Estimation of Distribution Algorithm for Protein Structure Prediction

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Abstract

Proteins are essential for maintaining life. For example, knowing the structure of a protein, cell regulatory mechanisms of organisms can be modeled, supporting the development of disease treatments or the understanding of relationships between protein structures and food attributes. However, discovering the structure of a protein can be a difficult and expensive task, since it is hard to explore the large search to predict even a small protein. Template-based methods (coarse-grained, homology, threading etc) depend on Prior Knowledge (PK) of proteins determined using other methods as X-Ray Crystallography or Nuclear Magnetic Resonance. On the other hand, template-free methods (full-atom and ab initio) rely on atoms physical-chemical properties to predict protein structures. In comparison with other approaches, the Estimation of Distribution Algorithms (EDAs) can require significant less PK, suggesting that it could be adequate for proteins of low-level of PK. Finding an EDA able to handle both prediction quality and computational time is a difficult task, since they are strong inversely correlated. We developed an EDA specific for the ab initio Protein Structure Prediction (PSP) problem using full-atom representation. We developed one univariate and two bivariate probabilistic models in order to design a proper EDA for PSP. The bivariate models make relationships between

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dihedral angles $\phi$ and $\psi$ within an amino acid. Furthermore, we compared the proposed EDA with other approaches from the literature. We noticed that even a relatively simple algorithm such as Random Walk can find the correct solution, but it would require a large amount of prior knowledge (biased prediction). On the other hand, our EDA was able to correctly predict with no prior knowledge at all, characterizing such a prediction as pure \textit{ab initio}.

\textbf{Keywords:} Estimation of Distribution Algorithm, Protein Structure Prediction, \textit{ab initio}, full-atom, van der Waals energy, Expectation-Maximization.

1. Introduction

From the discovery of a new disease until the development of its cure can take about ten years and involve costs of five billion dollars \cite{1}. One of the main reasons for this amount of money and time needed is the problem of finding the tertiary structure of the protein responsible for the disease. Most of the existing methods to determine the protein structures are experimental. They attempt to look at the proteins as they are present in nature. For instance, the X-Ray Crystallography experimental method shoots an x-ray beam into a protein crystal. Then it creates a diffraction map of hydrogen atoms and together with information of the primary structure of the protein, it is possible to construct the tertiary structure. The Nuclear Magnetic Resonance needs a solution of high concentration containing the target protein. This solution is submitted to a process that will excite the atom spins and, depending on how much the atoms will move to a different spin state, it is possible to construct the tertiary structure of a protein. Both of these experimental methods have disadvantages and they cannot always determine the tertiary structure of a protein \cite{2}.

Thus, in order to avoid the high costs and time needed by experimental methods, computational methods (\textit{in silico}) have been created. The method to determine the primary structure of proteins is well established nowadays, but as we know, the equivalent tertiary structure is not. Thus, from the se-
sequence of amino acids, computational methods attempt to look into the search space in order to find feasible protein configurations. We know that proteins in nature stabilize with the minimum energy state. Therefore, computational methods look for protein configurations that represent the minimum energy state. It means that we are predicting the protein structure. This problem is known as Protein Structure Prediction (PSP). There are two main computational approaches to predict proteins. One uses knowledge of proteins that were determined using other methods, as experimental, to infer the new ones (called template-based). Other, as used in this work, is called template-free. It does not use information about any existing structures to predict the new ones. Instead, it uses energy potentials so it can evaluate how good a configuration is. Then, when a configuration with a lower energy value is found, we will have a hypothesis that we found the correct structure. However, the search space of protein configurations is huge and this task cannot be performed using exact methods [3].

Our goal is to find the correct set of the dihedral angles $\phi, \psi, \chi$’s (Section 2.1), i.e. the variables of the problem, that will yield the protein configurations with the lowest possible energy. Thus, this can be treated as an optimization problem in which we want to minimize the energy of a protein configuration (our fitness) changing the dihedral angles (the variables). There are different ways that one can represent the protein configurations in the computer. One could use a coarse-grain representation of amino acid, called HP model [4], but it cannot represent the protein for practical purposes. Therefore, we use the full-atom model considering all atoms of the protein configurations [5].

Despite the efforts of the in silico methods in trying to find the correct protein configurations, an appropriate algorithm that can work properly in a pure ab initio way is still missing, which is especially relevant when one wants to predict a protein that has low similarity with the known structures. There are some Evolutionary Algorithms (EA) and other optimization approaches with useful results for PSP [6, 7, 8, 9]. The better the optimization technique, the better will be its chances of finding proper solutions. Nevertheless, this is not

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a trivial task since the optimization algorithm has to deal with the large search space of possible 3D protein structures in an efficient manner.

Thus, considering the characteristics of the PSP problem, we build an Estimation of Distribution Algorithm (EDA) \([10]\) specially designed for such a problem. Basically, an EDA constructs models of the distribution of variable values from promising regions of the search space in order to improve the optimization process. To sample better solutions, we need to have a good probabilistic model. First, we developed a Univariate model-based Optimization algorithm (UNIO). Secondly, taking advantage of properties of the problem we are tackling, we modeled the \([\phi, \psi]\) within the same residue as correlated, yielding a bivariate probabilistic model. We proposed two different ways to estimate a probability distribution for each pair of variables: (i) Kernel Density Estimation model-based Optimization (KDEO) \([11]\) and (ii) Finite Gaussian Mixtures model-based Optimization (FGMO) \([12]\). That yielded three novel different EDAs for PSP: UNIO, KDEO and FGMO.

In order to evaluate our probabilistic models, we measured three aspects: the quality of the protein configurations (RMSD, root-mean-square deviation) \([13]\), the computational time consumption and the energy values of the predicted structures (Section\([4]\)). As we expected, the bivariate models (KDEO and FGMO) were able to find better solutions in terms of the energy and RMSD. Furthermore, we made a comparison between KDEO and specifically other optimization approaches from the literature, Random Walk (RW) \([14]\), Monte Carlo (MC) \([15]\), Genetic Algorithm (GA) \([16]\) and Differential Evolution (DE) \([17]\). We discovered that all of these optimization techniques were able to find the correct protein configuration, of a specific case, when there is enough previous knowledge about a promising region of the search space. For instance, RW is known as a poor optimization technique, thus, in order to find the correct protein configuration, we introduced a bias, reducing the search space to a neighborhood of the dihedral angles of the native protein. On the other hand, the proposed EDA can, in general, find adequate solutions without search space reduction, that is, without bias. Therefore, we may concluded that the proposed EDA
is more adequate than other investigated approaches when predicting proteins with low similarity to known structures.

In the next section, we present the Protein Structure Prediction problem. In Section 3, we present our proposed EDA for PSP. Then, the results are shown in Section 4. Finally, the conclusions of this work are made in Section 5.

2. Protein Structure Prediction problem

Proteins are essential in almost all processes of life. They have different functions in living beings and the function of each protein is associated with their shape. For instance, a protein responsible for the transportation of other molecules has its shape appropriate to carry such a molecule along the organism until its destination, i.e. the hemoglobin, capable to transport oxygen. There are several other functions of proteins as, for instance, defense, control, regulation, breaking covalent binds etc. We say that a function of a protein is associated to its shape, also called tertiary structure [18].

When we know a protein responsible of causing a disease, we can develop treatments and inhibit that protein function. For instance, some viruses have the capability of cutting human cells, entering in it and then making copies of themselves spreading the disease along the rest of body. However, when we know the shape of the protein responsible for cutting the cell we can develop medicines, the complementary structure, which bind the virus in the medicine molecule instead of the human cell. This will probably lead to the death of the virus and reduce the effects of the virus on the organism [3].

Every protein has an identification that matches with its shape. This is called primary structure (or sequence of amino acids) and it stores all the residues chain of a protein. Nowadays, it is relatively simple to isolate the primary structure of a protein. However, finding the tertiary structure equivalent to the primary structure is a very complicated task. This is the main reason why the sequence database is growing fast [19] and the structures database is growing relatively slow [20].
The current methods to find the tertiary structure of proteins are expensive and require years of trial and error. The experimental methods X-Ray Crystallography (XRC) and Nuclear Magnetic Resonance (NMR) are by far the most used methods to determine protein structures. However, it is not always possible to use these methods, since they have disadvantages as, for instance, sometimes it is not possible to have a protein crystal and in this case, the XRC will be no longer available. There is no need to have a protein crystal in NMR but it only works for small proteins [13].

Based on the drawbacks of the experimental methods many computer methods that try to find the structure of proteins were developed. This is known as \textit{in silico} methods. Based on the sequence of amino acids, they look for feasible protein configurations in the search space and try to predict the protein structure. The \textit{in silico} methods are divided into two different ways of prediction. First, they are based on prior knowledge of proteins that were already determined by other methods, as XRC and NMR. They are promising techniques and there are many researches about these methods. One problem of these techniques is that their predictions can be biased toward the experimental methods. The second method is called template-free and does not make use of knowledge of other proteins. Instead, it uses energy potentials in order to compute the energy of the protein configuration. The energy potential is a way to describe how a protein configuration represents a protein in real life. Knowing that proteins stabilize in a state of minimum energy, we also want to find the configurations with lowest energy. In this work, we use a template-free approach with full-atom representation [21].

2.1. Protein representation

The protein configuration can often be represented using dihedral angles \( \phi, \psi, \chi \)'s. Each residue in a protein configuration chain has its own set of \([\phi, \psi]\) angles and the number of \(\chi\)'s depends on the type of residue. All these angles range from \(-180\) to \(180\) degrees, the search space range. The \(\phi\) angle is the dihedral angle between atoms \(N - C_\alpha\) within the same residue and the \(\psi\) is
related to the $C_\alpha - C$ angle. The set of all dihedral angles $\phi$’s, $\psi$’s, $\chi$’s from all residues of a protein configuration has all information needed to evaluate the protein. Thus, the variables of the problem can be understood as a vector of dihedral angles ranging from $[-180; 180]$ (Figure 1).

In order to compute the fitness of a protein configuration, we first need to convert the dihedral angles into Cartesian coordinates. Then, knowing the Cartesian coordinates of all atoms and their types it is possible to compute the energy (the fitness).

![Figure 1: An example of the dihedral angles $[\phi, \psi]$ from the sequence of amino acids using a full-atom representation (the side chains are hidden).](image)

2.2. The fitness function

Several potential energies contribute to the protein energy. There are bonded and non-bonded potential energies and each of them has a specific contribution to the molecule stabilization. We know that non-bonded potential energies have the largest contribution to the molecule energy. In our algorithm, called ProtPred \[7\], we have five bonded potential energies (bond stretching, angle bending, Urey-Bradley, improper dihedral and torsional angle) and four non-bonded potential energies (van der Waals, electrostatic, solvation and hydrogen bonds). Despite having all these potential energies implemented in our algorithm, we
decide to use only van der Waals energy in this paper, since it has the highest contribution to molecule energy. Besides, it makes it easier to understand the results so we can keep the focus on the evolutionary process. Thus, the fitness function is determined only by the van der Waals energy.

The van der Waals energy models the attraction and repulsion among atoms. In general, the Lennard-Jones potential (also known as Lennard-Jones 12 − 6) is used to compute the van der Waals energy of a protein configuration. The van der Waals energy changes according to the distance and the atom’s type (nitrogen, carbon, oxygen, hydrogen etc). Equation 1 describes the relative distance between two atoms \(i\) and \(j\), given the Euclidian distance between them \(d_{i,j}\) and the van der Waals radii constant \(R\) of atoms \(i\) and \(j\):

\[
v_{ij} = \frac{d_{i,j}}{R_i + R_j}.
\]

(1)

The van der Waals energy is very repulsive at short distances since the electron cloud between atoms starts to overlap. At this distance, the energy rapidly increases and tends to infinity. The equilibrium point, known as van der Waals contact, happens when the Euclidian distance between the atom pair is neither too far nor too close. This is the point of minimum energy of an atom pair. If the atom pair is too far from each other they will not have any type of interaction and the energy will tend to zero. We used a cutoff of 8 Å in order to avoid unnecessary computation, and to avoid dealing with large numbers we also set a tapering-off (see Figure 2). In case that \(v_{ij}\) is smaller than 0.8 we assume a constant \(C\). The Lennard-Jones potential used in our EDA for PSP is described by:

\[
f_{LJ}(v_{ij}) = \begin{cases} 
Av_{ij}^{-12} - Bv_{ij}^{-6} & \text{if } v_{ij} > 0.8, \\
C & \text{if } v_{ij} \leq 0.8,
\end{cases}
\]

where \(A\) and \(B\) are constants experimentally determined based on characteristics of the environment, and \(C\) is given by \(Av_{ij}^{-12} - Bv_{ij}^{-6}\) with \(v_{ij} = 0.8\).

The van der Waals energy is the sum of the Lennard-Jones potentials between all atom pairs in a molecule. There are \(\frac{n(n-1)}{2}\) interactions, where \(n\) is the
number of atoms, leading to:

\[
E_{vdw} = \sum_{i=1}^{n-1} \sum_{j=i+1}^{n} f_{LJ}(v_{ij}).
\]  

(3)

Figure 2: The modified van der Waals function we used, highlighting the tapering-off (when atoms are too close), the van der Waals contact (ideal distance) and the cutoff (they are too far).

Thus, in order to have the minimum global energy of a molecule it is necessary to find a compromise between the partial energies among atoms to get the least van der Waals energy. To find good van der Waals energy values we need to change the dihedral angles of our protein configuration. Changing the dihedral angles will lead to a change in Cartesian coordinates of atoms in the molecule as well. Thus, there is a set of dihedral angles that will imply the best positioning of atoms, yielding a global minimum in van der Waals energy.
3. Estimation of Distribution Algorithm for Protein Structure Prediction

In the previous section, we concretized the problem we want to tackle. In this section, we will describe how we can find promising solutions using EDAs, i.e., how can the EDA find a good set of dihedral angles that will express a low energy molecule.

The EDAs are a relatively new class of the EA. They are optimization techniques that use probabilistic models from a promising set of solutions in order to sample the offspring. In some cases, they also have the capability of accounting for correlations between variables. In the literature, the EDAs for binary and discrete variables are well described, since the probabilistic models and variable relationship are relatively easy to understand [24]. In fact, using a simplified representation of proteins, [24] showed that it’s possible to achieve relevant results with an EDA for the PSP problem.

However, dealing with dihedral angles (in the continuous search space) is relatively more difficult since it is not possible to map the combination of all variable values. Besides, the probabilistic model should be able to deal with multimodality, since the distributions of dihedral angles in the PSP problem are non-parametric. We also want something that can handle the variable relationship between $[\phi, \psi]$ within the same residues, so it has to be bivariate.

There are some real-valued EDAs in the literature as well. However, they would not be totally appropriate for the PSP problem. For example, the Univariate Marginal Distribution Algorithm (UMDA$_c$) [25] works in the continuous search space but cannot deal with multimodality neither variable relationship. The Bivariate Marginal Distribution Algorithm (BMDA) could model the variable relationship but not the multimodality aspect [26]. Since then, EDAs able to tackle the multimodality and the variable relationship aspects were developed, as the case of the EGNA [25], IDEA [27, 28] and PBIL [29]. The real-valued Bayesian Optimization Algorithm (rBOA) [30] can also handle both multimodality and variable relationship. It was shown that rBOA can outper-
form the mIDEA \cite{31} in several benchmark problems. However, the evidence that rBOA is able to predict protein structures using the full-atom representation has not been provided in the literature yet.

Knowing that the dihedral angles $\phi$ and $\psi$ within the same residue have a strong relationship, we designed an EDA in which there is no need to learn the variable relationship: we treat the dihedral angles $[\phi, \psi]$ of each residue as correlated variables. Besides, the statistical mechanisms used in previous EDAs as, for instance, the normal kernels and the mixtures of Gaussian distribution were used to build our new EDA for PSP full-atom. We already showed in a previous work that EDAs can successfully be applied in PSP using full-atom and \textit{ab initio} modeling \cite{32}. In this work, we show three different approaches that were developed. The detailed information about each of them is described in the next sections.

3.1. \textit{Univariate model-based Optimization}

The Univariate model-based Optimization (UNIO) is the simplest algorithm. It does not deal with variable relationships. Instead, it deals with multimodality in an efficient way. From the promising individuals (selected) the process of a one-dimensional kernel density estimation (KDE) is simulated for each variable involved in the problem. However, creating a kernel distribution for each problem variable would require high computational costs since it is necessary to iterate between all observations (the selected size) per variable.

KDE is based on the sums of the difference between the point of interest $x$ and all observations within a data set $(x_1, x_2, \ldots, x_n)$ over a bandwidth value $h$. In order to sample new values from our kernel distribution we first need to build a Probability Density Function (PDF). This can be done by calling the KDE function as many times as is required to fill the range of the $x$ values. That would require high computing time per generation. For instance, consider the selected size (of the EDA) is $n = 500$, and the number of points needed to build the PDF is $400$, then for $100$ dihedral angles ($100$-dimensional problem), we would need to call the KDE function at least $500 \times 400 \times 100 = 20,000,000$
at each generation of the EDA.

Thus, to keep the algorithm efficient, we extended the idea of the KDE in a different manner. Instead of creating the PDF from the kernels, we simply take a random observation from \( x \) and add a perturbation to it with the distribution \( N(0, 1) \).

Consider the selected set \( S_i^j \) where \( i = 1 \ldots n \) and \( j = 1 \ldots m \), being \( n \) the size of selected individuals and \( m \) the number of variables of the problem. We want to fill the offspring \( O_j^k \) where \( k = 1 \ldots o \) and \( o \) is offspring size. For a random point drawn from the selected \( S_i^j \) for a variable \( j \), we add some perturbation to it and put this new value into \( O_j^k \) for \( k = j = 1 \). Then we repeat this process for \( j = 1 \ldots m \) (for all variables of the first individual of the offspring) and then for all the remaining individuals \( (k = 1 \ldots o) \). The Algorithm 1 shows how this technique works.

**Algorithm 1** Random Points - It creates the offspring from selected.

**Require:** Selected individuals \( S \), Size of selected \( n \), Size of the Offspring \( o \), number of problem variables \( m \)

**Ensure:** Offspring \( O \)

\[
\text{for } j = 1 \text{ to } m \text{ do} \\
\quad u \leftarrow \text{Sample } o \text{ values from Uniform Discrete Distribution ranging from } [1, n] \\
\quad O_j \leftarrow S_j^u + N(0, 1)
\]

\[
\text{end for}
\]

We carried out an experiment to find out whether this kernel simplification is promising. We noticed that the simplification of KDE, indeed, does not have the same accuracy of the KDE, but it is much faster. So we are compromising a little of accuracy with a lot of reduction in computational cost.

Figure 3 shows an example of the comparison between our proposed simplification (Random points) and the traditional KDE. From a sample data set with three modes, we simulated new values using KDE and our proposed simplification. We repeated this procedure 30 times. The red line represents the true density. Density plots of the simulated values with KDE and our proposed
simplification are shown as well as the smoothed standard deviations.

Figure 3: Comparison of simulating data with KDE and Random Points: (a) a comparison with the true density of the data and (b) a time comparison.
3.2. Kernel Density Estimation model-based Optimization

Considering that all variables can interact with each other we could create an \( m \)-dimensional KDE. Nevertheless, that would be very difficult to handle. Then, we decided to use the two-dimensional Kernel Density Estimation, yielding the KDEO (Kernel Density Estimation model-based Optimization). It correlates the dihedrals angles \([\phi, \psi]\) of amino acids within the same residue. We know that these two variables are strongly correlated since rotations in \(\phi\) generally produce stereochemical constraints for its closest neighbor dihedral angle \(\psi\). Moreover, an implicit correlation is made with \([\phi, \psi]\) within the same residue every time one looks at the Ramachandran plot [33], since the plot itself shows a density map between angles \(\phi\) and \(\psi\).

In KDEO, the \(\psi\) values are generated conditional on \(\phi\). Firstly, the KDE is created for the \(\phi\) and a new value \(\phi'\) from its distribution is sampled. Then, a two-dimensional KDE map of \([\phi, \psi]\) is created. The closest value to \(\phi'\) in the two-dimensional KDE (in \(x\) axis direction) is taken to be the conditional KDE. Finally, a new \(\psi'\) value is sampled from the two-dimensional KDE (in \(y\) axis direction).

For the two–dimensional case, kernel density estimates are obtained via Equation 4:

\[
\hat{f}_h(x_1, x_2) = \frac{1}{n} \sum_{i=1}^{n} \frac{1}{h_1 h_2} K \left( \frac{x_1 - \phi_i}{h_1} \right) K \left( \frac{x_2 - \psi_i}{h_2} \right),
\]

where \(K(\cdot)\) is the kernel. In this work, we used the normal kernel

\[
K(x) = (2\pi)^{-1/2} e^{-0.5x^2}
\]

and the bandwidth \(\hat{h}\) is calculated using Equation 6:

\[
\hat{h} = \begin{cases} 
4 \cdot 1.06 \cdot \hat{\sigma} \cdot n^{-1/5} \text{ if } \hat{\sigma} < t, \\
4 \cdot 1.06 \cdot t \cdot n^{-1/5} \text{ if } \hat{\sigma} \geq t,
\end{cases}
\]

where \(4\) is a multiplicative factor [34, 35] and \(t\) equals
\[
t = \frac{Q_3 - Q_1}{1.34},
\]
(7)

where \(Q_3\) and \(Q_1\) are the third and first quartiles, respectively.

All of these processes require many computational resources. In order to speed up the evolutionary process we first check if the \([\phi, \psi]\) has a normal distribution using the Anderson-Darling test. If the p-value is large than 5\%, then the KDEO is bypassed and a bivariate normal distribution is used instead. Otherwise, the KDEO is used. This strategy is interesting because at the beginning of the evolutionary process we can have many modes. However, as the evolutionary process starts to converge the kernel is not necessary anymore for all pairs \([\phi, \psi]\) (see Algorithm 2).

**Algorithm 2** Two-dimensional Kernel Sampling - It creates the Offspring from selected

**Require:** Selected individuals \(S\), the number of protein residues \(r\), size of the new data \(o\)

**Ensure:** Offspring \(O\)

for \(i = 1\) to \(r\) do

\(\phi \leftarrow \text{Get } \phi \text{ from residue } i \text{ from } S\)

\(\psi \leftarrow \text{Get } \psi \text{ from residue } i \text{ from } S\)

if p-value from Anderson-Darling test of \([\phi, \psi]\) < 0.05 then

\(\phi' \leftarrow \text{Random Points } (\phi, o)\)

\(P \leftarrow \text{2D Kernel Density Map } (\phi, \psi)\)

\(\psi' \leftarrow \text{Sample from PDF } P \text{ conditional on } \phi'\)

else

\(\phi', \psi' \leftarrow \text{Sample } o \text{ values from two-dimensional Gaussian } N([\mu_\phi, \mu_\psi], \Sigma_{\phi\psi})\)

end if

\(O^i \leftarrow \phi', \psi'\)

end for

For instance, similarly to \([\phi, \psi]\), let \(x_1\) and \(x_2\) be two vectors of data dis-
Figure 4: Two-dimensional KDE example: (a) the data \( x_1, x_2 \), (b) the univariate distribution of \( x_1 \) (top), the kernel map created for \( x_1, x_2 \) (middle) and the distribution of \( x_2 \) given the value of \( x_1 \) (bottom).

First, a new vector \( x'_1 \) is sampled from the independent distribution of \( x_1 \). Then, the two-dimensional KDE map is created for the pair \( [x_1, x_2] \). For each new point of \( x'_1 \) new points of \( x'_2 \) are sampled conditional to the previous value of \( x'_1 \). Looking at the Figure 4b, let’s assume that \( x'_1 = 3.5 \) (top). As the two-dimensional KDE map must be represented using discrete variables for \( x_1 \) and \( x_2 \), we need to pick the closest point where \( x_1 = 3.5 \) (middle) and use this point as the conditional distribution (bottom). Finally, for each sampled \( x'_1 \) we sample new \( x'_2 \) values\(^1\).

### 3.3. Finite Gaussian Mixtures model-based Optimization

As a KDE alternative, we develop an EDA called Finite Gaussian Mixtures model-based Optimization (FGMO). Despite having the implementation of the \( m \)-dimensional FGMO we used only the two-dimensional in this paper. The

\(^1\)The complete animation can be accessed at:

http://icrserver.icmc.usp.br/~daniel/ani/kde2d/
FGMO combines mixtures of Gaussian distributions in order to estimate the density of data non-parametrically. Each mixture component \( k \) (cluster) has its own mean \( \mu_k \), variance matrix \( \Sigma_k \) and the mixture weight \( \pi_k \). From a given number of mixtures \( K \), we want to know their set of parameters \( \theta = [\mu, \Sigma, \pi] \) for all mixture components. The Expectation-Maximization (EM) is often used to estimate the parameters of Gaussian mixtures [36]. In order to get the estimated \( \hat{\theta} \), the EM algorithm takes a starting setting of parameters \( \hat{\theta}_0 \) and iterates between E-Step and M-Step. The algorithm converges when the log-likelihood between two iterations is less than a defined value (1.5 in our experiments). The E-Step updates a probability matrix \( w_{j,k} \) via:

\[
 w_{j,k} = \frac{\hat{\pi}_k f(x_j, \hat{\mu}_k, \hat{\Sigma}_k)}{\sum_{l=1}^K \hat{\pi}_l f(x_j, \hat{\mu}_l, \hat{\Sigma}_l)},
\]

where \( f(x, \hat{\mu}, \hat{\Sigma}) \) can be defined as

\[
 f(x, \hat{\mu}, \hat{\Sigma}) = \frac{1}{(2\pi)^{p/2}\Sigma^{1/2}} \exp \left\{ -\frac{1}{2} (x - \hat{\mu})^T \hat{\Sigma}^{-1} (x - \hat{\mu}) \right\},
\]

and the M-Step updates the parameters via

\[
 \hat{\pi}_k = \frac{1}{n} \sum_{i=1}^n w_{ik},
\]

\[
 \hat{\mu}_k = \frac{\sum_{i=1}^n w_{ik} x_i}{\sum_{i=1}^n w_{ik}},
\]

\[
 \hat{\Sigma}_k = \frac{1}{\sum_{i=1}^n w_{ik}} \sum_{i=1}^n w_{ik} (x_i - \hat{\mu}_k)(x_i - \hat{\mu}_k)^T.
\]

For each pair \([\phi, \psi]\) from the selected set we run a two-dimensional EM for a given number of mixture components \( K \), yielding the estimates \( \hat{\theta} \). From \( \hat{\theta} \) we randomly select a mixture component using a uniform distribution with weights \( \pi_k \) and sample the new \([\phi', \psi']\) values at once, using their parameters (see Algorithm 4). The \( \hat{\theta} \) is then used to sample all the different offspring individuals. Then, for the \([\phi, \psi]\) of the next residue we estimate a new \( \hat{\theta} \) and use
these new parameters to sample the values for this next residue. This process continues until it reaches the last residue of the molecule.

For instance, consider a protein with 5 residues. The selected individuals will have 5 \([φ,ψ]\) pairs \([φ_1,ψ_1]\); \([φ_2,ψ_2]\); \([φ_3,ψ_3]\); \([φ_4,ψ_4]\); \([φ_5,ψ_5]\]), where \(i = 1 \ldots n\) being \(i\) the individual index in the selected set and \(n\) is the total number of selected individuals. For the pair \([φ_1,ψ_1]\) we perform a complete EM algorithm and get \(\hat{θ}_1\). It contains the \(µ_k, π_k\) and \(Σ_k\) for the mixtures \(k\). Then, the pair \([φ'_1,ψ'_1]\) is generated by sampling values from \(\hat{θ}_1\). Next, \(\hat{θ}_2\) is estimated using \([φ'_2,ψ'_2]\) and then a bivariate mixture with new parameter \([φ'_2,ψ'_2]\) is simulated. This continues until the residue number 5. It is important to notice that in this case, there are five EM algorithms running at each generation. This can have high computational costs since the EM is an iterative process. In order to speed up the algorithm, we replaced the general PDF (Equation 9) by a specific expression for the two-dimensional case, as shown in Equation 13:

\[
f(x_1, x_2) = \frac{1}{2\pi\sigma_1\sigma_2\sqrt{1-\rho^2}} \exp \left\{ -\frac{z}{2(1-\rho^2)} \right\},
\]

\[
z = \frac{(x_1 - \mu_1)^2}{\sigma_1^2} + \frac{2\rho(x_1 - \mu_2)(x_1 - \mu_2)}{\sigma_1\sigma_2} + \frac{(x_2 - \mu_2)^2}{\sigma_2^2}.
\]

Besides, before computing the FGMO we check whether the standard deviation of both \(φ\) and \(ψ\) is small (say 0.01). If true, then we bypass the EM algorithm and use a bivariate normal Gaussian instead (see Algorithm 3).

During the EM iterations, we also needed to treat special cases to avoid division by zero in Equation 8 that may happen when there is a very far outlier with a very small standard-deviation in the data set. In this case, we stop the EM iterations and use the last valid \(\hat{θ}\).

Figure 5 shows an example of how the FGMO works. At the first iteration, the parameters are set to the initial condition. Then, according to the EM iterations, the parameters tend to converge. Figure 5b shows the parameters after the convergence. Finally, new values are sampled (Figure 5c) from its fitted mixtures model.
Algorithm 3 Two-dimensional Finite Gaussian Mixtures based-model Optimization - Sample the offspring using FGMO

Require: Selected individuals $S$, the number of protein residues $r$, size of the offspring $o$, mixtures components $K$

Ensure: Offspring $O$

for $i = 1$ to $r$ do
    $\phi \leftarrow$ Get $\phi$ from residue $i$ from $S$
    $\psi \leftarrow$ Get $\psi$ from residue $i$ from $S$
    if Standard deviation of $\phi > 0.01$ and Standard deviation of $\psi > 0.01$ then
        $\hat{\theta} \leftarrow$ Expectation-Maximization ($\phi; \psi; K$)
        $[\phi'; \psi'] \leftarrow$ Sample $o$ individuals from fitted model $\hat{\theta}$
    else
        $\phi', \psi' \leftarrow$ Sample $o$ values from two-dimensional Gaussian $N(\mu_\phi, \mu_\psi, \Sigma_{\phi\psi})$
    end if
    $O^i \leftarrow [\phi'; \psi']$
end for

Algorithm 4 Two-dimensional Finite Gaussian Mixtures - Simulating

Require: The fitted model $\hat{\theta}$ and the size of the new data $n$

Ensure: Simulated $s$

$x \leftarrow$ Sample from distribution $U(0, 1)$
$c \leftarrow$ Cumulative sum of $\hat{\theta}_x$
$k \leftarrow 1$

while $x < c_k$ do
    $k \leftarrow k + 1$
end while

$s \leftarrow$ Sample from $N(\hat{\theta}_{\mu_k}, \hat{\theta}_{\sigma_k})$
Figure 5: FGMO example. (a) The first iteration of the EM algorithm. The original data slightly overlapped with two mixture components. The red dots represent the means of each mixture and the contour curves their densities, (b) the last iteration of the EM and (c) the sampled values.

4. Results

The results show that the three proposed EDAs (UNIO, KDEO and FGMO) performed properly for PSP. For each proposed method, we evaluated three different aspects: (1) the computational cost, measuring the overall time of a prediction; (2) the van der Waals energy value of the best individual at the last generation and (3) the RMSD, which represents how similar a solution is to the native protein.

We have to be careful when evaluating (3) since we are only using van der Waals energy in our fitness function. In this case, we are bypassing other potentials that are less relevant to stabilize the molecule. Adding other potentials in pure \textit{ab initio} PSP would require a much more complex algorithm, i.e. the Multi-Objective method proposed by [9]. This paper shows that we can find adequate pure \textit{ab initio} protein configurations using only van der Waals and the proposed EDAs. We compared the KDEO against other optimization methods from the literature, as Random Walk (RW) [14], Monte Carlo (MC) [15], Genetic Algorithm (GA) [37] and Differential Evolution (DE) [17]. Appendix A shows a pairwise comparison using the Wilcoxon test [38] for all evaluated methods considering the three aspects: van der Waals energy, RMSD and running time.
4.1. Experimental setup

The experiments were run in the 20 nodes cluster of the Laboratorio de Computacao Reconfiguravel at the ICMC-USP. Each node has an Intel Core i7 2.67 GHz processor with 4 physical processors, 8 considering Hyper-Threading technology. It also has 32 GB of RAM and the Operational System is Debian 4.6.3-14 64 bits. Moreover, each node has two network adapters. One is used for the file system (NFS) and other to communication operations in MPI [39].

We have selected four small proteins from the Protein Data Bank (PDB) [40], all containing α-helices. This may be favorable for the optimization model used that is based on van der Waals only. Table 1 shows the PDB ID, the number of residues and the problem length ($m$). Figure 6 shows the shape and the $[\phi,\psi]$ dihedral angles distributions of each native structure.

| Protein | Residues | Problem length |
|---------|----------|----------------|
| 1A11    | 25       | 95             |
| 2LVG    | 40       | 169            |
| 2KK7    | 52       | 229            |
| 2X43    | 67       | 268            |

The convergence is reached when either each of the tested algorithms reaches one million evaluations or the standard deviation of the population fitness falls below 0.0001.

4.1.1. Performance issues

ProtPred is entirely written in C language and it uses efficient libraries as GSL and CBLAS [41] to deal with most algebraic and statistical operations. Some functions were taken from statistical language R [42] and translated into C. The van der Waals energy uses an efficient implementation based on cell-lists, as we proposed in an earlier work [43]. Furthermore, the tested approaches have several different parameters to set, so we used MPI in order to distribute each
algorithm configuration throughout our cluster. Despite this, we needed about 12,000 hours of CPU time to get all the results shown in this paper.

4.2. Proposed Estimation of Distribution Algorithms

Before we run the sequence of experiments, we found a set of parameters of the EDA according to the probabilistic model used (Table 2). For each combination of parameters, the experiments were repeated 30 times with different seeds for each protein.

Figure 7 shows the results obtained for the smaller protein 1A11. Consider...
er the energy aspect only, the FGMO was best, although it has some high energetic outliers. The KDEO performed best when considering the RMSD. The running time results were partially surprising since a sophisticated probabilistic model as FGMO was faster than UNIO, in most cases. That happens because the selected size used by UNIO needed to be 20 times larger than FGMO.

In the PSP problem, a trade-off between energy and RMSD is sometimes difficult to obtain since the predicted energy can exceed the native energy value, when using only van der Waals energy. A scatterplot of van der Waals energy per RMSD (Figure 7d for protein 1A11) enables to evaluate such trade-off. We also highlighted in yellow the set of points that dominate other ones, in terms of multi-criteria optimization [44]. These solutions in yellow approximate a Pareto Front. The points from UNIO do not show up in the Front, meaning that neither the RMSD nor the energy aspects were better than FGMO or KDEO.

The results for the protein 2LVG with 40 residues are shown in Figure 8. Although FGMO obtained a large variance in the energy, it also got the smallest values found, while UNIO and KDEO had concentrated points. KDEO was the only method capable of finding RMSD values below 5.0 and reached an average better than UNIO and FGMO as well. The running time followed the same pattern as the 1A11 protein. KDEO was the slower among the three and FGMO was the faster.

In the scatterplot of Figure 8d, most of the left upper points in the Front belong to FGMO with smallest energy, and the left lower points belong to KDEO. This means FGMO was able to minimize better than KDEO, but KDEO found solutions closest to the configuration found in nature.

For protein 2KK7 with 52 residues (Figure 9), FGMO also had widely-spaced energy values. Considering only the average measure, FGMO would be the worst, but it was able to find the best energy value. It means that FGMO can reach better solutions than others, but for some reason, it is getting stuck at some local optimum, worsening the average value. Considering the RMSD aspect, KDEO was the only one able to find solutions with RMSD below 7.0. The running times required by UNIO and FGMO become closer due to fact
Figure 7: EDA for protein 1A11 with the three proposed methods: (a) energy plot; (b) RMSD plot; (c) time needed to run; and, (d) the scatter plot between energy and RMSD and the Pareto Front highlighted in yellow.
Figure 8: EDA for protein 2LVG with the three proposed methods: (a) density plot of the energy; (b) RMSD plot; (c) time needed to run; and, (d) the scatter plot between energy and RMSD and the Front highlighted in yellow.
that this protein is larger than 1A11 and 2LVG are. More points belonging to KDEO appear in the van der Waals energy per RMSD scatterplot. The top left points in the Front belong to FGMO. They had the best energy but an inferior RMSD.

Finally, the results for the largest protein used in the experiments, 2X43 with 67 residues follow the same trend as the previous proteins (Figure 10). Considering the energy aspect only, the UNIO and KDEO were, in average, better than FGMO is. However, Figure 10a shows that two outliers of FGMO are below $-410$, i.e. less than all other points from UNIO and KDEO. We believe that happened due to the sensitiveness of FGMO when setting the same number of mixture components throughout the evolutionary process, which somehow was benefited by the entirely evolutionary process. Apart from the KDEO running time, UNIO and FGMO had similar results. It seems that the running time of FGMO would be higher than UNIO for large proteins.

The scatterplot from Figure 10d shows a more defined agglomeration of points. KDEO produced two points in the Front. On the other hand, FGMO returned one point in the Front, although most of the points are around $-300$ of energy with RMSD between 15 and 25.

The Table 3 shows a summary of all experiments for all the four proteins evaluated. We can see that FGMO was the algorithm that got the best energy values; the KDEO the best protein structures; and UNIO the best running times. The KDEO also got the worse running time among all evaluated algorithms. However, it is interesting to see that a relatively complex algorithm as the proposed FGMO was faster than RW for most of proteins evaluated.

4.3. Comparison with the Native energy

We compared all non-bonded energies (van der Waals, electrostatic, solvation and hydrogen bonding) of each of the best predicted protein configuration against the native structure of each used protein. In order to compute the energy of native proteins, we converted the XYZ Cartesian coordinates from the PDB file into dihedrals and then used these values as input of one fitness function.
Figure 9: EDA for protein 2KK7 with the three proposed methods: (a) density plot of the energy; (b) RMSD plot; (c) time needed to run; and, (d) the scatter plot between energy and RMSD and the Pareto Front highlighted in yellow.
Figure 10: EDA for protein 2X43 with the three proposed methods: (a) density plot of the energy; (b) RMSD plot; (c) time needed to run; and, (d) the scatter plot between energy and RMSD and the Pareto Front highlighted in yellow.
Table 3: Best values out of 30 runs found for each protein and for each method. The columns Energy, RMSD and Time are not related to each other, meaning that the values in a row may come from different running. The best values are highlighted.

| Protein | Method | Energy (kcal/mol) | RMSD (Å) | Time (min.) |
|---------|--------|------------------|----------|-------------|
| 1a11    | RW     | -40.4            | 5.160    | 24.6        |
| 1a11    | MC     | -113.3           | 6.335    | 36.0        |
| 1a11    | GA     | -144.7           | 4.709    | 39.0        |
| 1a11    | DE     | -167.9           | 2.976    | 37.1        |
| 1a11    | UNIO   | -148.9           | 3.677    | 24.4        |
| 1a11    | KDEO   | -156.9           | 2.464    | 42.6        |
| 1a11    | FGMO   | -162.5           | 2.338    | 24.6        |
| 2lvg    | RW     | -59.7            | 7.981    | 52.8        |
| 2lvg    | MC     | -169.0           | 9.768    | 55.5        |
| 2lvg    | GA     | -211.8           | 9.153    | 61.1        |
| 2lvg    | DE     | -270.7           | 7.847    | 47.3        |
| 2lvg    | UNIO   | -233.0           | 5.605    | 34.4        |
| 2lvg    | KDEO   | -255.4           | 2.953    | 79.1        |
| 2lvg    | FGMO   | -268.8           | 5.593    | 42.7        |
| 2kk7    | RW     | -53.6            | 8.213    | 68.6        |
| 2kk7    | MC     | -192.3           | 11.959   | 64.7        |
| 2kk7    | GA     | -263.7           | 12.911   | 85.6        |
| 2kk7    | DE     | -301.4           | 9.391    | 49.0        |
| 2kk7    | UNIO   | -275.8           | 6.655    | 43.3        |
| 2kk7    | KDEO   | -324.6           | 4.676    | 78.2        |
| 2kk7    | FGMO   | -345.3           | 5.995    | 49.9        |
| 2x43    | RW     | -50.7            | 8.283    | 92.0        |
| 2x43    | MC     | -214.4           | 9.412    | 90.0        |
| 2x43    | GA     | -302.0           | 7.507    | 104.4       |
| 2x43    | DE     | -354.2           | 8.092    | 62.4        |
| 2x43    | UNIO   | -325.5           | 9.257    | 56.3        |
| 2x43    | KDEO   | -394.4           | 7.245    | 128.3       |
| 2x43    | FGMO   | -413.1           | 9.710    | 82.8        |

call of ProtPred.

Figure 11 shows a comparison between the configuration with the best energy value found by the proposed EDA and the native configuration. For all the four
proteins evaluated, the EDA managed to minimize the van der Waals energy more than native energy. This was expected because only van der Waals energy was minimized. However, in native proteins, the electrostatic energy (charge-charge) was better than the structure found by the proposed EDA. Thus, if we attempt to decrease the electrostatic energy of the protein configuration obtained by our EDA, the van der Waals energy would probably increase. The solvation and hydrogen bonding energies of the EDA were also higher than the native were, apart from the hydrogen bonding of protein 2X43. Therefore, a multi-objective algorithm would be a more appropriate strategy in order to deal correctly with all these energies as proposed by [9].
4.4. Comparison with other Evolutionary Algorithms

We showed that FGMO can produce more diversified solutions, yielding promising energy values. However, KDEO was able to find a better compromise between energy and RMSD. For this reason, we selected the KDEO to be compared against other heuristics from the literature. The parameters used with the other heuristics are defined in Table 4.

| Table 4: Parameters used with RW, MC, GA and DE. |
|-----------------------------------------------|
| RW   | MC | GA | DE |
| Pop. Size | 100 | 1000 | 2000 | 1000 |
| Sel. Pressure | 1.2 |
| Tournament |
| Step | 2 |
| Cross. Rate | 0.3 | 0.2 |
| Mutation Rate | 0.5 |
| Mutation Factor | 0.5 |
| Differential weight | 0.2 |

Figure 12 shows the scatterplot between van der Waals energy and RMSD for all proteins used in this work. As we expected, RW was the worst for all cases and MC was the second worst. Then, GA was worse than DE, although GA got some points mixed with DE for the two larger proteins used (2KK7 and 2X43). Finally, the KDEO was better for all cases considering only the RMSD.

For the smaller protein 1A11 (Figure 12a), DE was able to find better energy values than KDEO, although the KDEO got a better RMSD. Then, for the protein 2LVG (Figure 12b), DE also found better energy values. However, the best RMSD found by DE was 7.847 and most of the RMSD values found by KDEO are below 7.775. For protein 2KK7 (Figure 12c) the KDEO managed to find a better energy and RMSD than all other heuristics, while most of the RMSD of other heuristics are higher than 8. Finally, for protein 2X43 (Figure 12d) the KDEO also found better solutions with many points close to
-400 energy and RMSD close to 10. No other heuristics managed to appear in the Pareto Front. Therefore, it looks that the EDAs becomes more efficient as the protein size increases.

Each protein has its own set of parameters (population size, selection pressure etc) that works better, but in these experiments, we fixed the same parameters for all four proteins, so that we can also evaluate how the parameters would be for an unknown set of proteins without needing to calibrate them before.

Finally, we ran another experiment in order to show the strength of the EDA against other heuristics. We created a hypothetical chart that correlates the heuristic and the prior knowledge level needed to the heuristic be successful.
The vertical axis is ordered by the prior knowledge level, i.e. how much the search space needs to be reduced so that the heuristic can hit the correct solution. P1 means that the heuristic uses the whole search space when creating the initial population (as we have done in all previous experiments, where it ranges from \((-180, +180)\) in \(\phi\) and \(\psi\) directions). Then we subsequently split the search space using 5 degrees offset toward the right solution, in this case, the dihedral angles of protein 1A11 (look at Figure 6b and see that the middle of the red lines is \([\phi, \psi] = [-65, -35]\)). There were 49 divisions and all the levels are shown in Figure 13b.

Even a poor heuristic, as RW, managed to find the correct solution. However, the prior knowledge level needed by RW was so high that for any input it will give almost the same output. We denote this prior knowledge level as P5, since RW would fail for any larger search space, that is, this is the maximum search space we can use so that RW finds the correct solution. Using a slightly higher search space, MC managed to find the correct solution. We call this level P4 and RW has failed at this level. Next, GA found the correct solution with level P3. At this level, MC and RW have failed. In the next level, DE managed to find the right solution when other heuristics as GA, MC and RW have failed. We denote this level as P2. Finally, the EDA (KDEO) using no prior knowledge (using the whole range of the search space) managed to find the correct solution while all other heuristics have failed. That means the EDA was able to predict the correct protein configuration with no bias.

Furthermore, it is interesting to notice that the main difference between heuristics RW, MC, GA, DE and EDA (KDEO in this case) lies in the amount of information they extract from their populations in order to sample new solutions. That is, RW does not use any information from its population so the evolutionary process is completely random. MC uses information about one individual in order to sample new ones. GA uses information about two individuals to compose the new ones, and DE uses information from three individuals. Thus, we have 0, 1, 2, and 3 individuals that are being used for RW, MC, GA and DE, respectively, to sample new solutions. We believe that this is related
to the strength of the heuristics as they appear in literature. That means, in general, DE works better than GA, which works better than MC, which works better than RW.

Therefore, as we know, the amount of information EDAs can extract from promising individuals in order to generate the offspring is much higher than for DE. That happens because EDAs are designed to work with a set of selected individuals, which can have promising information to drive the evolutionary process toward promising solutions. This is why we believe EDAs are suitable for any type of combinatorial problem. Besides, when dealing with problems where we already know how variables could behave, it becomes more effective for an EDA to find the correct solution. This is what we have been doing with the PSP problem, associating the correlation between close variables together and creating probabilistic models from a set of promising individuals.

5. Conclusion

We develop three different probabilistic models for an Estimation of Distribution Algorithm specific for the ab initio Protein Structure Prediction with real-valued variables. We refer to these three methods as UNIO for the Univariate model-based Optimization, KDEO for the two-dimensional Kernel Density Estimation model-based Optimization and FGMO for the Finite Gaussian Mixtures model-based Optimization, yielding three different EDAs for PSP. The difference among these three EDAs is how they extract representative information from the selected individuals and use this information to sample the offspring.

The first, UNIO, is the simplest one. It is similar to UMDA but can handle problems with multiple modes instead of one, in an efficient manner. Secondly, knowing that dihedral angles $\phi$ and $\psi$ of amino acids have a strong correlation we modeled the pair of dihedral angles $[\phi, \psi]$ as correlated for our two-dimensional probabilistic models: KDEO and FGMO. Results have shown that accounting for correlation enables us to find better protein configurations in the search
Figure 13: Comparison of protein configurations between five different heuristics for protein 1A11: (a) chart showing the success and the failure according to the prior-knowledge level and, (b) the minimum search space needed for each heuristic to get success in their predictions.
In this work, only the van der Waals energy was used to compose the fitness function, so we can focus on the probabilistic model. Thus, we tried to use only $\alpha$-helices in our predictions where van der Waals energy alone might represent the atomic interactions well. For the largest protein used in the experiments, 2X43, the RMSD is still not as good as we expected. We believe that this may occur because van der Waals energy alone cannot handle correctly the loop between the two helices. On the other hand, for the three smaller proteins 1A11, 2LVG and 2KK7 the configurations we found were good enough, producing a low RMSD.

We also compared our EDA (more specifically, the KDEO) with other heuristics from literatures as Random Walk, Monte Carlo, Genetic Algorithm and Differential Evolution. We noticed that the number of solutions used to compose new ones constitute a critical step in designing a good optimization algorithm. For example, the weakest heuristic RW does not use any information about the evolutionary process to infer new solutions, so it is completely random. The MC (usually better than RW) uses information about one single solution from the original population in order to infer new ones. The GA (usually better than MC) makes use of two individuals from the population in order to compose the offspring. The DE (usually better than GA) uses information about three individuals to infer the new ones. Thus, we noticed a trend in the number of individuals used to compose the offspring and the strength of the optimization algorithm. This is why we believe EDAs are better and more efficient optimization techniques, since they use a set of promising solutions (the selected individuals) and try to extract some representative statistical knowledge from this set (the probabilistic model) in order to, finally, infer better solutions, leading the whole evolutionary process toward promising directions. We made this comparison with the EDA against RW, MC, GA and DE to show why an EDA can perform better than these previous algorithms. Actually, we are not taking the RW as a serious competitor against the EDA. We just wanted to emphasize the importance of using several potential solutions (the mechanism that the
EDAs do) to build the new ones, against other approaches that do not do this as illustrated in Figure 13a.

Until now, we had not seen an EDA for PSP with ab initio and full-atom representation approaches. So we decided to use some knowledge of the behavior of proteins, modeling the correlation $[\phi; \psi]$ and developing our specific EDA aimed at the PSP problem.

At the first glance, comparing the energy aspect between the two-dimensional probabilistic model-based techniques, KDEO and FGMO, it looks that KDEO is better in most of the cases. However, we noticed that the number of mixture components used in FGMO has a high effect on the energy values. Then, we still need to develop an automated and efficient way to estimate the ideal number of mixture components for each pair $[\phi; \psi]$. Although FGMO needs an iterative algorithm in order to estimate the parameters (EM algorithm), the computational efficiency stays close to UNIO. The main reason for this is because we are using the optimized PDF function (Equation 13 instead of Equation 9) and a small set of selected.

Besides possible computational and statistical improvements, still the fitness function could be improved by adding other non-covalent energies as solvation, hydrogen bonding and electrostatic energy to our experiments. However, before adding these energies one needs to ensure that such energies are as efficient as possible, as already achieved for van der Waals and solvation energies in previous works [45]. Otherwise, it would be the bottleneck of the whole algorithm.

We also know that the energy effects acting in proteins are contradictory and sometimes operate on different scales. Thus, the next step is to bring the Multi-Objective approach from the GA from [9] to our EDA.

Appendix A. Statistical analysis

We performed the pairwise Wilcoxon test in order to determine whether there is relevant difference between the evaluated methods. The comparison was performed for every combination of methods for all the four evaluated proteins.
(1A11, 2LVG, 2LKK7 and 2X43). The p-values that are greater than 0.05 are highlighted in bold and they show that there is no significance difference between them, according to the Wilcoxon test. Table A.5 shows the test for the van der Waals energy values. In this table, there is a single value in bold, indicating that, according to the Wilcoxon test, most results concerning van der Waals energy were different for all proteins evaluated. Table A.6 shows the test for the RMSD for the best protein configuration found over all runs for each method. Although the Wilcoxon test for RW-KDEO and DE-KDEO got a high p-value for almost all proteins, it is possible to see by looking at Figure 12 that KDEO was the only capable method of finding RMSD values below 7.5 for protein 2LVG (Figure 12b) and below 8.0 for protein 2KK7 (Figure 12c). Finally, Table A.7 shows the comparison between p-values for the running times. For the protein 2X43, the running time between MC-KDEO produced no significant difference; and for proteins 2LVG, 2KK7, 2X43 the running time between GA-KDEO also did not produce significant difference. That means that KDEO can be used for PSP at a similar computational cost as a MC or a GA.

Table A.5: P-value comparison with pairwise Wilcoxon test evaluating the van der Waals energy

|       | 1A11     | 2LVG     | 2KK7     | 2X43     |
|-------|----------|----------|----------|----------|
| RW-MC | 6.3e-10  | 5.9e-10  | 5.8e-10  | 5.9e-10  |
| RW-GA | 6.3e-10  | 5.9e-10  | 5.8e-10  | 5.9e-10  |
| RW-DE | 6.3e-10  | 5.9e-10  | 5.8e-10  | 5.9e-10  |
| RW-KDEO | 6.3e-10 | 5.9e-10  | 5.8e-10  | 5.9e-10  |
| MC-GA | 6.3e-10  | 5.9e-10  | 5.8e-10  | 5.9e-10  |
| MC-DE | 6.3e-10  | 5.9e-10  | 5.8e-10  | 5.9e-10  |
| MC-KDEO | 6.3e-10 | 9.5e-09  | 5.8e-10  | 5.9e-10  |
| GA-DE | 6.3e-10  | 5.9e-10  | 5.8e-10  | 1.6e-07  |
| GA-KDEO | 6.3e-10 | 7.3e-07  | 5.8e-10  | 5.9e-10  |
| DE-KDEO | 4.3e-09 | 3.1e-09  | 5.8e-10  | 5.9e-10  |
| UNIO-KDEO | 2.8e-07 | 2.7e-07  | 5.8e-10  | 5.9e-10  |
| UNIO-FGMO | 3.2e-06 | **2.0e-01** | 3.7e-04  | 1.5e-06  |
| KDEO-FGMO | 4.1e-02 | 3.8e-02  | 1.3e-04  | 4.2e-07  |
Table A.6: P-value comparison with pairwise Wilcoxon test evaluating the RMSD

|          | 1A11   | 2LVG   | 2KK7   | 2X43   |
|----------|--------|--------|--------|--------|
| RW-MC    | 7.3e-04| 5.4e-07| 2.1e-04| 1.4e-01|
| RW-GA    | 1.0e+00| 5.8e-05| 7.2e-07| 2.2e-03|
| RW-DE    | 1.0e+00| 1.9e-02| 9.4e-02| 1.0e+00|
| RW-KDEO  | 1.8e-02| 5.9e-01| 1.0e+00| 1.0e+00|
| MC-GA    | 9.3e-03| 1.0e+00| 8.6e-01| 1.4e-01|
| MC-DE    | 3.4e-03| 6.0e-04| 9.5e-02| 7.8e-03|
| MC-KDEO  | 1.3e-04| 4.9e-07| 1.2e-02| 2.8e-01|
| GA-DE    | 1.0e+00| 4.2e-02| 3.0e-03| 2.2e-04|
| GA-KDEO  | 8.4e-03| 1.7e-05| 1.8e-04| 4.5e-03|
| DE-KDEO  | 8.0e-02| 1.5e-03| 8.6e-01| 1.0e+00|
| UNIO-KDEO| 2.9e-02| 2.2e-01| 1.0e+00| 1.0e+00|
| UNIO-FGMO| 4.0e-02| 1.0e+00| 1.0e+00| 3.2e-03|
| KDEO-FGMO| 3.3e-03| 5.2e-02| 1.0e+00| 9.2e-03|

Table A.7: P-value comparison with pairwise Wilcoxon test evaluating the running time

|          | 1A11   | 2LVG   | 2KK7   | 2X43   |
|----------|--------|--------|--------|--------|
| RW-MC    | 3.6e-16| 3.6e-16| 2.9e-13| 1.1e-15|
| RW-GA    | 3.6e-16| 1.2e-11| 2.1e-09| 2.0e-08|
| RW-DE    | 3.6e-16| 8.8e-01| 7.9e-01| 1.0e+00|
| RW-KDEO  | 3.6e-16| 1.4e-13| 1.8e-06| 6.8e-16|
| MC-GA    | 6.3e-01| 5.8e-04| 7.1e-01| 1.0e+00|
| MC-DE    | 6.2e-01| 1.4e-07| 1.0e-09| 9.0e-14|
| MC-KDEO  | 8.8e-14| 4.4e-02| 5.4e-05| 2.8e-01|
| GA-DE    | 1.0e+00| 4.4e-02| 2.9e-07| 2.4e-08|
| GA-KDEO  | 2.7e-13| 2.4e-01| 5.8e-02| 1.0e+00|
| DE-KDEO  | 6.6e-12| 1.6e-03| 9.1e-04| 6.8e-16|
| UNIO-KDEO| 1.5e-12| 1.4e-11| 1.8e-07| 3.6e-16|
| UNIO-FGMO| 1.7e-06| 1.9e-05| 9.6e-02| 1.0e+00|
| KDEO-FGMO| 1.0e-15| 8.3e-12| 3.5e-10| 6.8e-16|
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