ESCHER: A New Docking Procedure Applied to the Reconstruction of Protein Tertiary Structure

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ABSTRACT Evaluation of Surface Complementarity, Hydrogen bonding, and Electrostatic interaction in molecular Recognition (ESCHER) is a new docking procedure consisting of three modules that work in series. The first module evaluates the geometric complementarity and produces a set of rough solutions for the docking problem. The second module identifies molecular collisions within those solutions, and the third evaluates their electrostatic complementarity. We describe the algorithm and its application to the docking of cocry stallized protein domains and unbound components of protein-protein complexes. Furthermore, ESCHER has been applied to the reassociation of secondary and supersecondary structure elements. The possibility of applying a docking method to the protein structure prediction is discussed. Proteins 28: 556–567, 1997. © 1997 Wiley-Liss, Inc.

Key words: molecular recognition; automated docking; protein domains; secondary structure elements

INTRODUCTION

Numerous automated methods for the prediction of protein complex formation have been developed.1,2 They are generally based on the rigid body approximation3–5: some rely on a geometric criterion and on a simplified representation of the protein surface, “soft docking” methods,6–8,10 whereas others combine a shape complementarity search with subsequent energy refinement.11,12 Many algorithms can successfully reconstruct a protein complex starting from the experimentally determined conformation of the components in the bound state. In some cases, some success, albeit with lower accuracy, has been obtained by docking elements whose structures have been determined separately.8,10,11

In the rigid body approach, the position and orientation of a protein is completely described by three translational and three rotational degrees of freedom. This approach is, therefore, widely used in automated methods, when molecular flexibility is not expected to affect much the geometry of the interacting structures.

On the other hand, when the docking problem involves the recognition between a rigid protein and a flexible ligand, the rigid body approximation is not applicable, and more conformational degrees of freedom have to be taken into account.14,15 The basic principles underlying rigid body and flexible docking are common, but they generally require different computational techniques.

We explore here the possibility of applying a new soft docking rigid body method to components of intermediate size when compared with the “classical” targets of the docking methodologies discussed above. We chose to dock protein components, such as protein domains and secondary and supersecondary structures, and to explore the limits of the rigid body approximation when applied to a series of test cases displaying different sizes, structural complexities, and root-mean-square (rms) deviations from the structures of the protein in their bound conformations. The ultimate goal of this approach is the assembly of a whole protein tertiary structure starting from its secondary structure elements.

The reconstruction of a protein structure starting from its secondary structure elements has already been tried on a very small number of proteins, generally relying on knowledge-based potentials16–18 as well as on other methods.19–21 To tackle this problem, a novel docking program was used as a tool. To this end, we have substantially modified the PUZZLE8 procedure to achieve a more accurate description of the surface of small proteins and more effective matching and scoring criteria. Furthermore, the new program, ESCHER, takes advantage of two filter modules that reject solutions resulting in steric (BUMPS) or electrostatic (CHARGES) clashes. The addition of these modules allows less stringent criteria for geometric complementarity to be considered and, consequently, contributes to a further “softening” of the docking procedure.

Abbreviations: PDB, Protein Data Bank; rmsd, root-mean-square deviation

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ESCHER was applied to a series of two-domain proteins, by separating the domains and by attempting to reconstruct the native protein from the two-domain crystallographic structures, and to protein complexes starting from their unbound components. Furthermore, we defined a limited set of test cases in which the docking partners were represented by secondary and supersecondary structures separated from their protein. The docking experiments were performed with crystallographic structures and structural models. This was done to assess whether and how minor deviations from the idealized secondary structure backbone conformation or lack of knowledge of the exact side-chain conformation could affect the docking results.

METHODS

Overview

ESCHER is a rigid docking program, written in C language, assembled from three modules that work in series: SHAPES, BUMPS, and CHARGES. The first module takes as input the solvent-accessible surface of the target and probe peptides and provides a list of 500 solutions, ranked on the basis of geometric complementarity. Each solution is identified by the translations and rotations to be applied to the probe to match the target. The second module, BUMPS, analyzes the molecular collisions within the solution complexes, and CHARGES evaluates their electrostatic complementarity. All the solutions generated by SHAPES are subjected to the BUMPS filter and grouped together whenever their deviation from the member of the cluster with the highest geometric complementarity is lower than 2.5 Å. Finally, each group is represented by the solution with the best CHARGES value.

SHAPES Module

SHAPES analyzes two peptide structures, evaluates their complementarity in all possible orientations, and proposes a number of docking solutions, each identified by the Tᵢ translations and the Rᵢ rotations to be applied to the second element (the probe) with respect to the first element (the target). Each structural element is cut in parallel slices, and each slice is described as a polygon with sides of identical length. The polygons representing the target are compared with the probe polygons, and regions of complementarity are evaluated. A complete search in the rotation space is done by rotating the probe in all possible orientations with respect to the target.

The flowchart of the geometric module is reported in Figure 1. Each box in the flowchart corresponds to one paragraph in the following detailed SHAPES description.

(b-1) Structural elements description

The solvent-accessible surface of each protein structural element is generated from its atomic coordinates by using a probe radius of 1.8 Å (a density of 10 dots/Å²). Each surface is cut in parallel slices 1.5 Å thick, orthogonal to the Z axis (Fig. 2a). Each slice is transformed into a polygon (a few hundred of vertices) by an "intelligent contouring" algorithm (Fig. 2b,c,d).

(b-2) Starting position in Z translations

The probe polygons are translated along the Z axis until the probe top polygon is at the level of the target bottom polygon (Fig. 3a).

(b-3) Polygon matching

Each polygon of the target can now be compared with the corresponding probe polygon belonging to
the same Z plane. Each one of the $m$ probe polygon sides is superposed to each one of the $n$ corresponding target polygon sides, by applying the necessary translations (Tx and Ty) and rotation (Rz). A measure of geometric complementarity ("complementarity score") is associated to each one of the $mn$ different superpositions and to the corresponding Rz, Tx and Ty transformation (Fig. 3b); the complementarity score is the number of corresponding consecutive polygon vertices whose distance is lower than 1.6 Å (Fig. 3c). Whenever a small cavity is present in the "interface" between two polygons, the complementarity of the preceding or of the following polygon sides still can contribute to the geometric score.

(b-4) Match grouping

SHAPES solutions are generated by grouping the polygon matches coming from different Z planes but representing the same complementary region. To group polygon matches and define a complementary region extending through different Z planes, a
three-dimensional space is defined in which a polygon match is represented by a dot with the corresponding Tx, Ty, and Rz coordinates. In this space, we define a grouping grid with a cell size of 3 Å along the Tx and Ty axes and 12° along the Rz axis. Whenever a new polygon match is found, its complementarity value is added into the corresponding element of the grouping grid.

At the end of the match grouping step, the indexes of the grid element that scored the highest complementarity value represent the Tx, Ty, and Rz transformations to apply to the probe to correctly match the highest number of polygon pairs. These Tx, Ty, and Rz values, together with the Tz, Rx, and Ry explored stepwise as explained below, define a solution complex, associated with the complementarity value stored into its grouping matrix element.

(b-5) Z Translations

All the polygons of the probe are translated by $-1.5$ Å along the Z axis, thus shifting the pairs of polygons that are matched in the subsequent "polygon matching" phase. This corresponds to a stepwise exploration of the Tz degree of freedom, where $\Delta Tz = 1.5$ Å. This step is executed until the probe top polygon reaches the level of the target bottom polygon. All the probe polygons are therefore compared with the target protein polygons.

(b-6) X-Y rotations

In this step, the probe is rotated around the X and Y axis to analyze all the possible orientations with respect to the target. Rx and Ry are explored stepwise: $-180° \leq Rx < 180°$, $-90° \leq Ry < 90°$, with $\Delta Rx = \Delta Ry = \Delta R$. This is equivalent to exerting a complete search in the space of all possible target-probe relative orientations, by rotating one of the two peptides by $\Delta R$ steps around the X and Y axis. In the docking experiments described, $\Delta R = 10°$. This value can be changed, when necessary, according to the probe dimensions. The complete exploration of the rotation space around the Z axis (Rz) is exerted continuously within the polygon pairs comparison. At the end of this step, the probe is described into a new set of polygons.

(b-7) Target orientation

The cylindrical symmetry inherent to the kind of description adopted is very convenient to transform a three-dimensional surface matching problem into a simplified two-dimensional polygon comparison but offers a very poor description of the structural element poles. One of the two elements (the probe) is rotated in all possible orientations and is therefore optimally described at least once. The problem may concern the target element, which remains in the same orientation during the whole procedure. One may overcome this problem describing the target element in more than one set of parallel polygons and repeating the comparison algorithm on the minimal number of polygon sets which one considers necessary. Cases studied to address this problem indicate that, in general, two orthogonal sets of polygons are sufficient to detect a region of geometric complementarity in at least one of the target orientations tested (data not shown).
In all of the test cases discussed below, the target has been described only once in an optimal orientation (with the interaction site parallel to the Z axis). We also verified that the results of the procedure are essentially independent of the orientation of the target in the reference system (a target rotation around the X or Y axis of up to 45° does not affect the results).

**BUMPS Module**

SHAPES module evaluates complementarity on the basis of distances, which are intrinsically positive numbers. Consequently, it cannot distinguish between polygon perimeters that diverge or interpenetrate following or preceding a complementary region. Furthermore, a docking solution with a good “local” match may be accompanied by a steric clash in another part of the proposed complex. The solution list of the geometric module may, therefore, include sterically impossible structures, whose components share regions of shape complementarity but also overlap somewhere else.

The BUMPS module analyzes the distance between each atom of the target structural element and each atom of the probe in the complexes proposed by SHAPES and evaluates a “bumps” value.

\[
\text{BUMPS} = \sum_{a} \sum_{b} B_{ab}
\]

where

\[
B_{ab} = \begin{cases} 
0 & \text{if } R_{ab} > R_a + R_b \\
R_a + R_b - R_{ab} & \text{if } R_{ab} \leq R_a + R_b
\end{cases}
\]

\(R_a\) and \(R_b\) are the Van der Waals radii of the atoms of the target and of the probe, and \(R_{ab}\) is the distance between the atoms.

A complex proposed by the SHAPES module is discarded if the “bumps” value exceeds a certain threshold. The threshold may vary for different docking tests: it should be set high when docking models of proteins that have been crystallized separately, because conformational changes may take place on binding. On the other hand, the threshold could be decreased when docking models derived from co-crystallized structures. The threshold was kept uniformly at 50 Å for all docking experiments (high stringency), except for those with the common model structures (110 Å, low stringency). In the analysis of the molecular collisions within models of co-crystallized complexes, one should expect in principle a low or very low bump value. In our procedure, however, we have always taken into account the intrinsic resolution of the SHAPES module (see below).

**CHARGES Module**

CHARGES provides an estimate of the “electrostatic complementarity” of the solution complexes produced by SHAPES.

Each atom of the two structural elements is assigned to one among six different atom types: positively charged (N\(_{\text{H}}\), N\(_{\text{H}}\)2), negatively charged (O\(_d\), O\(_e\)), hydrogen-bond donor (backbone N, N\(_e\)2), hydrogen-bond acceptor (backbone O, O\(_d\)1), hydrogen-bond donor or acceptor (O\(_g\), Ser, Thr, Tyr), and apolar (everything else).

The distance between each atom of the target structural element and each atom of the probe structural element is measured. A “charge value” is calculated for atom pairs whose distance is lower than a fixed cutoff (Table I).

\[
\text{CHARGES} = \sum_{a} \sum_{b} C_{ab}
\]

where

\[
C_{ab} = \begin{cases} 
0 & \text{if } R_{ab} > R_{\text{cutoff}} \\
A_{K_a K_b} & \text{if } R_{ab} \leq R_{\text{cutoff}}
\end{cases}
\]

and

\[
R_{\text{cutoff}} = \begin{cases} 
4.0 \text{ Å} & \text{if both target and probe atoms are apolar} \\
3.4 \text{ Å} & \text{otherwise}
\end{cases}
\]

\(R_{ab}\) is the distance between atoms a and b; A is a matrix containing the values reported in Table I.

Each complex proposed by the module SHAPES is then discarded if the charge value is below a certain threshold. As already discussed for BUMPS, CHARGES threshold can be varied for different docking experiments: it is set lower (–110, low

<table>
<thead>
<tr>
<th>Atom type</th>
<th>(+)</th>
<th>(–)</th>
<th>(Hb-d)</th>
<th>(Hb-a)</th>
<th>(Hb-da)</th>
<th>(0)</th>
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<td>–3</td>
<td>+3</td>
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<td>–3</td>
<td>+3</td>
<td>+3</td>
<td>–2</td>
</tr>
<tr>
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<td>–1</td>
<td>–2</td>
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<td>+2</td>
<td>–1</td>
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<td>–2</td>
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<tr>
<td>H-bond donor/acceptor (Hb-da)</td>
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<td>+2</td>
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<td>–1</td>
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<tr>
<td>Apolar (0)</td>
<td>+1</td>
<td>+1</td>
<td>+1</td>
<td>+1</td>
<td>+1</td>
<td>+1</td>
</tr>
</tbody>
</table>
stringency) when docking common models, and higher (-50, high stringency) when docking all other types of structures.

The distances considered and the values assigned to the different potential electrostatic interactions do not mean to mirror real atomic interactions: the values shown in Table I and the cutoff considered were determined empirically by running the CHARGES module on a small database of protein complexes (trypsin/trypsin inhibitor, 3ipi; antibody/lysozyme, 2hfi; triose-phosphate isomerase dimer, 7tim) and on approximately 40 three-dimensional models of mutagenized Rop proteins. They have been set to allow for the correct prediction (in 96% of the cases) of the mutant dimerization phenotype. Atom classification assigned in Table I is similar to others.5,13

Output of the ESCHER Procedure

ESCHER execution time varies with the size of the input structures: on a Silicon Graphics with an R8000 75MH processor, it may vary from a few minutes for two 30 amino acid long α-helices up to a few hours for large domains (more than 300 amino acids each).

The ESCHER procedure output is a list of solutions, each identified by the translations and rotations to be applied to the probe with respect to the target. In all of the docking experiments described, the number of listed solutions has been set to 500. This number has been increased to 5000 for the docking of unbound proteins, with a modest increase in calculation time.

Each solution is associated with the output scores of the three ESCHER modules described: a geometric complementarity score coming from SHAPES, a molecular collisions score from BUMPS, and an evaluation of electrostatic complementarity from CHARGES. The solutions with acceptable BUMPS and CHARGES values are retained and ranked according to their SHAPES score.

When SHAPES identifies an extended region of good complementarity, it usually generates a number of solutions in the same region. Therefore, all the solutions generated are grouped together on the basis of similarity; this is done by taking the solution with the highest geometric complementarity value and defining all other solutions having an rms deviation < 2.5 Å (5.0 Å for low-stringency conditions) from this structure as belonging to one cluster. The grouping continues with the best solution from those remaining and forms the next cluster by using the same procedure. Within each group, solutions are ranked according to their CHARGES value.

The resolution of the ESCHER procedure depends on two major factors: the size of the grouping matrix element (which affects the Tx, Ty, and Rz coordinates) and the stepwise exploration of the Z translation and of the X and Y rotations. The T and R coordinates identifying the solutions proposed should, therefore, be considered more properly as T ± ΔT and R ± ΔR. The expected error is different according to the coordinate considered: ΔTz = 0.75 Å; ΔRz = 5°; ΔRx varies from 0° to 6°; ΔTx and ΔTy vary from 0 to 1.5 Å.

In some cases, when docking secondary structure elements, a loose distance constraint was applied to discard solutions not compatible with the primary sequence of the analyzed protein. In the experiment with the two Rop helices, we defined a 14.5 Å distance threshold between the α-carbons of residues 28 and 32; this threshold corresponds to the distance between the α-carbon of the N-terminal and C-terminal residues of a 5-residue long peptide in an extended conformation.

Construction of the Common, Similar, and Backbone Structural Models

To simulate the experimental cases in which only information on the secondary structure is known, we used three types of structural models in which the amino acid side-chain or backbone atoms position are different from those found in the crystal structure: common, similar, and backbone.

Common model structures (Fig. 4a) were obtained from the crystallized element coordinates by substituting the conformation of each amino acid side chain with the most common rotamer in a database of amino acid rotamers24 (on average, 4.5 rotamers/amino acid). Common models can therefore be built on the backbone atom coordinates without any information on side-chain conformation. They display an average rms ~ 0.80 Å with respect to the original crystal structures.

Similar model structures (Fig. 4b) have been obtained by replacing each side chain with the most similar rotamer in the rotamer database. These models can be built only when the original crystal structure is known. The similar model structures display an average rms ~ 0.30 Å with respect to the corresponding crystal structure.

The backbone model structures (Fig. 4c) were built by substituting Rop helices backbone with geometrically regular α-helices (φ = -65°, ψ = -40°, Ω = 180°), by using the secondary structure option in the Biopolymer module of InsightII.25 Their rms deviation from the starting structures is ~ 0.55 Å. No energy refinement was applied to the model structures obtained.

Structural Elements Preparation

All of the target and probe structural elements docked by ESCHER in this work were obtained by splitting single-protein structures into two parts. The splitting procedure consisted of three steps. In the first step, each amino acid was defined as belonging to either the target or the probe structural elements. In the second, the protein main chain was
cut at the peptide bond between target and probe
amino acids, generating a number of peptide frag-
ments. In the last step, the target and probe struc-
tural elements were defined as the merged ensemble
of their fragments. When the protein had to be split
into two domains, the target and probe were defined
with automated criteria, when the protein had to
be split into secondary structure elements, the sepa-
ration was performed according to secondary struc-
ture assignment (see Table 4).

The smallest structural element was designed as
the probe. The starting orientation of the target has
always been chosen so that the interface was approxi-
mately parallel to the Z axis. In each docking exper-
iment, the two parts were positioned in the worst
possible relative orientation: the probe was rotated
5° around the X axis and 5° around the Y axis away
from the crystallographically defined orientation
(SHAPEs explores the rotation space around the X
and Y axis by 10° steps).

**Protein Structures Analyzed**

Twenty-three proteins of known structure were
used for our docking experiments from the Protein
Data Bank (PDB). For test runs with protein
domains, we used the following: porcine pancreatic
elastase (3est); papain (9pap); penicillopepsin (1ppn);
bovine liver rhodanese (1rhd); thremolysin (4tlm); bovine ribonuclease A (1rnd); yeast
guanylate kinase (1gky); bovine gamma a-crystallin
(2gcr); a cytochrome C2 (2c2c); a bacterial holo-D-
glyceraldehyde-3-phosphate dehydrogenase (1gd1); a
bacterial trypsin (1sgr); rat mast cell protease II
(3rp2); a bacterial dihydrofolate reductase (3drf);
alpha-lactalbumin (1alp); a protozoan calmodulin
(1dm); yeast phosphoglycerate kinase (3pgk); a bo-
vine prothrombin fragment (2pf2). These molecules
were selected for our tests because the definition of
their domains were derived by means of automated
procedures and were present in the literature.

For test runs with unbound components, we chose
one protease/inhibitor complex (cocrystallized struc-
ture: 1cho, corresponding structures crystallized
separately: 5cha/2ovo); and two antibody/antigen
complexes: 1) the antibody moiety of the 3hfz struc-
ture versus the lysozyme unbound structure 6lyz; 2)
the antibody moiety of the 2hfl structure versus 6lyz.

In these two last cases, solutions were compared
with the 3ahf and 2hfy molecules. For test runs
with secondary and supersecondary structure ele-
ments, we used the Rop protein (PDB code: 1rop); the
light chain of Hy-Hel 10 (3hfm); and lysozyme
(6lyz).

**RESULTS**

In an initial series of experiments (data not shown),
ESCHER has been successfully applied to the recon-
struction of protein complexes starting from the
crystal structure of their separated components:
trypsin/trypsin inhibitor (3tpi); antibody/lysozyme
(2hfl); Rop and triose-phosphate isomerase dimers
from their separated monomers (1rop and 7tim).

**Docking of Protein Domains**

We have applied ESCHER to the reconstruction of
two-domain proteins starting from their separated
domains. The proteins chosen represent a good data-
bases for the study of protein domain interaction:
some of them have an extended buried surface and a
large number of domain contacts, whereas others
show only a limited number of interdomain contacts
and a very small buried surface (Table I).

The target and probe domains were prepared as
described in Methods, starting from the deposited
PDB structure and following automatic criteria
for the target and probe definition.

In Table II, the results obtained with protein
domains are shown: for each domain pair, the num-

![Rop helix 1 (a1–a30) crystallographic structure is shown in bold. It is superposed to the common model structure (a), to the similar model structure (b), and to the backbone model (c).](image_url)
ber of proposed solutions is reported together with the ranking of the correct solution and its all-atom root-mean-square deviation (rmsd) from the crystallographic structure. In 9 of 18 docking runs, the correct solution was found and ranked first; in two cases, the correct solution ranked among the first few solutions; in the remaining seven cases, ESCHER was not able to find the correct solution within the first 500 solutions proposed by the SHAPES module. In all but one of these seven cases (the bacterial dihydrofolate reductase, 3dfr), the number of atomic contacts among the two domains is generally low. It is interesting to note that 1) the number of solutions proposed by the SHAPES module can be increased up to many thousands (if necessary), with a modest increase in calculation time and 2) that, on the other hand, the number of solutions that escape the BUMPS and CHARGES filters is generally very low: in case of a really blind test, this would allow further investigation of few solutions by more sophisticated methods.

There is no evident correlation between ESCHER efficiency and buried surface area in the domain-domain interface. A strong correlation could be detected between ESCHER efficiency and a value \( \phi \), defined as the product of the ratios between the buried and the whole surface of each domain:

\[
\phi = \frac{b_1 \cdot b_2}{t_{\text{tot}_1} \cdot t_{\text{tot}_2}}
\]

where \( b_1 \) and \( b_2 \) are the two domain buried surfaces and \( t_{\text{tot}_1} \) and \( t_{\text{tot}_2} \) are the two domains total surfaces. This value has no physical significance but can be used to predict ESCHER ability in rebuilding a complex of known structure. In the test cases that we analyzed, when \( \phi > 19 \times 10^{-3} \), ESCHER ranked the correct solution among the first few. For lower \( \phi \) values, the correct solution was not found among the first 500 solutions proposed by SHAPES.

**Docking of Protein Complexes Starting From Unbound Components**

We chose three cases: a protease-protease inhibitor and two antibody-antigen complexes. The results of the docking procedure were evaluated by superimposing the docked (originally unbound) molecules onto their crystal structures in the PDB complexes. These complexes (Table III, second column) were used as reference systems to calculate the rmsd from the Ca of the proposed solutions.

The protease-protease inhibitor complex was run by using the entire protein surfaces, and the procedure proposed only three solutions. A solution with an rmsd of 1.9 Å with respect to the defined reference system scored third.

In the immunoglobulin-lysozyme docking experiment, the entire antigen was docked onto the complementary determining region (CDR) of the immunoglobulin. The 3hfm experiment ended with only two

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**TABLE II. Protein Domains Results**

<table>
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<th>PDB code</th>
<th>Target</th>
<th>Probe</th>
<th>Buried surface (Å²)</th>
<th>( \phi \times 10^{-3} )</th>
<th>Rank (rmsd) of the first correct solution ( ^* )</th>
<th>Rank (rms) of the best solution</th>
<th>Number of solutions proposed</th>
</tr>
</thead>
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<td>263</td>
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<td>1 (1.8)</td>
<td>31</td>
</tr>
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<td>1gd1</td>
<td>1–148</td>
<td>149–332</td>
<td>2186</td>
<td>22.0</td>
<td>1 (2.0)</td>
<td>2 (1.5)</td>
<td>11</td>
</tr>
<tr>
<td>9pap</td>
<td>1–16, 113–208</td>
<td>17–112, 209–212</td>
<td>1387</td>
<td>23.7</td>
<td>2 (2.0)</td>
<td>2 (2.0)</td>
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<tr>
<td>1ppn</td>
<td>1–18, 111–212</td>
<td>19–110</td>
<td>1567</td>
<td>28.0</td>
<td>2 (2.1)</td>
<td>2 (1.5)</td>
<td>4</td>
</tr>
<tr>
<td>1sgt</td>
<td>16–28, 69–80, 103–124</td>
<td>29–68, 81–120</td>
<td>1583</td>
<td>31.2</td>
<td>1 (1.5)</td>
<td>1 (1.5)</td>
<td>9</td>
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<tr>
<td>3est</td>
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<td>16–29, 121–233</td>
<td>2186</td>
<td>38.9</td>
<td>1 (1.7)</td>
<td>1 (1.7)</td>
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<td>1–27, 123–216, 221–230</td>
<td>28–122, 231–243</td>
<td>2382</td>
<td>47.6</td>
<td>1 (1.6)</td>
<td>3 (1.4)</td>
<td>5</td>
</tr>
</tbody>
</table>

*All atoms rms is calculated by superposing the target domain of the docking solution complex onto the target domain of the protein crystal structure.
†Rank and rmsd of the first solution having an rmsd below 2.5 Å.
‡Identifies solutions not presenting the best CHARGES value of their cluster (see Methods).
TABLE III. Unbound Proteins Results

<table>
<thead>
<tr>
<th>Reference</th>
<th>Rank (rms) of the first correct solution</th>
<th>Rank (rms) of the best solution</th>
<th>Number of solutions</th>
</tr>
</thead>
<tbody>
<tr>
<td>2ova5cha</td>
<td>1cho 3 (1.9)</td>
<td>3 (1.9)</td>
<td>3</td>
</tr>
<tr>
<td>2hfl/6lyz</td>
<td>2hfl</td>
<td>—</td>
<td>0</td>
</tr>
<tr>
<td>3hfm/6lyz</td>
<td>3hfm 2 (1.0)</td>
<td>2 (1.0)</td>
<td>2</td>
</tr>
<tr>
<td>2hfl/6lyz</td>
<td>2hfl 11 (1.12)</td>
<td>11 (1.12)</td>
<td>11</td>
</tr>
<tr>
<td>3hfm/6lyz</td>
<td>3hfm 1 (1.0)</td>
<td>1 (1.0)</td>
<td>1</td>
</tr>
</tbody>
</table>

*The rmsd of the unbound molecules with their corresponding reference molecules is obtained after superposing the Ca atoms of the unbound ligand (receptor) on those of the complexed reference ligand (receptor).

†Antibody CDRs versus entire lysozyme molecule. We selected a subset of the antibody solvent accessible surface, whose distance from the antigen is less than 6 Å in the cocrystallized structure; regions of geometric complementarity were evaluated if sharing at least 1 Å² with the defined subset.

‡Antibody CDRs versus the lysozyme antibody binding site. We selected a subset of the lysozyme solvent accessible surface, whose distance from the antibody is less than 6 Å in the cocrystallized structure; complementarity has been evaluated for regions sharing at least 1 Å² with this defined subset. In this case, approximately half of the antigen molecule has been considered in the docking test.

§Rank and rmsd of the first solution having an rmsd below 2.5 Å.

The described results show that generally a low rms can be obtained in a long list of possible solutions and that the choice of selecting only a few solutions, including the correct one, is usually paid in terms of higher rmsd. Both procedures, the geometry-based suite and PUZZLE, generally require four to five times less cpu time than ESCHER.

Docking of Secondary and Supersecondary Structural Elements

The experiments described in the previous section show that 1) ESCHER can reconstruct whole proteins starting from their separated crystal domains when the interdomain contact area is sufficiently large with respect to the total surface area and 2) ESCHER is able to dock unbound protein components whose structures display differences with respect to their conformation in the crystallographic complex.

Next, we explored the feasibility of docking secondary and supersecondary structure elements, where the ϕ value is generally higher than 19 × 10⁻³ (Table IV).

We chose three docking cases, in which the docking partners were as follows: 1) a secondary structure element separated from its structural environment and the remaining part of the protein. We selected alpha helices extracted either from Rop, a four-helix bundle protein (1α/3α), or from the alpha and beta protein lysozyme (1α/αβ); 2) two supersecondary structures forming a protein domain: the beta sheets components of an immunoglobulin variable domain (1β/1β); and 3) two secondary structure elements: the two helices forming the Rop monomer (1α/1α).

We performed docking experiments starting either from the crystal structures or from approximate molecular models. Three different types of models (Fig. 4) were utilized as described in Methods: 1) similar models are derived from crystal structures by substituting each amino acid rotamer with the most similar rotamer from a small database of amino acid rotamers. Their rmsd from the crystal structures is approximately 0.30 Å. 2) In backbone models, backbone dihedral angles are approximated to
When the crystallographic structures are used, the correct solution always scores first. When using low-stringency conditions (Table VI), ESCHER produces a low rms solution in the first ranking position in the 1α/αβ and 1β/ββ test cases. In the other two cases tested, no correct solution was obtained. When using low-stringency conditions (Table VI), the correct solution scores among the first ones in three of the four tested cases. In low-stringency conditions, a solution is considered correct when its backbone rmsd from the crystal position is lower than 5 Å.

Analysis of the solutions obtained with low-stringency conditions reveals that, in the 1α/αα and in the 1β/ββ cases, common model structures display a substantially lower degree of electrostatic complementarity compared with the corresponding crystal structure; in the 1α/αβ case, both molecular clashes and lowering of electrostatic complementarity contribute to the lack of low rms solutions in the proposed solutions. We report the results obtained under both conditions for discussion.

Finally, a control test was performed in which a 30 amino acid long Rop helix is docked onto the lysozyme domain 2, which normally shelters a 15 amino acid helix. ESCHER geometric module predicted the formation of a Rop/lysozyme “mosaic” protein, with a good geometric complementarity. However, the electrostatic module gave a very poor score of the electrostatic complementarity between the two protein portions and discarded it.

In all the docking runs described, the target and the probe were treated as if they were independent molecules; nevertheless, distance constraints could be applied, thus discarding physically impossible solutions. In the 1α/αβ and 1α/αα cases, when using similar or common model structures, a number of proposed solutions was antiparallel to the correct ones. They were, therefore, discarded because amino acids close in the primary sequence were too far in the proposed solution.

**DISCUSSION**

ESCHER is a new rigid body docking algorithm that was developed after PUZZLE. ESCHER differs from PUZZLE in the simplified description of the protein surface as well as in the matching and scoring criteria. Both procedures rely on a geometric
approach, but ESCHER has been endowed with a molecular clashes (BUMPS) and an electrostatic (CHARGES) module, which are used as a selective filter on the geometrically favorable solutions. The CHARGES electrostatic module allows a lower resolution definition of the protein surfaces and less stringent criteria of geometric complementarity. Typically, when docking unbound components, thousands of solutions are proposed by the geometric module, but only a few, including the correct one, survive screening with the electrostatic module. This illustrates the importance of adding this module to the procedure.

The application of ESCHER to a database of two-domain proteins indicates that its performance correlates with the ratio between the buried surface area upon domain-domain complex formation and the total surface \( \phi \). When starting from the crystallographic coordinates of the target and probe domains with \( \phi \) greater than \( 10 \times 10^{-3} \) (see Table I), the correct docking solution always scores among the first few.

ESCHER has been applied to the unbound components of a small database of protein-protein complexes. The low resolution description of the protein surface permits recognition of correct protein interfaces even when side-chain movements upon binding substantially alter the geometric and electrostatic complementarity of the interface.

In comparison with other published docking procedures, \(^\text{8,9,10,34}\) ESCHER generally requires longer computational times but can propose low rms solutions among a very restricted number of possible choices. This allows one to rank the small number of possible solutions by applying more sophisticated computational methods.

We have also explored the feasibility of applying ESCHER to the reassociation of secondary and supersecondary structure elements within a protein structure. We simulated a helix-helix and a helix-three helices interaction within a four-helix bundle (Rop); a helix and the remaining part of the protein in an alpha-beta protein (lysozyme) and two beta-sheets within a beta protein (an immunoglobulin variable domain).

Initially, these docking experiments were performed with secondary structure elements extracted from the crystallographic structure, without altering the conformation of the side chains and maintaining a rigid alpha-carbon backbone. In this ideal situation, in which the crystallographically determined complementarity is strictly maintained, ESCHER proved to be very effective. In a realistic docking experiment, however, the conformation of the side chains in the complex is not known. Thus, to explore the versatility of ESCHER and to simulate the assembly of a three-dimensional model from the definition of its secondary structure, we applied the ESCHER procedure to the docking of secondary structure elements after changing the conformation either of the amino acid side chains or of the backbone. The results obtained with the similar and backbone models are very satisfactory. With common models (Table V), the results were satisfactory in half of the cases tested: changing the rotamer conformations to the most common ones found in the database considerably lowered the geometric complementarity of target and probe elements. It was, therefore, necessary to lower the stringency of the BUMPS and CHARGES selective filters (Table VI) to achieve acceptable results in three of the four docking experiments.

The performance obtained in the common 1\(\alpha/1\alpha\) case deserves more discussion: this approach does not seem to be applicable to elements, such as single alpha helices or beta strands. Work is in progress to see whether, in these difficult cases, our procedure could be supported and complemented by different methods, such as the use of backbone-dependent rotamer libraries\(^\text{35,36}\) or the identification of correlated mutations.\(^\text{37}\)

The relatively low resolution of the best solutions in the experiments performed with common models, however, are more appropriately to be compared with resolutions typical of methods that predict structure from sequence rather than with docking methods. This confirms that, especially when considering small docking elements, the side-chain conformation is not irrelevant to the definition of the geometric and electrostatic complementarity. The success of the experiments with the similar models, however, reassures that, by exploring combinatorially a small library of side-chains conformations, the probability of predicting the "correct solution" is very high.

Other applications for the ESCHER procedure can be envisaged. The structure of a growing number of large proteins is approached by experimentally determining the conformation of their separate domains. We propose ESCHER to represent a useful tool to assist in the task of assembling the complete protein starting from the crystallographic structures of its domains.

It is often found that the structure of newly discovered proteins can be reliably predicted by homology modeling. Alignment of the two primary sequences, however, may reveal that the protein of unknown conformation contains peptide insertions with respect to the protein in the database. Whenever the secondary structure of the inserted peptide can be predicted, ESCHER can help in docking the orphan peptide onto the "core" structure determined by homology modeling.

ESCHER has already been exploited as a tool to predict the formation of protein-protein complexes. In one application, starting from the coordinates of the crystallized proteins, we have predicted the conformation of the complex between the TP7 antibody and its cognate antigen, Thermus aquaticus DNA polymerase.\(^\text{38}\)
Furthermore, ESCHER has been used as a tool to design and evaluate Myc mutants that can homodimerize (manuscript in preparation). Work is in progress to address the possibility of speeding up the procedure on parallel (or even massively parallel) machines.

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REFERENCES