  $p(x, S_i) = d^2(x, c_i) - r_i^2$  Thus,  $p(x, S_i) < 0, = 0, > 0$ , depending on whether x lies inside  $S_i$ , on the boundary of  $S_i$ , or outside  $S_i$ , respectively.

Henceforth, we shall be assuming that the atoms of a molecule are represented as spheres and will be using the terms *atom i* and  $S_i$  interchangeably. Let the radius of the probe-sphere be R. Then we define the *extended-radius sphere* for atom *i* to be  $\Psi_i = \langle c_i, r_i + R \rangle$ . This extended-radius sphere  $\Psi_i$  is the locus of the possible centers of the probe-sphere when it is in contact with atom *i*.

Define a chordale  $\Pi_{ij}$  of two spheres  $\Psi_i$  and  $\Psi_j$  as  $\Pi_{ij} = \{x | p(x, \Psi_i) = p(x, \Psi_j)\}$ .  $\Pi_{ij}$  is a plane perpendicular to the line joining  $c_i$  and  $c_j$ . If  $\Psi_i$  and  $\Psi_j$  overlap,  $\Pi_{ij}$  passes through the circle that is common to the boundaries of  $\Psi_i$  and  $\Psi_j$ . If  $\Psi_i$  and  $\Psi_j$  do not overlap, it lies between  $c_i$  and  $c_j$ . Define the halfspace  $H_{ij}$  as  $H_{ij} = \{x | p(x, \Psi_i) < p(x, \Psi_j)\}$ .

Thus, whereas  $\Pi_{ij} = \Pi_{ji}, H_{ij} \neq H_{ji}$ . Define the *feasible cell*,  $F_i$ , for atom *i* as  $F_i = \bigcap_j H_{ij}$ .

So far, the definitions that we have given for a *feasible cell* correspond exactly to the *power cells* of a power diagram as defined by Aurenhammer [1]. The reason we prefer to use a different term for these will be made clear in Section 3.3.

### 3.2 Determination of Neighboring Atoms

Determination of neighboring atoms can be done by spatial grid subdivision into voxels, and assigning atoms to appropriate voxels. We define the region of influence,  $\rho_i$ , for atom i to be the sphere  $\langle c_i, r_i + 2R + \max_{j=1}^n r_j \rangle$ . Then for computing the list of neighboring atoms,  $N_i$ , for atom i, one needs to find all the atoms that are close enough to affect probe placement on atom i. Formally,  $N_i = \{j | d(c_i, c_j) < r_i + 2R + r_j\}$ , or equivalently,  $N_i = \{j | \Psi_i \cap \Psi_j \neq \phi\}$ . Intuitively, an atom j is a neighbor to atom i, if it is possible to place a probe such that it is in contact with both  $S_i$  and  $S_j$  (without considering any hindrance due to other atoms). Since, by definition  $N_i \subseteq \rho_i$ , to compute the list of neighboring atoms for atom i, one needs to look at all the atoms that lie in the voxels that

intersect  $\rho_i$ . Let the average number of neighboring atoms be k. Note that k grows as  $R^3$  assuming that the atoms are uniformly distributed. In Figure 2 atoms  $j_1$  and  $j_3$  are neighbors to atom i, but atom  $j_2$  is not.



Figure 2: Determination of Neighboring Atoms

### **3.3** Determination of Surface Atoms

Here the aim is to determine the atoms that are buried in the interior of the molecule and would not therefore directly participate in the final definition of the smooth molecular surface. This step is not crucial to the linear time complexity of the overall algorithm but it helps in improving the execution times.

For every atom *i*, we first compute  $N_i$  as described in Section 3.2. Then we compute  $F_i = \bigcap_{j \in N_i} H_{ij}$ . If  $F_i = \phi$ , atom *i* is totally buried and cannot be a surface atom. This checking for nullity is done by Seidel's randomized linear programming algorithm that has linear time and is quite fast in practice [17].

It might be useful to point out how the feasible cells  $F_i$  we are computing are different from the power cells described by Aurenhammer [1]. The difference arises from the fact that each of these cells  $F_i$  are being computed independently of the other, and for each cell  $F_i$  we consider only the hyperplanes whose indices occur in  $N_i$ . This is best illustrated by Figure 3, where the two types of cells are shown. Thus, for construction of  $F_i$  we use only those chordales  $\Pi_{ij}$ , as defined in section 3.1, that arise from overlapping extended-radius spheres  $\Psi_i$  and  $\Psi_j$ . However, for forming the power cells  $C_i$ , we use all the chordales  $\Pi_{ij}$  regardless of whether  $\Psi_i$  and  $\Psi_j$  intersect or not.

In Figure 3, we show these differences for power cells and feasible cells defined over circles. The power cell  $C_3$  contains two edges and one vertex as does the corresponding feasible cell  $F_3$ . However, whereas the power cells  $C_1$  and  $C_2$  have two edges and one vertex, the corresponding feasible cells  $F_1$  and  $F_2$  have only one edge each, with no vertices.

We should also point out here that the feasible cells that occur on the boundary atoms are not compact (actually, they are neither closed nor bounded). Although the details of triangulation of these surfaces will appear elsewhere we would like to mention here that we use the vertices of  $F_i$ 



Figure 3: Difference between Power Cells and Feasible Cells

during triangulation. For this purpose we would like all  $F_i$  to be compact (closed and bounded) polytopes. To achieve this we bound the entire molecule by four extra planes forming a regular tetrahedron. These are included in  $N_i$ , for all *i*.

#### **3.4** Determination of Surface Patches

Determination of the vertices defining the convex spherical, concave spherical and toroidal patches is the most crucial (and time-consuming) part of the whole algorithm.

If one computes a three-dimensional  $\alpha$ -shape polytope for the set of atoms in a molecule, with  $\alpha$  = probe-radius, then the torii occur along the edges, the concave spherical triangular patches correspond to the faces, and the convex spherical patches correspond to the vertices of this polytope. The method given by Edelsbrunner [7] finds these edges by first computing the entire three-dimensional regular triangulation, an  $O(n^2)$  approach. We show here an equivalent method for computing the three-dimensional  $\alpha$ -hulls for molecules in parallel time  $O(k \log k)$  over n processors

To compute  $F_i$ , we compute the convex hull of the points dual to the  $H_{ij}$  in the dual-space, as described in [15]. This is an  $O(k \log k)$  time process. After this, we compute the dual of the convex hull to get the feasible cell  $F_i$ . The intersection of the feasible cell  $F_i$  with  $\Psi_i$  gives rise to a set of closed components on  $\Psi_i$ . Since  $F_i$  is convex, these components will be non-intersecting. Each of these closed components  $\partial c_j$ , divide  $\Psi_i$  into two connected regions, say  $R_{j_0}$  and  $R_{j_1}$ . For one of these, say  $R_{j_m}$ , it will be true that  $R_{j_m} = R_{j_m} \cap F_i$ . We define  $R_{j_m}$  to be the interior of the closed component  $\partial c_i$ . To find these components, we determine the edges and faces of  $F_i$  that intersect  $\Psi_i$ .

After a connected component has been determined on  $\Psi_i$ , it is easy to generate the surface patches. The arcs defining the boundary of this component determine the locus of the center of the probe while it is in contact with two atoms. These therefore are used to generate the toroidal patches. The vertices of this component, where two arcs intersect, define the positions of the center of the probe while it is in contact with three atoms. These are used to generate the concave spherical triangular patches. The interiors of the components on  $\Psi_i$ , correspond to the positions of the center of the probe while it is tangent to only atom *i*. These are used to generate the convex spherical patches.

This is shown in the Figure 4 for a component defined by three hyperplanes. The interior of this component is shown unshaded.



Figure 4: Determination of Molecular Surface Patches from Components

### 3.5 Parallelization

Our approach to computing the smooth molecular surface can be parallelized over all the atoms of the molecule. Since each of the steps as described above can be carried out independently for each atom, the complexity of our algorithm over n processors would be  $O(k \log k)$ . If the number of available processors p < n, we can allocate  $\frac{n}{p}$  atoms per processor to get a time complexity of  $O(\frac{nk \log k}{p})$ . These bounds hold in a CREW PRAM model of parallel computation.

### 3.6 Robustness

In the algorithms for computing the convex hull of a set of points, it is assumed that the points are in a general position, i.e. no more than d points lie on the same d-1 dimensional hyperplane. In reality this assumption often fails to hold, leading to problems. For example, planar benzene rings occur often in proteins, causing six carbon and six hydrogen atoms to be all coplanar.

One of the recent approaches to solving this problem has been to perturb the input point set slightly to avoid these degeneracies. We are using the deterministic perturbation scheme proposed by Emiris and Canny [10], which perturbs the  $j^{th}$  dimension of the  $i^{th}$  point as:

$$p_{i,j}(\epsilon) = p_{i,j} + \epsilon(i^j \mod q) 1 \le i \le n, 1 \le j \le d \tag{1}$$

where  $\epsilon$  is a symbolic infinitesimal and q is the smallest prime greater than n.

### 4 Results

Our implementation has been done on Pixel-Planes 5 [11], although it is general enough to be easily portable to any other parallel architecture. Table 1 shows our timings for computation and display of the molecular surface for various molecules for a probe-radius of 1.4Å. For these results we were using either a configuration of 13 or 26 Intel i860 graphics processors as shown below. At present, we are representing the molecular surface by triangles and the column *No. of Triangles* in the table below refers to the complexity of the computed surface. As can be seen the value of k is fairly constant for a given probe-radius over different molecules.

Molecule	No. of atoms	Times (in sec)		k	No. of Triangles
		13 procs	26 procs		
Crambin	396	0.46	0.25	45	$18.5 \mathrm{K}$
Felix	613	0.76	0.41	41	$36.5 \mathrm{K}$
Dihydrofolate reductase	3123	3.23	1.63	44	$95.5\mathrm{K}$
Superoxide dismutase	4386	4.51	2.25	46	$125.4\mathrm{K}$

Table 1: Smooth Molecular Surface Generation Times

Figure 5 shows the smooth molecular surfaces for Crambin with probe-sphere radii of (a)  $1.0\mathring{A}$ , (b)  $1.4\mathring{A}$ , (c)  $2.8\mathring{A}$ , and (d)  $5.0\mathring{A}$ . Figure 6 shows the smooth molecular surface for Dihydrofolate reductase with probe-sphere radius of  $1.4\mathring{A}$ . Figure 7 shows the smooth molecular surface for Superoxide dismutase with probe-sphere radius of  $1.4\mathring{A}$ .

### 5 Conclusions and Future Work

We have presented a parallel algorithm for computing the molecular surfaces in parallel time  $O(k \log k)$  over *n* processors. This is sufficiently general enough to be used for computation of  $\alpha$ -hulls and  $\alpha$ -shapes for a given value of  $\alpha$  as long as no two points are arbitrarily close. Our algorithm would give an order of magnitude improvement over the best known previous algorithms for molecules with large values of *n* - both in sequential as well as in parallel implementations.

At present we are not using any incremental temporal information in constructing these surfaces. Thus, if the atoms move slightly from their positions, the whole surface has to be recomputed from the beginning. Assuming the atoms of the molecule move along continuous trajectories, it should be possible to compute such surfaces (and indeed  $\alpha$ -hulls and  $\alpha$ -shapes) incrementally by using the information from previous time steps.

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(a)



(b)

Figure 5: Smooth Molecular Surfaces for Crambin



(c)



(d)

Figure 5: Smooth Molecular Surfaces for Crambin(contd)



 $Figure \ 6: \ Smooth \ Molecular \ Surface \ for \ Dihydrofolate \ reductase$ 



Figure 7: Smooth Molecular Surface for Superoxide dismutase

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