Inferential Optimization for Simultaneous Fitting of Multiple Components into a CryoEM Map of Their Assembly

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Models of macromolecular assemblies are essential for a mechanistic description of cellular processes. Such models are increasingly obtained by fitting atomic-resolution structures of components into a density map of the whole assembly. Yet, current density-fitting techniques are frequently insufficient for an unambiguous determination of the positions and orientations of all components. Here, we describe MultiFit, a method used for simultaneously fitting atomic structures of components into their assembly density map at resolutions as low as 25 Å. The component positions and orientations are optimized with respect to a scoring function that includes the quality-of-fit of components in the map, the protrusion of components from the map envelope, and the shape complementarity between pairs of components. The scoring function is optimized by our exact inference optimizer DOMINO (Discrete Optimization of Multiple Interacting Objects) that efficiently finds the global minimum in a discrete sampling space. MultiFit was benchmarked on seven assemblies of known structure, consisting of up to seven proteins each. The input atomic structures of the components were obtained from the Protein Data Bank, as well as by comparative modeling based on a 16–99% sequence identity to a template structure. A near-native configuration was usually found as the top-scoring model. Therefore, MultiFit can provide initial configurations for further refinement of many multicomponent assembly structures described by electron microscopy.

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Introduction

Structural description of macromolecular assemblies is essential for a mechanistic understanding of the cell. The scope of the problem is revealed by protein interaction studies: The yeast cell contains approximately 800 distinct core complexes of 4.9 proteins, on average, most of which have not yet been structurally characterized. The human proteome is likely to have an order of magnitude more distinct assemblies than the yeast cell. Therefore, there are thousands of biologically relevant assemblies whose structures still need to be determined.

Structural determination of macromolecular assemblies is a major challenge in structural biology. X-ray crystallography can provide structures of stable assemblies at atomic resolution. However, there are many other assemblies that are refractory to crystallographic determination. A low-resolution structure of these assemblies can be determined by
cryo-electron microscopy (cryoEM). The resolution usually ranges from 4 Å, where the backbone of the protein can be traced, to 30 Å, where only the outer envelope of the assembly is visible. The increasing numbers of atomic and cryoEM data sets have stimulated the development of computational techniques for fitting atomic structures of assembly components into a cryoEM density map of the whole assembly. The result is a pseudo-atomic model of the assembly that can reveal significant insights into its structure, dynamics, function, and evolution.

Here, we focus on determining the positions and orientations (i.e., placements) of multiple atomic component models within the assembly density. When the structure of a homologous assembly (template) is available, the placements of the components can be computed by fitting the template into the target assembly density, superposing the target component models on the corresponding template components, and refining the model. Alternatively, the component positions can be determined experimentally by protein-labeling methods relying, for example, on gold-labeled antibodies. However, when only a cryoEM map and component structures are available, a general method for solving the configuration problem is not yet available.

A sequential method for fitting multiple components into an assembly map has been described. The method starts by fitting of the largest component into the map, followed by iterative fitting of the largest remaining component into the unoccupied density until all components have been fitted. The fitting of a component into a given map can be performed manually using interactive visualization tools. More desirably, automated fitting methods that assess the placement of a component by a fit between the component and a segmented or complete density of the assembly can also be used; the fit is optimized over the translational and rotational degrees of freedom of a rigid component relative to the map. The sequential method is applicable if the remaining components can be unambiguously fitted into the unoccupied densities. Unfortunately, this condition is generally not satisfied, especially when the resolution is low, the number of components is large, and component models are inaccurate. For example, sequential fitting is not expected to work for the 19S proteasome with 20 component proteins, for the mammalian ribosome for which 30 of 80 proteins are not present in known archaeal or bacterial ribosomes, or for the ryanodine receptor isomorf 1 for which some domains are poorly modeled while no template is available for others.

Here, we describe a method named MultiFit for determining the configuration of multiple high-resolution component structures based on the quality-of-fit of each component into the density map, the protrusion of each component from the map envelope, and the shape complementarity between pairs of components. The combination of these terms reduces the ambiguity of the final solution, compared to using any individual term on its own. The task of sampling the configuration space is challenging because the placement of a component depends on the placement of other components. MultiFit tackles this combinatorial challenge by reformulating the problem as an inferential optimization over a discrete sampling space. In outline, a discrete set of possible placements for each component is first generated independently of other components. Next, the globally optimal combination of placements with respect to a scoring function is found by a combination of branch-and-bound search and the DOMINO (Discrete Optimization of Multiple Interacting Objects) inferential optimizer. The relative translations and orientations of pairs of components in the best-ranking configurations are then refined; specifically, a refined discrete sampling space is generated by pairwise geometrical docking between interacting components, and the optimal refined combination of placements is again found using DOMINO. We successfully validated the method on a simulated benchmark of six assemblies consisting of up to seven proteins each. In addition, for a more realistic test, we determined the configuration of four domains in the subunit of GroES–ADP7–GroEL–ATP7 chaperonin from Escherichia coli based on an experimentally determined map at a resolution of 23.5 Å. A near-native configuration scored best in four test cases, third best in two cases, and fourth best in the remaining case.

Below, we begin with a detailed description of general combinatorial optimization by DOMINO, followed by a formal definition of the component configuration problem and the MultiFit algorithm for a solution using DOMINO (Theory). We then demonstrate the performance of MultiFit on benchmark cases (Results). Finally, we discuss the implications of MultiFit and DOMINO for the structural characterization of large assemblies (Discussion).

Theory

Combinatorial optimization by DOMINO

DOMINO applies a divide-and-conquer approach to efficiently find solutions with the globally optimal score within a discrete sampling space (Fig. 1). The idea is to decompose the set of variables into relatively uncoupled but potentially overlapping subsets that can be sampled independently of each other, and then to efficiently gather the subset solutions into the global minimum. The strength of this approach derives from the decomposition procedure that helps reduce the size of the search space from exponential in the number of components in the whole system to exponential in the number of components in the largest subset. Next, we describe DOMINO’s application to the minimization of a scoring function $F$ corresponding to a sum of single-body terms $\{\alpha_i\}$ and pairwise terms $\{\beta_{ij}\}$:

$$F(y_1, \ldots, y_n) = \sum_i \alpha_i(y_i) + \sum_{i<j} \beta_{ij}(y_i, y_j)$$
where \(y_i\) are the variables being optimized (e.g., in MultiFit, these variables are the positions and orientations of the components). The scoring function \(F\) is represented by a graphical model \(G = (V,E)\). The graphical model \(G\) of the scoring function \(F\) is a graph whose nodes \(V\) correspond to the variables \(\{y_i\}\), and edges \(E\) connect pairs of nodes. The weight of a node corresponding to \(y_i\) is \(\alpha_i\), and the weight of an edge between nodes corresponding to \(y_i\) and \(y_j\) is \(\beta_{ij}\). Thus, the scoring function \(F\) is the sum of all node and edge weights.

The problem of finding the minimum of the scoring function \(F\) is equivalent to the maximum a posteriori problem in a graphical model. This problem is known to be NP-hard (nondeterministic polynomial—time hard) for an arbitrary graph \(G\). NP-hard is a class of decision, search, and optimization problems whose computing time increases at least exponentially with the number of optimized variables. When a graphical model has, at most, one path between any two given nodes (i.e., it does not contain cycles and thus is a singly connected...
graphical model or a tree), it can be efficiently optimized by the belief-propagation algorithm. Unfortunately, the belief-propagation algorithm is not guaranteed to converge into the globally optimal solution for graphs with cycles, such as the graphical models used for the MultiFit application. Therefore, to ensure that we find the global minimum of $G$ efficiently, we apply a divide-and-conquer approach. First, the variables to be optimized are decomposed into smaller relatively uncoupled but potentially overlapping subsets, using a junction tree construction algorithm (the decomposition step). Second, a discrete sampling space is generated for each variable (the variable sampling step; e.g., by uniform sampling). Third, the discrete states of the individual subsets are constructed and gathered into the globally optimal solution, using the belief-propagation algorithm (the gathering step). Graph theory provides

Fig. 2 (legend on next page)
efficient algorithms for decomposition (i.e., junction tree construction) and gathering (i.e., belief propagation). Next, we elaborate on each of the three steps.

In the decomposition step, the graphical model $G$ is converted into a tree $T$ whose nodes $U$ are potentially overlapping subsets of variables $y_i$ (Fig. 1). Importantly, for any two nonadjacent subsets in $T$ that share some variables, the subsets that connect them must also contain these variables (i.e., $T$ is a junction tree). For any junction tree, it is possible to gather the discrete states of individual subsets into the globally optimal solution using the belief-propagation algorithm. For maximum efficiency, we aim to decomposing the graphical model into the junction tree such that the size of the largest subset is minimal, which is an NP-hard problem. We use the minimum-degree method that was shown empirically to result in smallest subsets for sparse graphical models.28

In the variable sampling step, a discrete set of values for each variable is created. The details of this discretization may depend on the scoring function $F$. Most generally, uniform sampling over a relevant range of values can be used. A potentially better possibility is to use the union of the local minima of scoring functions spanned by the variables in the subsets containing the discretized variable.

In the final step (gathering step), the HUGIN version of the belief-propagation algorithm29 is applied to the junction tree $T$ to find the global minimum of $F$ (Fig. 1). The computational complexity of the HUGIN algorithm is $O(|U|L^2)$, where $s$ is the size of the largest subset of $T$, and $L$ is the number of values of a variable $y_i$. The belief-propagation algorithm is based on passing messages between the nodes (i.e., subsets of variables) of the junction tree. A subset is allowed to send a message to a neighbor subset if it has received messages from all of its remaining neighbor subsets.

Thus, propagation of messages is initiated in subsets connected only to a single subset (i.e., the leaf subsets) and proceeds to the neighboring subsets until some subset receives messages from all of its neighbors (i.e., the root subset). The content of a message to a target subset is a vector of the minimal values of the partial scoring function $F$ over the variables in all previously visited subsets and the target subset, for each combination of values of the remaining variables in the target subset; a partial scoring function over a subset of variables includes only those terms of $F$ that involve these variables. Messages from the root subset are then sent back to the other subsets, completing the message-passing process when the leave subsets receive back the messages from the root subset. For messages from the root subset, the partial scoring function is the scoring function $F$ (because all subsets have been already visited), and thus each subset that received a message from all other subsets can infer the values of its variables in the global minimum. The efficiency of message passing derives from enumerating combinations of values for only those variables that are shared between different subsets.

**MultiFit: Simultaneous fitting of multiple components into a density map of their assembly**

The goal is to find the positions and orientations (i.e., placements) of components (e.g., subcomplexes, proteins, domains, and secondary structure segments), represented at atomic resolution, within a cryoEM density map of their assembly. We express this structure characterization challenge as a combinatorial optimization problem. Next, we outline a representation of the modeled system, a scoring function, and an optimization algorithm.

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**Fig. 2.** MultiFit outline. The MultiFit algorithm is illustrated using an assembly between models of Rpb1 (red), Rpb2 (yellow), ARPC1 (light green), ARPC2 (blue), ARPC3 (gray), ARPC4 (dark green), and ARPC5 (purple) (PDB entry 1tyq). The component–template sequence identities and Cα RMSDs are indicated. The input to MultiFit is the assembly density map (gray mesh) and the atomic structures of the individual components (top left). The output is a ranked list of assembly models that optimize the MultiFit scoring function (one model is shown on the bottom right). (1) The anchor points (the seven labeled nodes) are constructed for the input density map by k-cluster; the nine gray edges indicate pairs of anchor points that are sufficiently close to allow components placed in their vicinity to interact with each other. (2) The sampling space of component placements is discretized by fitting each of the seven components around each of the seven anchor points (regions) and by selecting a number of top-ranking placements for each component in each region; the small colored spheres indicate placement centroids. (3) The optimal combination of component placements is found by optimizing the scoring function $S$ for each mapping of components to anchor points using DOMINO. (3.1) For efficiency, we replace the enumeration by a branch-and-bound procedure that eliminates some of the mappings and makes use of partial results. In the branch stage, we first decompose the anchor graph into an anchor junction tree using DOMINO’s decomposition algorithm (Fig. 1). The top 60% mappings of components to anchor points for each subset of anchor points (partial mappings) are found and stored by iterating over all possible partial mappings; the color of the circle indicates which component is mapped to the anchor point. The partial mappings are scored by partial scoring function $S$, including only the terms involving the mapped components. Complete mappings consistent with the stored partial mappings are generated efficiently with a hashing procedure (not described). (3.2) Next, for each of these complete mappings, the optimal combination of placements for the seven components is found by DOMINO; the color of the solid circles in the component junction tree indicates the component mapped to the corresponding anchor point in the anchor junction tree. The molecular model shown has a mapping score of 0 and a rank of 10th. (4) The 20 top-scoring coarse models are further refined. A refined sampling space is generated for each coarse configuration by docking pairs of its interacting components and by selecting only those placements that are approximately consistent with the initial coarse configuration. (5) DOMINO is applied again to find the optimal combination of placements for the seven components; the molecular model shown has a mapping score of 0 and a rank of 4th.
**Representation**

The assembly density map is represented by a three-dimensional grid in which every voxel is assigned an estimated density value. The components are represented by their atoms and remain rigid throughout the entire optimization process (Fig. 2).

**Scoring**

We evaluate potential configurations based on the quality-of-fit of individual components into the density map, the protrusion of each component from the map envelope, and the shape complementarity between pairs of components.

**Optimization**

The component configuration that optimizes the scoring function is identified by a combinatorial optimization protocol consisting of three stages: (i) anchor graph construction, (ii) coarse-grained sampling, and (iii) fine-grained sampling (Fig. 2). In anchor graph construction, the density map is discretized into regions, and the connectivity between them is calculated. In coarse-grained sampling, the sampling space is first discretized by fitting each of the components into each of the map regions and selecting a number of top-ranking placements for each component in each region. Next, a branch-and-bound search through all mappings of components to regions combined with DOMINO finds 20 top-scoring configurations. In fine-grained sampling, each of these top configurations is refined by DOMINO; a refined sampling space is generated for each coarse configuration by docking pairs of its interacting components and selecting only those placements that are approximately consistent with the initial coarse configuration.

**Scoring function for MultiFit**

The score of placements of $N$ components in an assembly density map is:

$$S(x_1, \ldots, x_N) = \sum_i \{ \varphi_1(x_i) + \varphi_2(x_i) \} + \sum_{i \neq j} \varphi_3(x_i, x_j)$$

$\varphi_1(x_i)$ is the quality-of-fit of $x_i$ into the assembly density map $D$. In the extreme case, the configuration that optimizes $\sum_i \varphi_1(x_i)$ may occupy only the highest-density region in the assembly density map. To overcome this problem, we add two geometric terms ($\varphi_2$ and $\varphi_3$) to the scoring function. The component protrusion term $\varphi_2(x_i)$ scores how well $x_i$ is placed inside the density envelope. The interaction term $\varphi_3(x_i, x_j)$ scores the pairwise shape complementarity between the structures $x_i$ and $x_j$.

**Quality-of-fit term**

The fit of a given structure $x_i$ into the assembly density map $D$ is usually assessed by a cross-correlation measure between the densities of $x_i$ and the assembly. Here, we use the “normalized fitting score” $C$ as implemented in Mod-EM (Eq. 2 in Topf et al.); the density of $x_i$ is simulated at the same resolution as the assembly density map $D$, using the uniform-sphere model. However, $C$ is insufficient for comparing placements of different components because small domains have a better chance of higher cross-correlation with the map. Thus, we calculate the quality-of-fit of a component into a map by expressing $C$ as a Z-score $(C_m)/s$, where $m$ and $s$ are, respectively, the mean and standard deviation of a reference distribution of $C$. The reference distribution is generated by optimally fitting randomly selected, similarly sized protein structures into simulated maps of randomly selected, similarly sized protein structures (F. Davis, M. S. Madhusudan, N. Eswar, A. Sali, and M. Topf, unpublished results).

**Interaction term**

The pairwise shape complementarity score between main-chain atoms of the structures $x_i$ and $x_j$ is calculated as the weighted sum of a reward for interaction surfaces and a penalty for steric clashes between the components. Specifically, the reward is the total number of surface atom pairs of $x_i$ and $x_j$ within a distance cutoff, and the penalty is a weighted sum of all clashing pairs of atoms of $x_i$ and $x_j$. To speed the calculation of the reward, we first classify atoms as buried or exposed by placing each component structure on a grid and dividing the grid into a surface and four core shells according to the closest distance from the molecular surface (the surface shell contains all grid points that are, at most, half of the map resolution away from the surface). The reward is calculated by indexing the surface atoms of $x_i$ in a geometric hash table, querying the hash table for each surface atom of $x_j$, and summing the number of hits to get the reward. To calculate the steric clash penalty, we determine the accessibility of each atom of $x_i$ (and $x_j$) using the grid of $x_i (x_j)$. If an atom in $x_i (x_j)$ is located within the surface ($k=0$) or the $k$th core shell of $x_i (x_j)$, we add $(k+1)\times 27$ to the penalty. The sum of the penalty score of $x_i$ with respect to $x_j$ and the penalty score of $x_j$ with respect to $x_i$ is divided by 2 to obtain the steric clash penalty. Due to fitting and modeling errors, the correct configuration of components might include some minor clashes between interacting components. These clashes are not significantly penalized because of the thickness of the surface shell and because of the evaluation of the favorable and penalty terms using only main-chain atoms. The choice of shell thickness and weight of the penalty score was chosen by trial and error.

**Component protrusion**

The protrusion of a component from the assembly envelope is defined to be the negative value of the shape complementarity score between the component surface and the assembly envelope. The assembly envelope is calculated by representing each density
voxel above a threshold as an atom and calculating the Connolly surface\(^{37}\) of this collection of atoms.

**Optimization for MultiFit**

**Construction of anchor graph**

The centroids of \(L\) approximately equally sized regions of density voxels are calculated from the density map \(D\) using a \(k\)-means clustering\(^{38}\) (\(k\)-cluster) that is similar to the QVOL procedure of Situs;\(^{39}\) a density voxel belongs to the region with the closest centroid. When \(L\) equals \(N\), and the components are of similar sizes, the centroids of the regions correspond approximately to the centroids of the \(N\) assembly components. These points are the nodes of the anchor graph. We then calculate the connectivity between the anchor points (i.e., the edges of the anchor graph); a pair of anchor points (\(a_i, a_j\)) is connected if their regions are in contact.

**Discretization step in coarse-grained sampling**

We construct a discrete sampling space of component placements, represented by a set of \(M\) placements (by default, 50) for each of the \(N\) components in each of the \(L\) regions. Thus, each set of placements for all components in region \(i\) \((A_i)\) contains \(M \times M'N\) “local” placements around its anchor point \(a_i\). Here, we set \(L\) to \(N\), although \(L\) can also be larger than \(N\).

In detail, for each component \(j\), the discrete sampling space is constructed as follows. Placements around each anchor point \(a_i\) are sampled by optimizing the normalized fitting score \(C\) in a cube surrounding the anchor point (the edge length of the cube is half the resolution of the map). This optimization is performed by Mod-EM,\(^{31}\) starting with a random starting orientation of the component centered at the anchor point. Next, the sampled placements for all anchor points are clustered based on their pairwise \(C^o\) RMSD values: The highest scored placement (by \(C\)) initiates the first cluster and is its pivot. The closest remaining placement either is joined with the first cluster for which its \(C^o\) RMSD with the cluster’s pivot is less than the threshold (half the resolution of the map) or initiates its own cluster otherwise. The process is repeated with the best-scoring nonclustered placement until all placements have been clustered. The best-scoring placement from each cluster is assigned to the set of placements \(A_j\) corresponding to the closest anchor point \(a_j\); each anchor point is assigned, at most, \(M'\) placements.

**Optimization step in coarse-grained sampling**

We find the optimal combination of placements of components by optimizing the scoring function \(S\) within the discrete sampling space constructed in the previous step. The global minimum of \(S\) is the minimum of the optimal solutions for each of the \(L! / N!\) mappings of components to anchor points \(\{\pi_k\}\), where \(\pi_k\) is a function that maps a component \(j\) to an anchor point \(i = \pi_k(j)\); formally, we solve

\[
\min_{x=1}^{\infty} \min_{\{y_1, \ldots, y_N\}} S(x_1, \ldots, x_N),
\]

where \(x_i\) are placements of component \(j\) in the set \(A_{\pi_k(j)}\) as constrained by mapping \(\pi_k\).

Naively, this optimization could be achieved by a nested double loop in which the outer loop consists of enumerating the mappings and the inner loop consists of applying DOMINO to the scoring function \(S\) constrained by the given mapping. However, enumerating over all possible mappings becomes computationally expensive as the number of components increases. To improve the efficiency of MultiFit, we replace the enumeration by a branch-and-bound procedure that eliminates some of the mappings and makes use of partial results (Fig. 2).

The scoring function \(F\) optimized by DOMINO for each mapping \(\min_{x=1}^{\infty} \min_{\{y_1, \ldots, y_N\}} S(x_1, \ldots, x_N)\) is a simplified \(S\) that does not contain uninformative interaction terms \(\varphi_3\) corresponding to physically noninteracting components (Fig. 2); specifically, we eliminate interaction terms between pairs of components that are mapped to unconnected anchor points. Importantly, it is this simplification that results in a relatively “sparse” graphical model \(G\), thus allowing it to be optimized efficiently by DOMINO.

**Discretization step in fine-grained sampling**

We construct a refined discrete sampling space for a coarse configuration found in coarse-grained sampling \((x_1^0, \ldots, x_N^0)\). The refined set of placements of component \(j\) is first initialized with the placements in \(A_{\pi_k(j)}\) as found in coarse-grained sampling. We then enrich this set of placements by sampling the binding of component \(j\) to neighboring components \([w]\) with PATCHDOCK.\(^{33}\) A PATCHDOCK-produced binding mode of component \(j\) to component \(w(x_j)\) is added to the refined set of placements of component \(j\) if (i) the distance between the centroid of \(x_i\) and the centroid of \(x_j^0\) is below half of the resolution of the map and (ii) \(x_j\) is consistent with the density map boundaries [i.e., if \(\varphi_2(x_j)\) is below a predefined threshold]. Finally, the refined set of placements of component \(j\) is reranked by the quality-of-fit score and clustered according to \(C^o\) RMSD (as described above).

**Optimization step in fine-grained sampling**

The optimal combination of component placements is found by DOMINO by optimizing the scoring function \(S\) within the refined discrete sampling space.

**Results**

**Benchmark with simulated maps**

**Benchmark**

We tested MultiFit on a benchmark of six simulated test cases. The assembly density maps
were simulated at 20 Å resolution using the PDB2VOL program of Situs, as well as models calculated by comparative modeling using MODELLER-9v3† based on related template structures with sequence identity ranging from 16% to 99%. The accuracy of the individual comparative models is quantified using Cα RMSD and native overlap to the corresponding native structure. Native overlap (NO3.5) measures the percentage of Cα atoms of the model that are within 3.5 Å from the corresponding Cα atoms in the native structure. The native overlap was calculated by superposing the model on the corresponding native structure using rigid-body least-squares minimization, as implemented in the model.superpose command of MODELLER-9v3.

We use three scores to quantify the accuracy of modeled configurations at different levels of resolution. First, the mapping score is the number of substitutions needed to convert the assessed mapping of components to anchor points into the native mapping of components to anchor points (the Hamming distance); the native mapping has a mapping score of 0. Second, the configuration score is the fraction of the components positioned correctly; we define a component as positioned correctly if the distance between its centroid and the corresponding reference centroid is smaller than half of the map resolution. Third, the assembly placement score is the average of its component placement scores, each of which is composed of a distance and an angle to the reference placement; the distance is calculated between the centroids of the placements, and the angle is the axis angle of the rotation matrix between the two placements. Because the components are kept rigid throughout the optimization process, the reference components used in the assessment of an assembly model are the component models superposed on the corresponding components in the native assembly (i.e., the reference placement). We chose not to use the Cα RMSD measure to assess assembly models because the significance of Cα RMSD values depends strongly on the number of assembly components and their sizes.

**Determining the configuration of Arp2/3**

To illustrate MultiFit, we first describe in detail a challenging application to Arp2/3 (Table 1, Figs. 2 and 3). The Arp2/3 complex of seven proteins is crucial for regulating the initiation of actin polymerization and the organization of the resulting filaments. A density map was simulated from the Arp2/3 crystal structure with ATP and Ca2+ (PDB entry 1TYQ). The atomic structures of the Arp2/3 components (proteins) were modeled using templates with sequence identity ranging from 16% to 99%.

**Table 1. Determining the configuration of the Arp2/3 assembly**

<table>
<thead>
<tr>
<th>Component (name, chain, residue range)</th>
<th>Component modeling Coarse-grained sampling</th>
<th>Fine-grained sampling</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Discretization (best placement score, fitting rank)</td>
<td>Optimization (best-scoring placement score)</td>
</tr>
<tr>
<td></td>
<td>Cα RMSD (Å)</td>
<td>NO3.5 (%)</td>
</tr>
<tr>
<td>Rpb1, A, 4 – 408</td>
<td>2q1nB, 4 – 370</td>
<td>40</td>
</tr>
<tr>
<td>Rpb2, B, 143 – 349</td>
<td>1nwkA, 140 – 334</td>
<td>48</td>
</tr>
<tr>
<td>ARPC1, C, 5 – 361</td>
<td>1erjC, 342 – 708</td>
<td>16</td>
</tr>
<tr>
<td>ARPC2, D, 1 – 274</td>
<td>1u2vF, 3 – 168</td>
<td>29</td>
</tr>
<tr>
<td>ARPC3, E, 1 – 169</td>
<td>2p9nE, 2 – 173</td>
<td>99</td>
</tr>
<tr>
<td>ARPC4, F, 3 – 186</td>
<td>1u2vD, 137 – 279</td>
<td>29</td>
</tr>
<tr>
<td>ARPC5, G, 11 – 150</td>
<td>2p9nG, 11 – 150</td>
<td>94</td>
</tr>
</tbody>
</table>

**Notes:**
- The percentage of sequence identity between the template and the component as calculated from their alignment used for comparative modeling.
- The Cα RMSD score and native overlap NO3.5 between the component model superposed on its native structure.
- The placement score of the placement found in the top-ranking assembly configuration.

† http://salilab.org/modeller
99%: the Cα RMSD error for these models varied from 0.4 Å to 21.4 Å, and their native overlap varied between 38% and 100%; we intentionally used inaccurate comparative models to benchmark the robustness of our method with respect to errors in the component conformations.

In the final output of MultiFit, the near-native model with an assembly placement score of (7.1 Å, 25°) was ranked 4th among all the sampled configurations. In coarse-grained sampling, this model was ranked 10th, with a configuration score of 4/7 and an assembly placement score of (10.8 Å, 136°). The centroids of the individual components were positioned in the vicinity of their native centroids; however, the orientations of some components were incorrect, resulting in steric clashes between components. In fine-grained sampling, the 20 top-scoring models were refined. The refinement procedure was able to resolve many of the clashes in the model, which in turn improved its global score, resulting into the final rank of 4th. Next, we elaborate on the individual steps of the optimization protocol.

In anchor graph construction, we computed seven anchor points from the density map. The average distance between the anchor points and the centroids of the corresponding reference components was 7.2 Å. We then identified pairs of anchor points that are sufficiently close to allow components placed in their vicinity to interact with each other. The procedure pruned 12 of the possible 21 pairs (i.e., 7·6/2). The remaining 9 pairs allowed identification of 9 of the 12 native contacts between the 7 components.

![Fig. 3. MultiFit results for Arp2/3. (a) An assessment of the final model with a mapping score of 0. The model has the fourth smallest value of the scoring function S (the fourth model in (b)). The modeled (color) and reference placements (gray) of the individual components are compared (Results); the corresponding placement scores are indicated below each comparison. (b) Five top-ranked models for Arp2/3. The atomic representations of the models are displayed on the top row. The bottom row shows the centroid and the rotation axis for each component; the corresponding rank, mapping score, configuration score, and assembly placement score are indicated below each model.](Image)
In the discretization step of coarse-grained sampling, we fitted each component into the neighborhood of each anchor point using ModEM. We assessed the accuracy of the discretization by the placement score of the best placement of each component (i.e., the placement with the lowest $C^\alpha$ RMSD corresponding to the reference). These best placement scores ranged from (2.5 Å, 30°) to (23.0 Å, 177°). As expected, as the model accuracy was assessed for the entire assembly did result in the correct placement occurring in the best-scoring assembly configuration. To validate the contribution of the shape complementarity score, we optimized a scoring function lacking this term ($S_\alpha$ in the scoring function $S$; Theory). The top-ranking configuration had a mapping score of 3, a configuration score of 3/7, and an assembly placement score of (42.5 Å, 94°). A model with a mapping score of 0 was not found in the top 50 solutions. This comparison demonstrates the positive contribution of the shape complementarity score to the accuracy of the generated assembly models.

### Benchmark

To assess MultiFit more comprehensively than is possible by a single example, we also applied it to a benchmark that included five additional simulated test cases. In all six simulated tests, a model with a mapping score of 0 was found within the top four solutions (Table 2); in fact, the model with the mapping score of 0 was the best-scoring model in all cases for which the structures of the individual components were modeled based on templates with sequence identities higher than 60%. The assembly placement score of the model with the mapping score of 0 ranged between (2.6 Å, 4°) and (7.1 Å, 25°). These results demonstrate the utility of MultiFit in predicting the configuration of atomic components in a low-resolution density map of their assembly. Next, we report the benchmark results at each of the five steps of the algorithm.

In anchor graph construction, the average distance between the predicted anchor point and the centroid of the corresponding reference component in the near-native configuration was between 4 Å and 7 Å. In the discretization step of coarse-grained sampling, a near-native configuration was sampled within the discrete sampling space in all test cases. However, this configuration was not necessarily ranked highly according to our scoring function due to steric clashes between interacting components.

In the optimization step of coarse-grained sampling, a model with a mapping score of 0 was found in the top 10 solutions in all test cases; in four of the six cases, it was the best-scoring solution. The assembly placement score of the model with a mapping score of 0 ranged from (2.6 Å, 4°) to (10.8 Å, 136°). The prediction accuracy depended on the
As the accuracy of the component models is decreased, the rank of the correct configuration and its placement score also become worse. The benchmark shows that coarse-grained sampling is able to determine component positions quite accurately, but frequently fails to result in accurate relative orientations. The main reason is the coarseness of the discrete sampling space, as demonstrated by the Arp2/3 and 1z5s examples. In the latter case, we obtained the near-native assembly [i.e., (5.9 Å, 113°)] with the native components and a less accurate configuration [i.e., (7.7 Å, 92°)] with distorted components.

In the discretization step of fine-grained sampling, the PATCHDOCK docking program was able to sample near-native interaction modes between pairs of components. However, these interactions were generally not ranked highly by PATCHDOCK. For example, in the 1z5s case with distorted components, the most accurate docking prediction of chains C and D against chain A ranked 405th and 138th, respectively.

In the optimization step of fine-grained sampling, the refined models were at least as accurate as the most accurate models generated in coarse-grained sampling, sometimes much more so. In particular, the accuracy of the relative orientations between pairs of interacting components improved. For example, in the 1z5s case with distorted components, the assembly placement score improved from (7.7 Å, 92°) to (6.4 Å, 62°). The refined model contained placements derived from the docking prediction of chains C and D against chain A. These placements were ranked 405th and 138th by PATCHDOCK; reweighing the placements by the normalized fitting score increased their ranks to 78th and 43rd, respectively. In the end, DOMINO correctly selected these placements for the final best-scoring configuration.

Benchmark with an experimentally determined map

To test the method in a realistic setting, we benchmarked it again by modeling the component configuration for an assembly with an experimentally determined cryoEM map.

**GroEL–GroES domains**

GroEL–GroES is a chaperonin that aids protein folding in *E. coli*. GroEL consists of two back-to-back rings of seven identical subunits, each of which contains three domains (i.e., the equatorial, intermediate, and apical domains). GroES is a ring of seven identical single-domain proteins that cap GroEL. We applied MultiFit to model the configuration of the four domains in an interacting pair of the GroEL and GroES subunits. Atomic coordinates for the four domains were obtained from a crystal structure of the GroEL–ADP–GroES complex (ADP state; PDB entry 1aon). The corresponding density was segmented from a cryoEM map of the bacterial GroES–ADP7–GroEL–ATP7 chaperonin determined component accuracy (Table 2). As the accuracy of the component models is decreased, the rank of the correct configuration and its placement score also become worse. The benchmark shows that coarse-grained sampling is able to determine component positions quite accurately, but frequently fails to result in accurate relative orientations. The main reason is the coarseness of the discrete sampling space, as demonstrated by the Arp2/3 and 1z5s examples. In the latter case, we obtained the near-native assembly [i.e., (5.9 Å, 113°)] with the native components and a less accurate configuration [i.e., (7.7 Å, 92°)] with distorted components.

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**Table 2. Benchmark**

<table>
<thead>
<tr>
<th>Assembly</th>
<th>Component modeling</th>
<th>Component configuration on a CryoEM Map</th>
</tr>
</thead>
<tbody>
<tr>
<td>Assembly (name, PDB entry)</td>
<td>Number of components</td>
<td>% Sequence ID (average (range))</td>
</tr>
<tr>
<td>Chaperonin GroEL, 1oel</td>
<td>Three domains</td>
<td>65 (60–72)</td>
</tr>
<tr>
<td>SUM - RanGAP1-Ubc9-Nup358/RanBP2 complex, 1z5s</td>
<td>Four proteins</td>
<td>100 (100–100)</td>
</tr>
<tr>
<td>SUM - RanGAP1-Ubc9-Nup358/RanBP2 complex, 1z5s</td>
<td>Four proteins</td>
<td>0.0 (0.0–0.0)</td>
</tr>
<tr>
<td>Dihydropyrimidine dehydrogenase, 1gte</td>
<td>Five domains</td>
<td>100 (100–100)</td>
</tr>
<tr>
<td>Archaeon Methanopyrus kandleri, 1e6v</td>
<td>Six proteins</td>
<td>61 (57–68)</td>
</tr>
<tr>
<td>Arp2/3 complex, 1tyq</td>
<td>Seven proteins</td>
<td>51 (16–99)</td>
</tr>
</tbody>
</table>

- **a**: The average, minimum, and maximum percentages of sequence identity between the assembly components and their templates.
- **b**: The average, minimum, and maximum C α RMSDs and native overlap NO3.5 between the modeled components and their corresponding component in the native assembly.
- **c**: The rank of a model with a mapping score of 0.
at 23.5 Å resolution (ATP state; EMDB ID 1046\textsuperscript{23}). The crystal structure of the ADP state was fitted to the density (as one rigid body) and used as reference for assessment. The main structural differences between the ATP and ADP states are the downward rotation of the intermediate domain and the counterclockwise twist of the apical domain.\textsuperscript{23}

The configuration with a mapping score 0 was ranked third, with an assembly placement score of (13.9 Å, 160°). A sampling space of approximately 14 million combinations was searched within 16 min of central processing unit time. The fine-grained sampling was able to generate a more accurate model with an assembly placement score of (11.0 Å, 84°). We note in passing that fitting all 49 domains (i.e., $3 \times 7 \times 2 + 7$) into the density of both rings would presumably benefit from the added information in the subunit–subunit interactions within and across rings; however, to test MultiFit in a more challenging setting, we deliberately modeled only a single symmetry unit consisting of three GroEL domains and one GroES domain.

Discussion

We described MultiFit, a computational method used for determining the positions and orientations (i.e., placements) of multiple atomic components in a cryoEM density map of their assembly. The problem is formulated in terms of combinatorial optimization, solved by our inferential optimizer DOMINO that guarantees the finding of the global minimum within a given discrete sampling space. The input is a density map and a set of atomic components that are kept rigid throughout the optimization process. For a given configuration of components, the scoring function measures the quality-of-fit of the atomic structures in the map, the protrusion from the map envelope, and the shape complementary between pairs of components. The optimization process consists of coarse-grained and fine-grained sampling stages. Each sampling stage starts with a discretization step achieved, respectively, by fitting and docking, followed by an optimization step that relies on DOMINO. Both DOMINO and MultiFit are available as part of Integrative Modeling Platform\textsuperscript{‡,48,49}

Accurate MultiFit predictions for seven test cases demonstrated its utility (Table 2). Specifically, our benchmark demonstrated the utility of MultiFit in predicting the configuration of components with known folds within a density map at resolutions between 20 Å and 23.5 Å; the average assembly placement score for the near-native configurations was (5.3 Å, 38°). MultiFit was able to determine the assembly configuration even in cases where the fitting scores were ambiguous. Examples include Arp2/3 (Table 1) and the 1z5s test case with distorted components (Table 2).

Next, we discuss (i) the benefits of simultaneous multiple component fitting, (ii) inaccuracies resulting from the discrete sampling space, and (iii) broad utility of combinatorial optimization.

Benefits of simultaneous fitting

Most methods for modeling assemblies in the context of a cryoEM map rely on a segmented assembly map and/or a model of the whole assembly. In the absence of such information, sampling the configuration space is computationally challenging, as the placement of each component may depend on the placements of other components. For example, the configuration of the Arp2/3 assembly with modeled components could not have been solved by iteratively fitting the largest remaining component in the unoccupied region using ModEM.\textsuperscript{31} Moreover, the configuration cannot be modeled accurately without the component protrusion and the interaction terms in the scoring function used by MultiFit. However, by considering the placements of all components simultaneously, the protrusion of a component from the assembly envelope, and the shape complementary between the interacting components, we were able to determine the assembly configuration with an assembly placement score of (7.1 Å, 25°).

Inaccuracies resulting from discrete sampling space

MultiFit prediction will be accurate when a near-native configuration exists in the discrete sampling space and corresponds to the global minimum of the scoring function. These two conditions depend, in turn, on the accuracy of the atomic models of the individual components and the choice of anchor points. Next, we elaborate on these two dependencies.

Accuracy of component models

The atomic models of the individual components might be inaccurate due to modeling errors, induced fit, and conformational selection.\textsuperscript{50} As the accuracy of the component models decreases, the discretized sampling space (either by fitting or by docking) is less likely to contain near-native placements (i.e., the sampling problem), and the global minimum is less likely to correspond to the most accurate sampled configuration (i.e., the scoring problem). In other words, these errors may affect the accuracy of the predicted assembly configuration due to scoring and sampling inaccuracies. One such example is the pair of 1z5s test cases (Table 2): The inputs to the first test case were the native components and the assembly density. The discretization steps of coarse-grained and fine-grained samplings resulted in near-native placements, and the top-ranked configuration detected by DOMINO had a relatively accurate assembly placement score of (5.0 Å, 67°). The inputs to the second test case were models with an average C\textsuperscript{\textalpha} RMSD error of 6.3 Å. The discrete sampling spaces

\[‡ \text{http://salilab.org/imp}\]
generated in the coarse-grained and fine-grained samplings contained less accurate placements. As a result, the utility of the scoring terms (especially the protrusion from the map envelope and the shape complementarity) decreased. The assembly placement score of the final assembly model with distorted component models was significantly worse (6.4 Å, 62°) than the assembly placement score of the assembly model with the native components. More accurate assembly models may be obtained by using a shape complementarity score that is less sensitive to component model errors and/or by an explicit treatment of the component conformations. To this end, techniques might be adopted from flexible fitting of a component into a density map32,51 and from flexible molecular docking.52,53

**Accuracy of anchor points**

Given the k-cluster algorithm, the utility of the anchor points is affected by the variances in the size and shape of the components (data not shown). The utility of the anchor points is also affected by the resolution of the map (data not shown). To obtain a discrete sampling space that contains a near-native configuration, we sample candidate placements of each component in the neighborhood of each anchor point. However, there are many assemblies for which the variation in component sizes is too large for reasonable neighborhood sizes. We intend to improve the utility of anchor point calculation by considering component sizes and density map segmentation.54,55

**Combinatorial optimization in structural biology**

Modeling challenges in structural biology can generally be expressed as optimization problems.48 These optimization problems often fall into a general class of NP-complete problems (Theory).56 Combinatorial optimization is a type of optimization in which the set of feasible solutions is discrete, and the goal is to find the best possible solution within this discrete set. Combinatorial optimizers have been suggested for various modeling tasks such as side-chain packing,57–59 threading,60 ab initio RNA folding,61 and prediction of quaternary structures of multiprotein complexes.62 These methods can, in principle, be reformulated as a combinatorial optimization of a scoring function represented by a graphical model, benefitting from graph theory techniques.24,25 Such a formulation has already been proposed for the side-chain packing problem.59

Our DOMINO method can, in principle, be applied to many problems in structural modeling, from low-resolution assembly modeling to side-chain refinement. Its strength derives from the junction tree algorithm that helps reduce the size of the search space from exponential in the number of components in the whole system to exponential in the number of components in the largest subset. More specifically, the computational complexity is $O(\frac{|U|L^2}{s})$, where $|U|$ is the number of subsets in the junction tree, $L$ is the size of the largest subset, and $s$ is the number of discrete values of a single variable in the graphical model. Fortunately, at the granularity level used in MultiFit’s application to protein assemblies in our benchmark, the theoretical complexity of the junction tree algorithm has not been a limiting factor. Nevertheless, in other applications that involve a dense graphical model of the scoring function and extensively sampled variable values, incomplete sampling of a discrete space may have to be accepted.

In conclusion, MultiFit and DOMINO can help bridge the gap between the atomic structures of the individual proteins and the cryoEM maps of their assemblies. In particular, they can provide initial configurations for further refinement of many multicomponent assembly structures described by electron microscopy.

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