University of Warsaw Faculty of Physics

Maciej Dziubiński

Development of selected mesoscopic physical models with the aid of machine learning methods and their applications in studies of molecular systems.

Doctoral Thesis TODO: na kierunku Fizyka TODO: w zakresie Biofizyki

> Praca wykonana pod kierunkiem Prof. dr hab. Bogdana Lesynga Zaklad Biofizyki, Instytut Fizyki Doswiadczalnej, Wydzial Fizyki, Uniwersytet Warszawski

Warsaw, September 2015

Chapter 1

Introduction

The primary aim of this dissertation were methods of extracting relevant information from multi-dimensional data describing (bio)molecular systems. In Chapter II we present our novel method of discovering quasi-rigid parts in proteins, which can interpret experimental as well as simulation data. Chapter III presents a very different problem of identifying parts of a molecular system that propel (or hinder) a given structural transition. In both these cases we applied unsupervised machine learning techniques (in particular, *clustering*), which is a methodology commonly used in finding patterns in unstructured, and seemingly chaotic, data.

In this introductory chapter we give an overview of the molecular dynamics simulation scheme and the machine learning approach. The discussion here is broad, and was intended to motivate the application of clustering techniques described in the remainder of the dissertation. We go into more details concerning particular molecular dynamics and machine learning algorithms in Chapters II and III.

1.1 Molecular dynamics

Molecular dynamics (MD) are a wide range of numerical methods designed for the study of molecular systems. Typically, the MD approach assumes a classical potential energy function U also known as the force field, which treats atoms as electrostatically charged points, and chemical bonds as springs and hinges. The arguments of the function U are configurations \mathbf{q} of a particular system, with values $U(\mathbf{q})$ emulating the potential energy of the system at that configuration \mathbf{q} . If we denote the set of all configurations of a given system by Ω , then $U: \Omega \to \mathbb{R}$. The force acting on atom i is then derived from U by taking a gradient with respect to position \mathbf{q}_i of that atom:

$$\mathbf{F}_i(\mathbf{q}) = -\nabla_i U(\mathbf{q}),$$

which requires that U is differentiable.

More accurate algorithms take into account quantum effects, making the simulation more reliable, but also immensely expensive in terms of computational time. However, although modeling intricate properties of molecular systems using a classical potential may seem an over-simplification, MD has achieved considerable success over the recent years, and is constantly improving.

Classical MD simulations are popular mainly because of their significantly lower computational cost than their quantum-mechanical counterparts. This allows for estimation of free energy profiles of structural transitions, even for large, biologically-relevant systems (such as enzymes, DNA, membrane systems, and other).

Free energy translates into experimentally observable properties. For example: the affinity of a ligand and an enzyme, or the chance of transferring an ion through a membrane channel. Free energy profiles offer insight into the process at hand, but also allow for utilizing feedback from experimental data to refine the modeling potential.

To run an MD simulations, one needs to choose the right force field, but also – and perhaps more importantly – the appropriate *sampling technique*. The underlying, well-established premise is that molecular systems in thermal equilibrium experience random impulses of momenta, which can be though of as a model of collisions with molecules of the surrounding world. In other words, the total energy of a system in thermal equilibrium is not conserved, but rather fluctuates around a certain mean value.

Consequently, configurations of a system are characterized by a probability density, which in the case of thermal equilibrium at T = const is the well-known $Boltzmann\ distribution$:

$$\rho_B(\mathbf{q}) = Z^{-1} \exp[-U(\mathbf{q})/k_B T], \tag{1.1}$$

where k_B is the Boltzmann constant, and Z is a normalizing factor of great importance, often referred to as the partition function, and is given by:

$$Z = \int_{\Omega} \exp[-U(\mathbf{q})/k_B T] d\mathbf{q}.$$

From this probabilistic perspective we can view an MD simulation as a sampling procedure, during which we *should* acquire configurations according to the Boltzmann distribution. Macroscopic properties of the system (affinity of a ligand to an enzyme, for example) are defined as expected values of appropriate microscopic quantities¹.

1.1.1 Sampling techniques and the potential of mean force

Underlying the simplistic, classical assumptions embedded in MD simulations are complex sampling techniques, used for estimating macroscopic properties of microscopic systems. MD simulations can, for example, mimic time evolution of a

$$S = -k_B \int \rho_B(\mathbf{x}) \log \rho_B(\mathbf{x}) d\mathbf{x}.$$

¹Or the probability density itself, as, for example, in the case of entropy,

system with the use of integration schemes such as the *Langevin dynamics*, which numerically integrates the following stochastic differential equation:

$$\mathbb{M}\ddot{\mathbf{q}} = -\nabla U(\mathbf{q}) - \gamma \mathbb{M}\dot{\mathbf{q}} + \sqrt{2\gamma k_B T \mathbb{M}} \mathbf{R}(t)$$
(1.2)

where M is a diagonal matrix of atom masses, and $\mathbf{R}(t)$ is a stationary Gaussian process with zero mean, and auto-correlation expressed by the delta function: $\langle \mathbf{R}(t)\mathbf{R}(t')\rangle = \delta(t-t')$. It is important to note that although the output of a Langevin dynamics simulation is a trajectory, most sampling techniques avoid modeling the time evolution of a system.

Potential of mean force

Vast samples of multidimensional points are impossible to interpret, unless we utilize some simplifying technique. Probably the most popular representation of a change or transformation occurring in a system is the so-called *collective variable*, along with the concept of the *potential of mean force* (PMF). A collective variable ξ is a function of the system's configuration \mathbf{q} , defined so that its value can be readily translated into the progress of a particular transformation. The PMF associated with a collective variable ξ , for a particular value ξ^* of that collective variable, is defined as follows:

$$A(\xi^*) := -k_B T \log \rho(\xi^*),$$

where ρ is a probability density of the configuration \mathbf{q} , such that $\xi(\mathbf{q}) = \xi^*$.

While the collective variable ξ serves as a means of reducing the dimensionality of a complex structural transition, the PMF provides a probabilistic interpretation of the process. Through the PMF we gain insight about the transformation's bottle-necks and ranges of values of the collective variable, which corresponding to the system's meta-stable states.

1.2 Unsupervised machine learning

Unsupervised machine learning is a class of numerical methods for extracting information from unstructured, multi-dimensional data. Roughly, these methods amount to reducing the dimensionality of the data. Stated differently, unsupervised machine learning tries to address the following questions²:

- Is there a clear way of visualizing the data?
- Can we partition the set into cohesive subgroups, distinctive from each other?

The collective variable discussed above was a method of dimensionality reduction, but one based on a probabilistic model derived from physical postulates.

²These are not the only questions taken up by unsupervised machine learning. Most notably, one of the more interesting, and fast-changing fields is the problem of *community detection*, which can be thought of as a more general problem than clustering (discussed further).

Data produced by MD simulations of biomolecules are multi-dimensional trajectories, points $\{\mathbf{q}_t\}_{t=1}^S$, where $\mathbf{q}_t \in \mathbb{R}^{3n}$ (S is the number of steps of the simulation, and n – the number of atoms of the system). Depending on the sampling technique **adapted** in the MD simulation, the t parameter may or may not be associated with time, as, for example, in the case of Monte Carlo sampling. Analogously to MD, configurations acquired from NMR experiments can be also thought of as a set of multi-dimensional points, $\mathbf{q}_t \in \mathbb{R}^{3n}$, in which the order imposed by the t parameter has very little to do with time.

Regardless of the source, we are faced with the problem of purifying the information hidden within our data. Unsupervised machine learning offers us two methods of simplifying the data: *principal component analysis* (for visualization and preprocessing) and *clustering* (for identifying distinctive subgroups of points).

Clustering is the task of finding distinct subsets (*clusters*) of a given set of objects, which share a high similarity within a given cluster, but low similarity across clusters. Inherently, the task requires that a measure of similarity between objects is given, or that a natural measure can be proposed. In fact, the similarity function determines the meaning behind any clustering, and often predetermines which clustering algorithms can and cannot be utilized in a given problem.

Once we define a similarity function, we are left with choosing the right clustering algorithm.

1.2.1 Adjacency matrix representation

It is important to introduce at this point the matrix representation of a set of objects and their similarity. If we enumerate the objects in our set using natural numbers 1, 2, 3, ..., N, and denote the similarity between objects i and j by w_{ij} , we can represent the relations between the objects using a matrix $W \in \mathbb{R}^{N \times N}$. Matrix W is referred to as the similarity (or: adjacency) matrix.

1.2.2 Dynamic domains as clusters

Small, globular proteins are often thought of as fairly rigid biomolecules. However, the set of low-energy, accessible configurations grows rapidly as the number of atoms in a system increases. Consequently, large molecular systems can undergo significant structural changes, involving coordinated, multi-step transitions. Due to immense complexity of these systems, such phenomena are difficult to describe, let alone to interpret.

In Chapter II we introduce a method of simplifying these incomprehensible transitions using a clustering scheme. The main idea of our method is to find parts of the protein which – although in movement with respect to each other – remain internally rigid. Or, more accurately, quasi-rigid, because small variations in distances between amino acids are omnipresent. These quasi-rigid parts of the protein are referred to in the literature as dynamic domains [3, 29, 31].

In our method of dynamic domains identification we adapted a clustering framework. We proposed a measure of strength of a contact between amino acids, and apply an appropriate clustering algorithm to identify quasi-rigid parts of a protein. The strength of a contact depends on structural variability of a given pair of residues. Having a set of objects (amino acids) and a "similarity measure" (contact strength between residues), we construct an adjacency matrix. This matrix is used as input for a particular, well-suited clustering procedure, one of the class of *spectral algorithms*, which is centered around finding eigenvectors of a matrix closely related to the adjacency matrix.

1.2.3 Molecular cogs as clusters

In Chapter III we present our newly-developed method of identifying *molecular* cogs – subsystems of a molecule which, for a given structural transformation, drive the transition forward or backwards. This is another application of the cluster analysis, although quite dissimilar to the one discussed in Chapter II.

For the purpose of identifying molecular cogs we proposed a "similarity function" between pairs of atoms, which indeed quantifies the contributions of that pair the the free energy change along a collective variable. With this, we again construct adjacency matrices, and discover clusters which correspond to groups of atoms sharing a common contribution to the free energy (positive or negative). However, the method of dynamic domains and molecular cogs identification differ not only at the stage of adjacency matrix construction, but also in the choice of the clustering algorithm.

Chapter 2

ResiCon: a method for the identification of dynamic domains, hinges and interfacial regions in proteins

2.1 Abstract

Structure of most proteins is flexible. Indentification and analysis of intramolecular motions is a complex problem. Breaking a structure into relatively rigid parts, so-called dynamic domains, may help comprehend the complexity of protein's mobility. We propose a new approach called ResiCon (Residue Contacts analysis), which performs this task by applying a data-mining analysis of an ensemble of protein configurations and recognizes dynamic domains, hinges and interfacial regions, by considering contacts between residues. Dynamic domains found by ResiCon are more compact than those identified by two other popular methods: PiSQRD and GeoStaS. The current analysis was carried out using a known reference set of 30 NMR protein structures, as well as MD simulation data of flap opening events in HIV-1 protease. The more detailed analysis of HIV-1 protease dataset shows that ResiCon identified dynamic domains involved in structural changes of functional importance. The ResiCon server is available at URL:

http://dworkowa.imdik.pan.pl/EP/ResiCon

2.2 Introduction

Proteins are not static. Nuclear magnetic resonance (NMR) spectroscopy ([37]) and the spin-echo spectroscopy ([9]) experiments show that. In several cases it was proven that flexibility may be crucial to protein functionality ([19], [27]). Although experimental methods provide only general clues about intramolecular motions, molecular dynamics (MD) simulations extend their reach by giving a higher resolu-

tion picture – both in space and time – of protein mobility. By studying an NMR ensemble or MD trajectory one may notice that it is composed of relatively rigid structural parts, often referred to as *dynamic domains* ([30]).

Domains in traditional sense are regarded as parts of the protein which are: conserved (in terms of evolution), autonomous (in terms of folding), and/or compact (in terms of tertiary structure). Such "static" domains are identified through sequence homology, structural analysis of a single configuration, or both. (For conventional methods of identifying protein domains based on multiple sequences or a single structure see e.g. ([43], [7]).) Conversely, dynamic domains depend on structural transitions performed by the protein.

A number of methods for identification of dynamic domains have been developed. The simplest procedures are based on normal mode analysis, which assumes a harmonic approximation of the potential energy function ([31]). More advanced approaches use the Gaussian Network Model and analyze correlations in motions between the residues of the protein ([57]). Many other approaches have also been developed ([3], [55], [5], [22], [41]), but all of them anticipate dynamic domains by analyzing a single structure of a protein. Another class of methods for dynamic domains assignment requires exactly two configurations (see [29], [36], [56]). However, the assumption that two "representative" structures encompass all relevant motions is rather speculative.

Experimental and in silico methods reach beyond single-structure representation, and are capable of producing numerous configurations of a given protein. Rather than inferring dynamic domains from one or two structures, a more natural approach would be to determine them based on an ensemble of configurations. GeoStaS is the only method known to us that analyzes a whole ensemble of configurations and assigns each residue to a dynamic domain ([44]). Although GeoStaS can analyze not only proteins but also nucleic acids, it fails to discover dynamic domains whenever they rotate with respect to each other. Alternative methods of analyzing ensembles of configurations assign residues to a static "core" or unstructured bundle (see [49], [33]).

The purpose of this study was to develop a novel methodology named ResiCon, capable of extracting dynamic domains from an ensemble of protein's configurations. ResiCon analyzes strengths of contacts between residues based exclusively on geometrical changes occuring in the provided set of structures. The set may be an NMR ensemble of configurations, or snapshots produced in the course of an MD simulation. ResiCon's main functionality is to identify dynamic domains, but it can also find hinges and interfacial (interdomain) regions.

2.3 Approach

ResiCon starts with identifying pairs of residues which are *in contact*. There are several definitions of contacts between amino-acid residues in the literature. We used the definition presented in ([16]) and adapted it to the case when more than one structure is given (see also [17]).

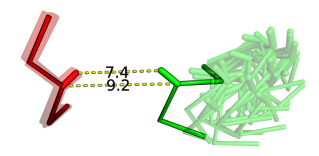


Figure 2.1: For each configuration in an ensemble, pairs of elements are constructed. In this picture residues are in contact because there exists a configuration for which the condition (2.1) is satisfied (the distance $d_C = 7.4\text{Å}$, and $d_{\alpha} - d_C = 1.8\text{Å}$).

Next, ResiCon constructs a virtual scaffold, connecting residues which are in contact with bars. Stiffness of a given bar reflects the estimated strength of the corresponding contact.

Finally, to identify dynamic domains, ResiCon carries out a partitioning (by computing minimal cuts) of the scaffold, cutting weaker and preserving stiffer bars. This partitioning is carried out by applying a spectral clustering algorithm presented in the following section.

The fundamental underlying assumption is that stability of rigid parts results from stable interactions between its residues. However, in our approach we do not analyze physical interactions between residues – they may be hydrophobic, electrostatic or significant in some other way. We simply assume that the measure of strength of a contact between residues is reflected by their geometrical variability across a given sample of structures.

2.4 Methods

Throughout this paper we use terms: model, configuration, structure and conformation interchangeably. We will refer to a set of structures acquired from an MD trajectory or NMR experiment as the ensemble of configurations or simply: an ensemble. Let us denote the number of structures in an ensemble by S.

Residue contact

For each pair of residues in every model we compute distances between C_{α} atoms (d_{α}) and between geometrical centers of side-chains R_{C} (d_{C}) (for glycine $R_{C} = C_{\alpha}$, and for alanine $R_{C} = C_{\beta}$). We say that two residues are *in contact*, if at least one configuration in the ensemble satisfies the condition:

$$(d_{\alpha} \le 6.5\text{Å})$$
 or $(d_{C} \le 8\text{Å} \text{ and } d_{\alpha} - d_{C} \ge 0.75\text{Å})$ (2.1)

Threshold values are the same as in definition of contact presented in ([16]) and relate to the range of distances in which physical interactions between residues

occur. The second sub-condition favours residues whose side-chains point towards each other (see Figure 2.1). Residues that are sequential neighbors are not taken into account.

We assign a quantitative value to the strength of a contact in terms of geometrical variability of the structural part associated with that contact. Such measure is required to capture not only changes in d_{α} , but also rotational shifts and alterations in the backbone in the vicinity of both residues that are in contact. To do so, we constructed structural parts, comprising sequential neighbors of the two residues in contact. ResiCon assigns a numerical value to the geometric variability by using the least root mean square deviation (RMSD) ([32]). Before elaborating on the details, we proceed with the following definitions.

Elements.

An element is a structural part of a protein centered around a given residue. It comprises five points, corresponding to the positions of the C_{α} atoms of the central residue, and its four sequential neighbors (two preceding and two following). For each model s in an ensemble, and each residue i, an element – denoted by E_i^s – is constructed. Residues for which an element cannot be built (e.g. N- and C-termini) are omitted.

Geometrical variability.

We consider pairs of elements, $E_{ij}^s := (E_i^s, E_j^s)$, and express structural deviation of a contact between two configurations r and s by RMSD of E_{ij}^r and E_{ij}^s . We use the following function to express the strength of a contact in terms of the whole ensemble:

$$G(i,j) := \max_{\text{pairs of states } (r,s)} \text{RMSD}(E_{ij}^r, E_{ij}^s).$$

The smaller the geometric variability, the stronger the contact.

We tested several statistics based on RMSDs of pairs of elements. This particular definition of G assumes that a strong contact is not "broken" in any pair of models. Conversely, a contact whose structural stability is breached at least once is assumed to be weak and not contributing to the stability of a given dynamic domain.

Note that conformational transitions may be rapid or insufficiently sampled. Therefore, defining geometrical variability in terms of some averaging statistic (e.g. mean, median) might lead to omitting significant structural changes occurring in a protein.

Contact matrix

We now describe a matrix representation of an edge-weighted graph, in which nodes correspond to residues. Because we used a spectral clustering algorithm which required that weights in the graph were in the interval [0,1], we needed to

renormalize the geometrical variability. We calculated the weight between node i and j using the following *contact function*:

$$D(i,j) := \begin{cases} 1 & |i-j| \leq 1 \\ L_{\alpha,\beta}(G(i,j)) & \text{residues } i \text{ and } j \text{ are in contact} \\ 0 & \text{otherwise} \end{cases}$$

where $L_{\alpha,\beta}$:

$$L_{\alpha,\beta}(x) := \left(1 + e^{\frac{x-\alpha}{\beta}}\right)^{-1}$$

is a logistic function. We refer to matrix $[D(i,j)]_{ij}$ as the contact matrix.

Parameters $\beta > 0$ and $\alpha > 0$ allow for the customization of ResiCon. The logistic function, $L_{\alpha,\beta}$, can be thought of as a rescaling transformation for the measure of geometrical variability, G. It has a simple interpretation – values of G exceeding α are smoothly cut-off, where the degree of "smoothness" is determined by β . All results presented in this paper were acquired with default values of α and β :

$$\beta_{\text{default}} := \frac{1}{\sigma(G)}$$
 $\alpha_{\text{default}} := \mu(G)$ (2.2)

where the σ and μ stand for standard deviation and mean taken over values of G for all pairs of elements.

Another feature of the logistic function becomes apparent if we consider an ensemble composed of (nearly) identical structures. This has two possible interpretations: the protein is very stable and no conformational changes exist, or that the provided ensemble does not represent such changes. Thus, ResiCon will assume that the contacts are strong – nearly as strong as the peptide bonds $(G \approx 0 \Rightarrow L_{\alpha,\beta} \approx 1)$. Consequently, the contact matrix becomes a so-called *contact map*, assigning binary values to pairs of residues (i.e. 1 if a contact occurred at least once, and 0 otherwise).

Clustering

The contact matrix may be treated as a *similarity matrix*, denoted W, and be used as input in a clustering procedure. Thus, we consider residues to be similar if they are likely to belong to the same dynamic domain. The identified clusters would then correspond to quasi-rigid structural parts.

The choice of a clustering algorithm is not a trivial task and for the identification of dynamic domains two crucial requirements need to be met. Firstly, contact matrices for various proteins vary in dimension and density and the clustering algorithm needs to perform well regardless of these variabilities. Secondly, the algorithm should facilitate an automated method of choosing the optimal number of clusters.

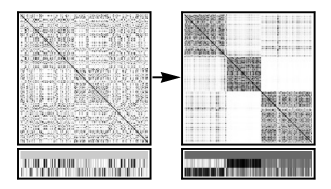


Figure 2.2: An example of a similarity matrix W and three eigenvectors of the stochastic matrix $D^{-1}W$ with two different orderings.

Agglomerative hierarchical clustering algorithms are one of the most popular approaches to clustering ([28]). They follow a greedy scheme to construct a dendrogram encoding distances between clusters. This dendrogram can be cut at a certain height, which determines the number of clusters. If the height parameter could be set so that for all similarity matrices we would obtain high-quality clusters, the hierarchical clustering would have been a good candidate for a clustering procedure. However, as we explain in the *Supplementary Materials*, estimation of this parameter is difficult, and to determine the number of clusters we would need to extend the conventional hierarchical clustering with a measure of cluster quality.

This was one of the reasons we have chosen a spectral clustering algorithm, which has an inherent indicator of a partitioning's quality.

Spectral clustering.

Clustering algorithms based on finding the eigensystem of the similarity matrix (or more often a matrix derived from it) are termed *spectral algorithms*. They perform a clustering by minimizing the cost of cutting a graph into subgraphs, which agrees with our intuition about finding quasi-rigid parts based on a contact matrix. Optimal clustering is achieved by discarding contacts with the lowest total weight (as few and as weak as possible) to achieve a partitioning into unconnected regions.

In the case of the clustering algorithm used in ResiCon optimal partitioning is decoded from eigenvectors corresponding to the largest eigenvalues of a stochastic matrix $D^{-1}W$, where $D:=diag(d(1),\ldots,d(n))$ and $d(i):=\sum_{j=1,j\neq i}^n w(i,j)$ (see [53]). This transformation of the similarity matrix ensures that the identified clusters tend to have similar sizes, which prevents from identifying singular nodes as clusters. Spectral algorithms make no assumptions on the shape of clusters, and, in contrast to the greedy procedures, are insensitive to the ordering of vertices.

In Figure 2.2 we present a 150×150 similarity matrix with rows and columns ordered in two different ways. The ordering on the right is dictated by the spectral clustering – elements 1–50 were assigned to the first cluster, 51–90 to the second,

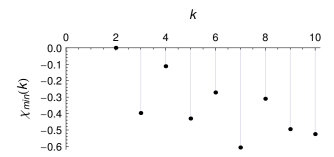


Figure 2.3: Values of the χ_{\min} indicator for k = 2, ..., 10, computed for HIV-1 protease. Because $\chi_{\min}(4)$ is closer to 0 than $\chi_{\min}(6)$, the optimal number of clusters is 4.

and 91–150 to the third. To find these three clusters we use the first three eigenvectors (corresponding to three largest eigenvalues) of the stochastic matrix $D^{-1}W$. The first eigenvector always has a constant value in all positions and corresponds to a trivial clustering into a single group. The second eigenvector encodes a partitioning into two groups: nodes 51–90 in the first, and the remaining nodes in the second group. The third eigenvector allows to discern the third cluster, composed of nodes 1–50.

Clustering algorithm.

We used a spectral clustering algorithm in which partitioning of a graph is expressed in terms of a membership matrix χ . A short description of the procedure is presented below, for details refer to ([53]).

Let Y denote the matrix containing k eigenvectors of the $D^{-1}W$ stochastic matrix, corresponding to k largest eigenvalues. The procedure computes a linear mapping \mathcal{A} from the eigenvectors Y to the membership matrix:

$$AY = \chi$$

The element χ_{ij} of this matrix represents the membership of the i^{th} node in the j^{th} cluster. Therefore, if n is the number of nodes in the graph and k is the number of clusters, then $\chi \in \mathbb{R}^{n \times k}$.

This algorithm has two important features:

- it computes the membership matrix χ which allows for overlapping clusters,
- it offers an indicator, called χ_{\min} , used to determine the optimal number of clusters.

Number of clusters.

ResiCon first checks if k = 1. To do so, a partitioning into two clusters is carried out. The spectral algorithm presented above finds an optimal cut ([53]) leading to

clusters A and B (two sets of indices, that correspond to nodes). We express the cost of such cut by:

$$f := \frac{\sum_{i \in A} \sum_{j \in B} w_{ij}}{\left(\sum_{k,l \in A} w_{kl}\right) \left(\sum_{k,l \in B} w_{kl}\right)}$$

where w_{ij} is the weight of the edge between nodes i and j. The validity of a clustering into clusters A and B was checked by asserting that its cost is less than a given threshold. If the criterion was met, we assumed that a clustering into two or more clusters existed. Default threshold for f used to produce all results presented in this study was 0.1. Above this value we observed that compact static proteins were partitioned into short (less than four residues long) segments, which we regarded as improper dynamic domains. On the other hand, lower values resulted in a single dynamic domain assignment in several cases where two domains were apparent.

If k > 1, the optimal number of clusters is determined with the use of the indicator presented in ([53]). Here, we give a short overview of the properties of χ_{\min} and propose a simple procedure for computing the optimal number of clusters. The indicator $\chi_{\min}(k)$ is defined as the minimal element of the membership matrix χ found by partitioning it into k clusters.

In the case of k=2 the indicator is always zero, $\chi_{\min}(2)=0$. For k>2 the indicator is less than zero, however if $\chi_{\min}(k)\approx 0$, the clustering into k clusters is the optimal one (Figure 2.3). Let us recall the notion of visualizing a clustering by a block-like similarity matrix. Roughly speaking, the value of $\chi_{\min}(k)$ resembles the deviation of the similarity matrix from the "pure" block-like form. However, it is difficult to decide which values of χ_{\min} are sufficiently close to zero to indicate the optimal number of clusters. Therefore, the problem at hand is: does the optimal number of clusters equal two, or more?

In our first approach, the optimal number of clusters was chosen based on a threshold – the optimal k was the one for which χ_{\min} was above a certain value. However, it was difficult to find the right threshold because values of χ_{\min} strongly depend on the number of nodes in the graph. Therefore, the following procedure was adapted in ResiCon:

- 1. Determine the cost of the optimal cut. If it exceeds the 0.1 threshold, the optimal number of clusters is k=1. Otherwise, assume that k>1 and continue the procedure.
- 2. Compute the values of χ_{\min} for partitionings into $3, 4, \ldots, M$ clusters.
- 3. Find $k_1 > 2$ for which χ_{\min} is closest to 0, and $k_2 > 2$ for which χ_{\min} is the second closest to 0.
- 4. If $\chi_{\min}(k_1) > 0.5 \chi_{\min}(k_2)$, then the number of clusters is $k = k_1$. Otherwise, k = 2.

The 0.5 constant in the fourth point means that we choose k_1 as the number of clusters, if $\chi_{\min}(k_1)$ is closer to 0 then to the next best χ_{\min} (i.e. $\chi_{\min}(k_2)$). The

maximal number of clusters M is set to 10 by default, but the user can specify a different number. All results presented in this paper were computed with M = 10.

In other words, ResiCon chooses k > 2 for which the indicator χ_{\min} is "relatively close" to 0. When no such value exists, a partitioning into two clusters is assumed to be optimal.

Hinges and interfacial regions

Hinges.

We define hinges as parts of the structure satisfying both of the following conditions:

- 1. they do not belong conclusively (in terms of membership as explained below) to any dynamic domain,
- 2. they are sequentially located between dynamic domains.

The first condition is tested in terms of membership: if a residue membership in any cluster does not exceed certain threshold χ_{hinge} it may belong to a hinge. The default value of the parameter is 0.65, but the user can specify a different value.

Interfacial regions.

A residue is assumed to compose an interfacial region if two conditions are met:

- 1. it does not belong to any hinge,
- 2. it was in contact with a residue that does not belong to the same dynamic domain at least once.

Results comparison

According to our knowledge, no expert curated database of dynamic domains exists. Also, we are not aware of any quality measure for the dynamic domains assignment. Therefore, we compared different methods by analyzing agreement between their results.

We used the measure presented in ([38]) called Variation of Information (VI) to analyze the results compatibility. It has the advantage of being a metric in the space of all partitionings of a given dataset. The downside of VI is that its values do not lie in a fixed interval (e.g. [0,1]), but instead have an upper bound that depends on the size of data. In our case data size equals the number of residues in a given protein. This means that values of VI for partitionings of one protein are not directly comparable to the values acquired for partitionings of another protein, with a different number of residues. Nonetheless, when considering a particular protein, the VI metric quantifies the agreement between different assignments of dynamic domains. Here we give an outline of the method, for details see ([38]).

Let us denote a clustering by C. It is composed of clusters – mutually disjoint subsets C_1, \ldots, C_k . That is, $C = \{C_1, \ldots, C_k\}$ such that $C_i \cap C_j = \emptyset$ for all pairs

i, j. Assume that the numbers of points in consecutive clusters are n_1, \ldots, n_k . Then, the probability that a random point from the dataset belongs to the ith cluster is

 $P(i) := \frac{n_i}{n},$

where n is the number of all points in the set. Note that $\sum_{i=1}^k n_i = n$. Analogously, let another clustering of the same set of points $\mathcal{C}' = \{C'_1, \ldots, C'_{k'}\}$ be composed of clusters with $n'_1, \ldots, n'_{k'}$ points. By $n_{ij'}$ we will denote the number of points assigned to cluster i in clustering \mathcal{C} and cluster j' in \mathcal{C}' . Then,

$$P(i,j') := \frac{n_{ij'}}{n}$$

is the probability of randomly choosing a point that belongs to both clusters.

The \mathcal{VI} measure is defined in terms of entropy and joint entropy of the probability distributions defined above. That is, if the entropy of clustering \mathcal{C} is expressed by:

$$H(\mathcal{C}) := -\sum_{i=1}^{k} P(i) \log_2 P(i),$$

and the joint entropy of two clusterings is given by:

$$H(\mathcal{C}, \mathcal{C}') := -\sum_{i=1}^{k} \sum_{j'=1}^{k'} P(i, j') \log_2 P(i, j'),$$

then the variation of information of the two clusterings is defined as:

$$\mathcal{VI}(\mathcal{C}, \mathcal{C}') := 2H(\mathcal{C}, \mathcal{C}') - H(\mathcal{C}) - H(\mathcal{C}')$$

Quality of dynamic domains

We consider dynamic domains to be structural parts of the protein, which move with respect to each other, but remain internally rigid. In order to assess which method for dynamic domains identification is better, a measure was required that would quantify the quality of a given assignment. We did not find such a scoring function in the literature, and propose the following geometrical measure called total geometrical variability:

$$Q := \sum_{i=1}^{k} \max_{\text{pairs of states } (r,s)} \text{RMSD}(D_i^r, D_i^s),$$

where D_i^s is the set of C_{α} atoms comprising the domain D_i in the state s, and k is the number of domains. Smaller values of Q indicate higher quality.

Note that typically, if a domain is structurally rigid, the maximal RMSD for that domain is smaller than the sum of maximal RMSDs of its two subsets. Therefore the proposed measure favors large, compact domains. It is also worth noting that for a trivial dynamic domain (single residue) the RMSD is undefined. We set its value to 0, although this artificially reduces Q (see $Quality\ analysis$).

	R	G	P1	P2
R	0	1.47	3.29	0.21
G	 1.47	0	3.76	1.51
P1	3.29	3.76	0	3.29
P2	0.21	1.51	3.29	0

Figure 2.4: Values of VI for partitionings of the 1d1d protein molecule. A pair of results produced by PiSQRD which had the highest VI are denoted by P1 and P2.

2.5 Results and discussion

We compared ResiCon with two recent methods: GeoStaS ([44]) and PiSQRD ([41]). The latter method represents the class of methods which identify dynamic domains by analyzing a single structure. We used a test set comprising 30 NMR-resolved protein structures exhibiting significant mobility, which was previously used in ([49]), ([33]), and in ([44]). The set was initially proposed to examine the efficacy of a method of identifying structurally-stable cores in flexible proteins. These structures often contain a single rigid core with significant geometrical distortions present in peripheral regions (as indicated in [49] and also observable in ResiCon's results).

We have also used ResiCon to analyze a canonical test case – the HIV-1 protease molecule – using an MD trajectory computed with a coarse-grained force field RedMD ([23, 24]) as input data. This example shows that ResiCon is capable of finding dynamic domains of a protein whose functionality depends on flexibility and mobility of its rigid parts.

We also explain the so-called *zebra effect* – a peculiar result produced by Resi-Con, observed in several cases. This effect is especially strong when a suboptimal numer of clusters is chosen.

Comparative analysis

ResiCon and GeoStaS are both designed to work on an ensemble of structures, and produce a single partitioning into dynamic domains. Both methods impose no assumptions about sampling and order of provided conformations. ResiCon uses maximal local distortions computed over all pairs of frames, indifferent to over- or undersampling of configurations, as long as they are present in the ensemble.

However, the PiSQRD server by default analyzes a single structure and estimates the so-called *low-energy modes*, which are the eigenvectors of the structural covariance matrix (under the canonical ensemble, i.e. assuming Boltzmann distribution of configurations). These low-energy modes are assumed to carry the information relevant to dynamic domains identification. We observed that the choice of an input structure influences the results significantly, but there is no definite criterion for choosing the *right* structure for the analysis. The PiSQRD server provided with a PDB file containing NMR models by default finds dynamic domains for the first model.

This might introduce a bias unfavorable for PiSQRD's performance. We, there-

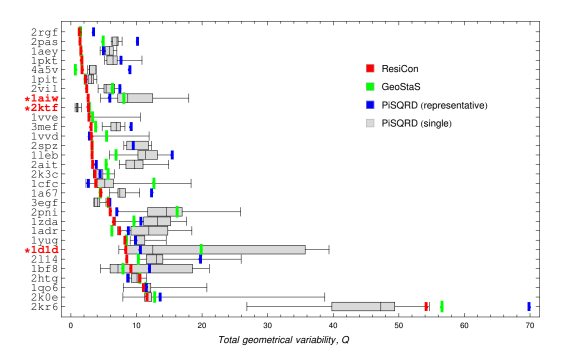


Figure 2.5: Box and whiskers plot of the dynamic domains quality score Q for ResiCon, GeoStaS, PiSQRD. In blue are Q values for PiSQRD's dynamic domains determined from structural covariance matrices (see Supplementary Materials).

fore, decided to examine results produced by PiSQRD for every structure in the ensemble in order to compare ResiCon against its full capacity. The best dynamic domains were chosen and presented together with the results produced by ResiCon and GeoStaS in Table 2.1.

PiSQRD also gives a possibility of providing a set of low-energy modes extracted from a structural covariance matrix estimated from a set of structures, but this procedure is not straightforward and requires additional assumptions (see *Supplementary Materials*). We scrutinize the quality of dynamic domains found by PiSQRD from user-provided low-energy modes in section *Quality analysis*.

Because of a large number of models, it was not possible to provide a graphical representation for each partitioning. We therefore focused on values of the agreement measure \mathcal{VI} between results produced by the three methods.

To familiarize the Reader with values of the measure \mathcal{VI} , we take a look at two most distant partitionings produced by PiSQRD (denoted by P1 and P2) for an exemplary 1d1d protein, and how they relate to the results given by ResiCon (R) and GeoStaS (G) (Figure 2.4). The clustering P2 seems to be very similar to the one produced by ResiCon, while P1 gives a sliced-and-diced picture of the protein's mobility. It seems that a partitioning produced by PiSQRD depends on physical properties embedded in a particular configuration. Although NMR models carrying information about dynamic domains may exist, others lead to chaotic partitionings. In order to produce a single clustering, PiSQRD would need a procedure

for interpreting physical properties based on an ensemble of configurations.

Nearly 50% of the assignments produced by PiSQRD for 1d1d are coherent with the clustering given by ResiCon (see Supplementary Materials). On the other hand, the result given by GeoStaS differs from all results produced by PiSQRD. In fact, ResiCon and PiSQRD are more coherent than GeoStaS and PiSQRD, which can be expressed by the mean of \mathcal{VI} . However, it would be naive to use the mean value as an indicator of self-consistency of PiSQRD. Among partitionings produced by PiSQRD, N(N-1)/2 agreements were calculated (where N is the number of models of 1d1d), whereas the comparison of ResiCon or GeoStaS with PiSQRD gave N values of \mathcal{VI} . Consequently, the average value of \mathcal{VI} between PiSQRD clusterings is not directly comparable with the average value of \mathcal{VI} , e.g. for ResiCon vs. PiSQRD.

Therefore, to give a better picture of the divergence of results we exploit the fact that \mathcal{VI} is a metric in the space of all clusterings. For a given partitioning we computed the radius of a ball centered at that partitioning, encompassing a certain fraction of the results. In Table 2.2 we provide values of radii of balls which encompass 25, 50 and 75% of the results. In the case of PiSQRD, we constructed balls (for a given percentage of results) centered at each partitioning, and computed the mean values of their radii. The mean of radii does not carry the bias mentioned earlier. Additionally, histograms of \mathcal{VI} for each protein can be found in Supplementary Materials.

Quality analysis

The box-and-whisker plot in Figure 2.5 provides a concise picture of quality scores Q (see Methods) of the dynamic domains assignments. It clearly shows that beyond a few exceptions ResiCon gives the best results. There are 7 notable exceptions: 4a5v, 2ktf, 3egf, 1adr, 1bf8, 2htg and 2kr6.

The single dynamic domains found by PiSQRD (blue in Figure 2.5) were produced using low-energy modes, computed as eigenvectors of a structural covariance matrix estimated by superimposing all models in an ensemble on a representative structure. In the *Supporting Materials* we explain how this representative structure is chosen, and show qualities of dynamic domains found by PiSQRD using different methods of estimating the structural covariance matrix. It should be emphasized, however, that these dynamic domains strongly depend on the method of estimating the structural covariance matrix, which is not part of PiSQRD's functionality. Therefore, these results should only be treated as an additional insight into what the user can expect from a more complex analysis of NMR structures.

In the case of 1adr GeoStaS gave partitionings with lower Q, by introducing a trivial domain (cutting off C- and N-termini – see Supplementary Materials). On the other hand, for 4a5v GeoStaS identified a single domain, which gave a lower value of Q than any other partitioning.

For proteins 2ktf, 3egf and 2htg, PiSQRD produced results with lower Q than ResiCon, by finding numerous small quasi-rigid fragments. However, for 2ktf more than 25% of assignments found by PiSQRD contained a trivial, single-residue

domain (indicated in red, with an asterisk). Although our measure does not penalize for this, we consider such behavior undesirable. Also, note that proteins 3egf and 2htg comprise 53 and 27 residues accordingly. It seems that in case of small proteins ResiCon often identifies a single domain, which does not necessarily result in the lowest Q.

The 2kr6 protein is an interesting example. It contains a flexible linker composed of more than 30 residues. As a consequence, partitionings with high values of Q are observed. Only PiSQRD was able to produce lower values of Q, by cutting the linker into many shorter parts. The example of the 2kr6 protein shows, that ResiCon does not consider unstructured regions to be separate dynamic domains. Instead, residues constituting linkers and lacking long-distance contacts, are assigned to dynamic domains which are closest in sequence.

Comments

Although PiSQRD's capability of analyzing a single structure may be considered an advantage, results presented in Table 2.2 and Figure 2.5 show that there is a large discrepancy for different configurations of the same protein. Nevertheless, based on the histograms of \mathcal{VI} presented in Supplementary Materials and the values of radii in Table 2.2, we conclude that in most cases results given by ResiCon and PiSQRD are mutually more coherent, than GeoStaS and PiSQRD (e.g. 1cfc, 1qo6, 1vvd, 1vve, 1yug, 2k3c). Notable exceptions are: 2rgf, 2pas, 3mef and 1zda. For these proteins ResiCon did not find any significant structural transitions and achieved the lowest value of Q (see Figure 2.5) by assigning a single domain.

Dynamic domains found by ResiCon are generally larger (particularly: 2k3c, 1d1d, 2k0e and 2kr6). Conversely, GeoStaS often allocates flexible N- and C-terminal parts (e.g. 1adr, 1qo6, 1vve, 3egf) as quasi-rigid parts. The size of dynamic domains is also the main difference between ResiCon and PiSQRD. In case of small, static proteins (such as 1aey, 1pkt, 1pit, 2spz, 2ait, 3egf and 1zda), PiSQRD identifies numerous small and often trivial dynamic domains. ResiCon on the other hand detects no significant conformational changes (by analyzing an ensemble), and assigns a single dynamic domain. We observe that in many cases ResiCon identified a single domain which had the lowest value of Q among all partitionings (see 2rgf, 2pas, 1aey, 1pkt, 1pit, 2vil, 1aiw, 3mef, 2spz, 1leb, 2ait, 2pni, 2l14). In these cases ResiCon correctly detected that no significant transitions were present in the protein. Therefore, unlike PiSQRD and GeoStaS, ResiCon can reliably indicate whether conformational changes occur in an ensemble of structures.

It is also noteworthy that ResiCon does not employ any post-processing procedures. This keeps the algorithm simple and clean, but results in a discontinuity of certain partitionings (see 4a5v, 1bf8 and 1vvd), which we refer to as the zebra effect. Although this seems to be an artifact of the clustering algorithm, we will take a closer look at this effect and show that it may also carry valuable information referring to the protein's dynamics.

HIV-1 protease

An analysis of an MD trajectory of the HIV-1 protease showcases ResiCon's capabilities. This protein undergoes substantial conformational changes associated with opening/closing of its structural parts, so-called *flaps* ([27]). A database of X-ray-resolved structures, representing configurations which the protease can attain is available ([52]).

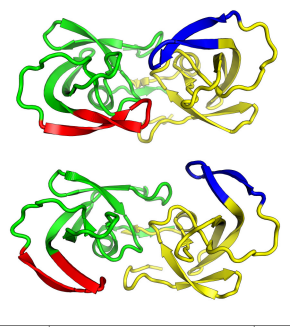
We examined a set of configurations of the HIV-1 protease acquired from a simulation carried out using the RedMD package ([24]). This coarse-grained force field was designed to simulate intramolecular motions in proteins and nucleic acids. It has correctly predicted the flap-opening motion in the HIV-1 protease, which is known biological fact ([27]), and was independently confirmed by an all-atom MD simulation ([46]). Roughly, about 1% of the trajectory seems to exhibit significant conformational transitions (flap opening events). This is a typical scenario in MD simulations, where a transition between meta-stable states is swift and fairly short. We needed a set of representative structures in order to find dynamic domains using ResiCon. From the whole trajectory, we extracted a set of 200 configurations using a generic procedure facilitating the Principal Component Analysis (PCA), implemented in the R programming language (in the bio3d package ([25])) – see Supplementary Materials.

Values of $\chi_{\rm min}$ for different numbers of clusters are given in Figure 2.3. The optimal number of clusters according to our procedure is 4. Figure 2.6 depicts the dynamic domains found by ResiCon and two representative configurations of the protease, as well as results from GeoStaS and PiSQRD. Because the sample of configurations was drawn according to the Boltzmann distribution, we were able to straightforwardly estimate the structural covariance matrix and provide PiSQRD with well-founded low-energy modes, and acquire a high-quality partitioning into two domains. Results produced by GeoStaS were acquired from the whole trajectory of the protease.

Dynamic domains identified by ResiCon have the highest value of the Q measure. Note that ResiCon does not try to minimize Q, but to produce a clustering with the optimal value of the χ_{\min} indicator. Consequently, at the cost of a slightly higher Q (ca. $1\mathring{A}$) we arrive at a partitioning which corresponds to the biologically relevant sub-division of the protein. It is also worth mentioning that ResiCon's partitioning into k=2 clusters incidentally leads to an identical assignment as the one produced by PiSQRD.

Motions of the flaps between the closed and open states are crucial in the functionality of the HIV-1 protease. It is also known that throughout their motion they remain quasi-rigid ([20]). Therefore, using ResiCon, we successfully extracted a simplified picture of the mobility of HIV-1 protease, which agrees with experimental knowledge.

Alternative partitioning into 6 clusters (see Figure 2.7) also deserves interest. Apart from the flaps, additional dynamic domains resembling "arms" carrying the flaps can be observed. Note that conversely to the case of the four dynamic domains, the quasi-static regions are similar, but not ideally reflective. This indicates



Method	Clustering	Q
GeoStaS		9.500
ResiCon		10.409
PiSQRD		9.189

Figure 2.6: Two representative conformations of HIV-1 protease with flaps being closed and open, and summary of results produced by GeoStaS, ResiCon and PiSQRD. Structures are colored according to the dynamic domains detected by ResiCon.

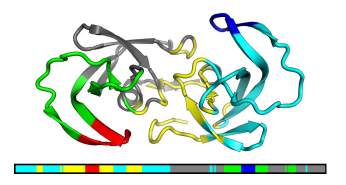


Figure 2.7: More subtle division emerges when the number of clusters is set to 6.

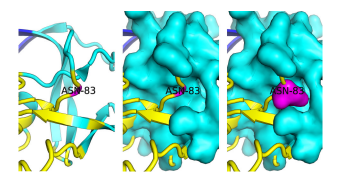


Figure 2.8: From the analysis carried out by ResiCon one can see that throughout the trajectory ASN-83 (magenta) not only remains close to residues in the N-terminal arm domain (gray) but also fills a cavity in that dynamic domain.

that motions of the two centrally symmetric sub-units of the HIV-1 protease in the provided trajectory were slightly different.

It can be also seen that domains are discontinuous. Especially residues belonging to the N-terminal lobe (yellow) and to the arm (cyan) are interleaved. In the next section we will analyze this effect, and show that this partitioning also carries valuable information.

The zebra effect

Dynamic domains found by ResiCon may include residues that seem to be pulled out from another quasi-rigid fragment. This is indicated as discontinuities (stripes) in Figure 2.7. Let us take a look at an example of such an extracted residue and try to understand the source of this effect. ResiCon assigned the ASN-83 residue to the arm (gray) dynamic domain (see Figure 2.8). However, its sequential neighbors belong to the lobe domain (yellow). The reason of this discontinuity is that, although ASN-83 has peptide bonds with its yellow neighbors, they are outweighted by contacts with the gray residues (see Figure 2.8). The discontinuity suggests that throughout the trajectory ASN-83 remained docked in the cavity of the arm domain, while its peptide bonds formed an axis of a hinge.

It seems that the zebra effect is not accidental, and that it can be used to find residues which act as "pivot points". However, this effect is volatile and depends on small fluctuations in the input. Residues are assigned to dynamic domains based on the fuzzy membership matrix. At certain positions values of membership to different domains may be almost equal, and when discretized cause emergence of stripes. Therefore, to strengthen the identification of these "pivot point" residues, a more thorough analysis of the membership matrix is required.

2.6 Conclusions

Typically, protein structures are flexible and mobile, and their conformational changes may be crucial in facilitating signaling and metabolical processes in which they participate. Breaking a structure into dynamic domains may be compared to discovering gears and levers connected by cogs and pegs analogous to those found in classical machines. Such analyses allow us to better understand molecular mechanisms responsible for biological functions (e.g. [50]), facilitate MD simulations in large time-scales with simplified forcefields (e.g. [48]), or discover potential binding sites when designing inhibitors (e.g. [58]).

We have presented a universal tool for discovering dynamic domains in proteins. ResiCon is capable of analyzing a single structure, or an NMR ensemble of structures provided in a PDB format. It can also be applied to the set of independently obtained structures (e.g. X-ray crystallographic structures obtained under different conditions). In any case, it provides a complete set of results comprising partitioning of the molecule into dynamic domains with additional highlighting of residues composing hinges and interfacial regions. ResiCon also provides an indicator of partitioning quality, and suggests the optimal number of domains.

We tested our method using the reference set of NMR structures. It is comparable or better than the recently developed GeoStaS and PiSQRD. Apart from giving more compact dynamic domains, it is also capable of distinguishing structures composed of a single quasi-rigid region.

We have made ResiCon available online.¹ To make the analysis feasible and limit the number of uploads, queries may be rerun with changed parameters. Also, all queries and results are stored on our server.²

¹http://dworkowa.imdik.pan.pl/EP/ResiCon

²These two features require registration for security reasons.

PDB code	Method	Clustering	Q	PDB code	Method	Clustering	Q
	GeoStaS		1.430		GeoStaS		5.678
2rgf	ResiCon		1.275	2k3c	ResiCon		3.676
	PiSQRD		1.421		PiSQRD		3.422
	GeoStaS		4.926		GeoStaS		12.658
2pas	ResiCon		1.398	1cfc	ResiCon		3.780
	PiSQRD		6.141		PiSQRD		2.294
-	GeoStaS		1.520		GeoStaS		4.478
1aey	ResiCon		1.520	1a67	ResiCon		4.548
	PiSQRD		4.455		PiSQRD		5.887
	GeoStaS		1.670		GeoStaS		5.700
1pkt	ResiCon		1.670	3egf	ResiCon		5.713
	PiSQRD		5.147		PiSQRD		3.499
	GeoStaS		0.678		GeoStaS		16.195
4a5v	ResiCon		1.698	2pni	ResiCon		5.994
	PiSQRD		2.561		PiSQRD		6.993
	GeoStaS		2.177		GeoStaS		9.610
1pit	ResiCon		2.209	1zda	ResiCon		6.549
	PiSQRD		2.171		PiSQRD		6.798
	GeoStaS		6.329		GeoStaS		6.241
2vil	ResiCon		2.414	1adr	ResiCon		7.417
	PiSQRD		4.146		PiSQRD		7.139
	GeoStaS		8.066		GeoStaS		8.479
1aiw	ResiCon		2.636	1yug	ResiCon		8.230
	PiSQRD		4.527		PiSQRD		9.033
	GeoStaS		2.822		GeoStaS	•	19.862
2ktf	ResiCon		2.693	1d1d	ResiCon		8.368
	PiSQRD		0.636		PiSQRD		7.313
	GeoStaS		3.299		GeoStaS		10.284
1vve	ResiCon		2.767	2114	ResiCon		8.525
	PiSQRD		2.767		PiSQRD	•	10.411
	GeoStaS		3.770		GeoStaS		7.947
3mef	ResiCon		3.086	1bf8	ResiCon		9.153
	PiSQRD	_	4.765		PiSQRD		4.519
	GeoStaS		5.439		GeoStaS		10.514
1vvd	ResiCon		3.110	2htg	ResiCon		10.514
	PiSQRD		2.821		PiSQRD		8.665
	GeoStaS		3.244	1qo6	GeoStaS		11.403
2spz	ResiCon		3.212		ResiCon		11.057
	PiSQRD		8.071		PiSQRD		7.985
	GeoStaS		6.885		GeoStaS		12.771
1leb	ResiCon		3.234	2k0e	ResiCon		11.604
	PiSQRD		5.874		PiSQRD		7.927
	GeoStaS		5.370		GeoStaS		56.578
2ait	ResiCon		3.310	2kr6	ResiCon		54.160
	PiSQRD		3.871		PiSQRD		26.834

Table 2.1: Summary of results produced by GeoStaS, ResiCon and PiSQRD. Dynamic domains shown for PiSQRD are those for which the lowest value of Q was achieved.

	ResiCon vs. GeoStaS	ResiC	on vs. Pis	SQRD	D GeoStaS vs. PiSQRD			PiSQRD vs. PiSQRD		
	\mathcal{VI}	r _{25%}	r _{50%}	r _{75%}	r _{25%}	r _{50%}	r _{75%}	$\langle r_{25\%} angle$	$\langle r_{50\%} \rangle$	$\langle r_{75\%} \rangle$
2rgf	0.62	0.45	0.48	0.57	0.26	0.26	0.32	0.01	0.14	0.20
2pas	1.94	2.43	2.45	2.47	2.00	2.08	2.11	0.61	0.96	1.23
1aey	0.00	1.99	2.33	2.37	1.99	2.33	2.37	0.77	1.10	1.52
1pkt	0.00	2.07	2.24	2.36	2.07	2.24	2.36	1.38	1.63	1.84
4a5v	1.00	1.08	1.12	1.19	1.53	1.54	1.60	0.24	0.29	0.38
1pit	1.59	1.76	2.05	2.14	2.42	2.55	2.74	1.35	1.70	1.94
2vil	1.89	1.19	1.39	1.89	2.31	2.49	2.67	1.11	1.36	1.57
1aiw	1.64	1.66	1.80	2.13	2.58	2.69	2.77	1.30	1.53	1.90
2ktf	0.80	1.69	2.01	2.06	2.15	2.36	2.48	0.93	1.17	1.74
1vve	1.05	0.00	0.00	0.00	1.05	1.05	1.05	0.62	0.62	0.62
3mef	0.32	1.30	1.43	1.50	1.21	1.32	1.40	0.64	0.86	1.09
1vvd	1.24	0.21	0.21	0.21	1.26	1.26	1.26	0.31	0.31	0.31
2spz	0.13	1.42	1.62	1.94	1.37	1.59	1.96	1.28	1.60	1.73
1leb	0.61	2.05	2.27	2.41	2.33	2.37	2.45	0.92	1.21	1.52
2ait	0.71	1.26	1.65	1.82	1.49	1.67	1.99	0.99	1.31	1.52
2k3c	1.86	0.82	0.97	0.98	1.61	1.69	1.69	0.16	0.42	0.63
1cfc	1.60	0.41	0.44	0.97	1.41	1.48	1.86	0.77	0.92	1.30
1a67	1.07	1.90	1.96	2.06	2.22	2.38	2.69	1.58	1.93	2.12
3egf	0.14	1.30	1.40	1.43	1.29	1.41	1.45	0.63	0.91	1.14
2pni	1.60	1.16	1.58	1.78	2.00	2.25	2.35	1.18	1.44	1.65
1zda	0.79	1.37	1.55	1.96	0.83	1.28	1.78	1.26	1.54	1.75
1adr	0.84	0.80	1.30	1.81	1.11	1.64	2.22	1.19	1.57	1.94
1yug	1.79	0.90	1.00	1.31	1.98	2.12	2.15	0.70	0.99	1.29
1d1d	1.47	0.00	0.21	3.15	1.47	1.51	3.80	1.08	3.08	3.17
2114	0.55	0.49	0.57	0.75	0.91	0.98	1.18	0.61	0.71	0.95
1bf8	1.61	0.74	1.27	3.22	1.69	1.89	4.20	1.13	1.74	2.99
2htg	0.00	1.36	1.58	1.91	1.36	1.58	1.91	0.84	1.13	1.30
1qo6	0.88	0.63	0.72	0.88	0.87	0.98	1.18	0.57	0.71	0.94
2k0e	0.97	0.59	0.66	0.82	1.05	1.16	1.31	0.72	0.87	1.03
2kr6	1.11	0.79	0.81	0.84	0.85	0.87	0.93	0.61	0.68	0.82

Table 2.2: Discrepancies in assignments expressed by radii of balls encompassing 25%, 50% and 75% of results. The ordering of the results is the same as in Figure 2.5, i.e. best-scoring results are first.

Chapter 3

Towards the identification of molecular cogs

Computer simulations of molecular systems allow determination of microscopic interactions between individual atoms or groups of atoms, as well as studies of intramolecular motions. Nevertheless, description of structural transformations at the mezoscopic level and identification of causal relations associated with these transformations, is very difficult. Structural and functional properties are related to free energy changes. Therefore, in order to better understand structural and functional properties of molecular systems, it is required to deepen our knowledge of free energy contributions arising from molecular subsystems in the course of structural transformations.

The method presented in this work quantifies the energetic contribution of each pair of atoms to the total free energy change along a given collective variable. Next, with the help of a genetic clustering algorithm, the method proposes a division of the system into two groups of atoms referred to as molecular cogs. Atoms which cooperate to push the system forward along a collective variable are referred to as forward cogs, and those which work in the opposite direction as reverse cogs.

The procedure was tested on several small molecules for which the genetic clustering algorithm successfully found optimal partitionings into molecular cogs. The primary result of the method is a plot depicting the energetic contributions of the identified molecular cogs to the total Potential of Mean Force (PMF) change. Case-studies presented in this work should help better understand the implications of our approach, and were intended to pave the way to a future, publicly available implementation.

INTRODUCTION

Molecular dynamics (MD) simulation methods and advanced algorithms for calculating free energy bring us closer to predicting the physical properties of biomolecules [54, 12, 15]. However, computer simulations are not limited to interpreting experimental results. *In silico* one may also process MD data which can provide much more detailed information than that accessible in any experiment. The key to a deeper understanding of complex molecular systems is the extraction of valuable information from data produced in such simulations.

Biomolecules carry out their functions through conformational transitions between meta-stable states. Such phenomena can be simplified and described by a selected reaction coordinate which, in many cases, is a collective variable [13]. A valuable result of many simulation procedures (such as Umbrella Sampling [51], Thermodynamic Integration [34], Adaptive Biasing Force [15], and others[26]), is a free energy profile, the so-called Potential of Mean Force (PMF) [34]. In this study, the discussion is limited to one-dimensional PMFs, although it should be noted that the aforementioned methods can be formulated more generally and produce multi-dimensional free energy profiles. Of course, a one-dimensional PMF may be an oversimplification of what is going on in a complex system, but a meaningful collective variable usually leads to a free energy profile which makes the complicated transition more comprehensible [40, 12]. However, what the PMF does not provide is the information about what drives transitions between meta-stable states.

Various attempts to understand and describe the internal mechanics of molecular systems have already been reported [2, 4, 11, 35, 45, 1, 47]. We did not, however, find any general-purpose approach for studying a broader spectrum of cases, and in particular a methodology explaining the cause of a transition in terms of free energy contributions arising from certain parts of a molecule.

In this study we propose a new approach of analyzing the tendencies of a molecular system to undergo a selected structural transition. The main idea is to look at a shift along a collective variable as an effect of two opposing tendencies generated by interactions within the molecule. Our method indicates two groups of atoms – referred to as molecular cogs – which, through cooperative interactions, are the source of these tendencies. For this purpose, we construct undirected graphs with weights between nodes expressed by energetic contributions to the free energy change arising from pair-wise interactions between atoms. These graphs are then partitioned into subsystems – corresponding to molecular cogs – using a genetic clustering algorithm. We present results for small, model systems which served as case-studies for the identification of molecular cogs and for testing if their functioning agrees with our intuitions.

METHODOLOGY

Decomposition of the Helmholtz free energy, A, was investigated in the past, most notably by Karplus and coworkers [6, 8]. The aim was the determination of contri-

butions to the free energy coming from components comprising the potential energy of the system. These potential energy components might come from different interaction types, or from cooperation between subsystems of the whole molecule. The strategy of expressing the potential energy of a system as a sum of terms, $U = \sum_i U_i$, and computing the contribution of each of these terms to the free energy is ineffective because additivity in the potential energy components does not imply additive contributions to the entropy[18].

Two attempts of describing contributions to the free energy coming from parts of the system are worth mentioning. The first one was an approximate approach based on Free Energy Perturbation, in which higher order terms were neglected [8]. These terms are not, however, negligible, which severely limits the applicability of this method. An alternative route, employing Thermodynamic Integration, was also explored [6]. However, this approach is also limited, namely – values of the free energy contributions depend on the choice of the integration path.

We propose a different approach, in which pair-wise energetic contributions to the free energy are readily attained. Alas, as in the aforementioned methods of free energy decomposition, our current formulation struggles with a description of the entropic contributions, and at the present is not included in our method. The current implementation is primarily applicable to small molecules for which the role of entropy in structural transitions is negligible (see an example of a PMF in the *Results* section).

Our analysis originates from the following formula [10]:

$$A'(\xi^*) = \langle m_{\xi} \nabla U \cdot \mathbb{M}^{-1} \nabla \xi \rangle_{\xi^*} - \langle \mathbf{v} \cdot \nabla (m_{\xi} \nabla \xi) \mathbf{v} \rangle_{\xi^*}, \tag{3.1}$$

where A' is the derivative of the free energy with respect to the collective variable ξ , \mathbb{M}^{-1} is a diagonal matrix of inverse masses, and \mathbf{v} are velocities. The term m_{ξ} is defined by:

$$m_{\xi} := \left[\sum_{i} m_{i}^{-1} \left(\frac{\partial \xi}{\partial \mathbf{x}_{i}} \right)^{2} \right]^{-1} \tag{3.2}$$

and can be interpreted as inertia of an effective point mass moving along the ξ coordinate.

On the right-hand side of Equation (3.1) we have, respectively, energetic and entropic contributions to the free energy change, both expressed as conditional averages, with the collective variable fixed at ξ^* . In the *Supporting Information* we briefly explain how these averages are estimated *via* constrained Molecular Dynamics (cMD) simulations, and highlight an important limitation of this method.

Note, that if we consider the potential energy as a sum of interaction components:

$$U = U^{ele} + U^{vdw} + U^{bond} + \dots = \sum_{i}^{I} U^{I}$$

the energetic contribution in Equation (3.1) maintains this additivity:

$$\langle m_{\xi} \nabla U \cdot \mathbb{M}^{-1} \nabla \xi \rangle_{\xi^*} = \sum_{i}^{I} \langle m_{\xi} \nabla U^I \cdot \mathbb{M}^{-1} \nabla \xi \rangle_{\xi^*}, \tag{3.3}$$

as was noted by Chipot et al. [13].

In our numerical experiments we used a force field with the following set of interaction types:

- *ele* (electrostatic interactions);
- vdw (van der Waals interactions);
- bond (2-body bonded interactions);
- angl (3-body angle interactions);
- tors (4-body torsional interactions).

We shall also consider:

- nbd (non-bonded interactions, the sum of ele and vdw interactions);
- conf (conformational interactions, composed of bond, angl and tors interactions);
- total (the sum of nbd and conf interactions).

Decomposition of the energetic contribution

Our purpose was to identify the molecular cogs, i.e. sets of atoms, which cooperate and push the whole system forward/backward along a reaction coordinate. We approached this problem by converting it into the task of finding clusters in a graph. Nodes in such a graph correspond to atoms, whereas edges represent cooperation of pairs of atoms. A natural measure of such cooperation can be introduced by taking into account the energetic contribution to the free energy in Equation (3.1).

Electrostatic, van der Waals and two-atom chemical bond interactions can be readily transformed into weights in the graph. For example, electrostatic interactions between atoms α and β lead to the following contribution:

$$c_{\alpha\beta}^{ele}(\xi^*) := \frac{1}{m_{\alpha}} \left\langle m_{\xi} \frac{\partial U^{ele}}{\partial \mathbf{x}_{\alpha}} \cdot \frac{\partial \xi}{\partial \mathbf{x}_{\alpha}} \right\rangle_{\xi^*} + \frac{1}{m_{\beta}} \left\langle m_{\xi} \frac{\partial U^{ele}}{\partial \mathbf{x}_{\beta}} \cdot \frac{\partial \xi}{\partial \mathbf{x}_{\beta}} \right\rangle_{\xi^*}. \tag{3.4}$$

Alas, n-body potential energy components cannot, in general, be decomposed into a sum of pair interactions. However, because the free energy contributions are of the form $\nabla U \cdot \mathbb{M}^{-1} \nabla \xi$, such decomposition is not required.

To clarify, let us consider a 3-body potential energy component, e.g. $U^{angl}(\mathbf{x}_{\alpha}, \mathbf{x}_{\beta}, \mathbf{x}_{\gamma})$. The weight $c^{angl}_{\alpha\beta}$ of an edge joining nodes α and β represents the contribution coming from atoms α and β which interact $via\ U^{angl}$, while the γ atom is kept at a fixed position. By asserting that the sum of pair-wise contributions is equal to the total contribution arising from $U^{angl}(\mathbf{x}_{\alpha}, \mathbf{x}_{\beta}, \mathbf{x}_{\gamma})$, i.e.:

$$\sum_{i>j} c_{ij}^{angl}(\xi^*) = \sum_{i=\alpha,\beta,\gamma} \frac{1}{m_i} \left\langle m_{\xi} \frac{\partial U^{angl}}{\partial \mathbf{x}_i} \cdot \frac{\partial \xi}{\partial \mathbf{x}_i} \right\rangle_{\xi^*},$$

we arrive at the following definition of the cooperation term:

$$c_{\alpha\beta}^{angl}(\xi^*) := \frac{1}{2} \sum_{i=\alpha,\beta} \frac{1}{m_i} \left\langle m_{\xi} \frac{\partial U^{angl}}{\partial \mathbf{x}_i} \cdot \frac{\partial \xi}{\partial \mathbf{x}_i} \right\rangle_{\xi^*},$$

where the 1/2 means that the overall angl contribution is evenly distributed between the cooperation terms: $c_{\alpha\beta}^{angl}$, $c_{\beta\gamma}^{angl}$ and $c_{\alpha\gamma}^{angl}$.

For the general case of an n-body energy component, U^{I} , we propose the following definition of the cooperation term between atoms α and β :

$$c_{\alpha\beta}^{I}(\xi^{*}) := \frac{1}{n-1} \sum_{i=\alpha} \frac{1}{m_{i}} \left\langle m_{\xi} \frac{\partial U^{I}}{\partial \mathbf{x}_{i}} \cdot \frac{\partial \xi}{\partial \mathbf{x}_{i}} \right\rangle_{\xi^{*}}.$$

Note that we propose to distribute the cooperation evenly among all pairs of atoms involved in the interaction I. With this definition we constructed graphs for all interaction types, I.

A matrix, $C^I(\xi^*)$, such that $[C^I(\xi^*)]_{\alpha\beta} := c^I_{\alpha\beta}(\xi^*)$ constructed for a particular value of the collective variable ξ^* and the interaction type I is referred to as the transient cooperation matrix for I (see Figure 3.1). For convenience, the transient cooperation matrix for the total interaction type does not contain any superscipt:

$$\mathcal{C}(\xi_i) := \sum_I \mathcal{C}^I(\xi_i).$$

Note that the sum $\sum_{\alpha>\beta} [\mathcal{C}(\xi_i)]_{\alpha\beta}$, is the overall energetic contribution to $A'(\xi_i)$ in Equation (3.1).

Comment on the entropic contribution

The second term on the right-hand side of Equation (3.1) does not explicitly depend on U. It is possible to propose pair-wise contributions of this quantity in order to construct matrices which in turn could be used as input for the genetic clustering procedure described below. However, calculating these entropic contributions would require computation of the Hessian of the collective variable (the matrix of second-order derivatives with respect to atom coordinates) in each step of the simulation, which would make the computational cost of the procedure prohibitively high. We speculate on a possible solution of this problem in the Conclusions section.

Clustering

Data clustering is the procedure of finding disjoint subsets (clusters) of objects that share a high affinity, and are dissimilar to objects from other clusters. To carry out a clustering procedure we need a pair-wise affinity measure; thus, the data set is often represented by a weighted graph. Such a graph is encoded by an affinity matrix with elements corresponding to weights between nodes. In this

study we were looking for groups of atoms whose cooperation decreased (increased) the free energy, thus pushing (pulling) the whole system forward (backward) along a reaction coordinate. Thus, we took the pair-wise energetic cooperation between atoms introduced earlier for the affinity measure.

We will refer to the clusters identified in a transient cooperation matrix as transient molecular cogs. The free-energy-reducing group is referred to as forward cogs, and the second group (which, conversely, increases the free energy gap) as reverse cogs. We, therefore, assume that a step taken by the molecule along a reaction coordinate is a consequence of a resultant tendency generated by the molecular cogs. But cross-cooperation between atoms from different groups can also occur, and we designate this unwanted effect 'gear grinding' (see the Results section).

Affinity Propagation algorithm

A considerable challenge associated with clustering is that different procedures produce different results. Therefore, it was crucial that the *right* clustering algorithm was chosen. We anticipated that the optimal algorithm should recover molecular cogs with as little gear grinding as possible, but we could not estimate how large this effect might be. To gain some insight into how molecular cogs might look like, we chose a clustering algorithm which is known to be successful in other applications.

Most clustering algorithms assume a non-negative affinity measure, whereas cooperation between atoms can be negative as well as positive. Among those algorithms which accepted negative values, Affinity Propagation (AP) was the procedure found to be the most promising[21]. The AP algorithm searches for *exemplars*, i.e. nodes within the graph around which non-exemplars are grouped, thus forming a cluster. It is an iterative procedure, where, in each step, "messages" between nodes are exchanged to designate exemplars and their followers.

In our case, the sign of cooperation is arbitrary i.e. it depends on the direction of the reaction coordinate, ξ . It was crucial to ensure that molecular cogs found for a reaction coordinate $\xi' := -\xi$ were the same as those found for ξ . We resolved this problem by carrying out two clusterings with the AP method: one for $\mathcal{C}(\xi)$, and the second one for $-\mathcal{C}(\xi)$, in which the sign of all elements is changed, such as produced for ξ' . Description of the AP algorithm and our method for merging two clusterings can be found in *Supporting Information*.

One of the drawbacks of the AP method, similarly as in other clustering procedures, is that it requires several parameters, which influence the outcome. It appeared that small variations in parameters lead to markedly different results, and it was difficult to find a single set of parameters suitable for all cases (see the discussion in the *Supporting Information*). Nevertheless, the AP method gave us a first estimation of how molecular cogs look like, and it appears to be sufficiently good to initialize the genetic clustering procedure (see the *Genetic clustering* section).

Objective function

Molecular cogs identified by the AP algorithm were laden with low gear grinding, but that was not always a valuable finding. We noticed that in cases where no good partitioning was achievable, the AP method produced one-element clusters, which we considered to be artificial and incorrect.

The AP algorithm performs a clustering which maximizes the so-called *net similarity* [21]. This objective function, although helpful in many applications, does not carry any physical meaning. Note that cooperation, which we used to express affinity between atoms, translates into free energy differences, and thus into the tendency of the whole system to move along a collective variable. Molecular cogs should not only be laden with low gear grinding, but also cooperation generated by the cogs should cover the overall free energy change as much as possible.

To clarify, given a cooperation matrix C, the sum of all negative (positive) elements is the overall tendency of the system to go forward (backward) along a reaction coordinate. It is desired for forward (reverse) cogs to cover as much of this overall tendency as possible. When there is only one atom in a cluster, there is no coverage, even though gear grinding might be small.

Let us denote by FC the set of atom indices, which were assigned to the forward cogs, and analogously by RC the set of atoms assigned to the reverse cogs. Given a cooperation matrix, $C = [c_{ij}]$, the following three quantities are useful in measuring the quality of molecular cogs:

• Forward Cogs Rate:

$$FCR := \frac{\sum_{i,j \in FC} c_{ij}}{\sum_{c_{ij} < 0} c_{ij}}$$
 (3.5)

• Reverse Cogs Rate:

$$RCR := \frac{\sum_{i,j \in RC} c_{ij}}{\sum_{c_{ij} > 0} c_{ij}}$$
(3.6)

• Gear Grinding Rate:

$$GGR := \frac{\sum_{\substack{i \in FC \\ j \in RC}} c_{ij}}{\sum_{\substack{c_{ij} > 0 \\ \sum_{c_{ij} < 0}}} + \frac{\sum_{\substack{i \in FC \\ j \in RC}} c_{ij}}{\sum_{\substack{c_{ij} < 0 \\ \sum_{c_{ij} < 0}}} c_{ij}}$$
(3.7)

We assumed that both positive and negative elements exist in the cooperation matrix, C, i.e. that $\sum_{c_{ij}<0} c_{ij} \neq 0$ and $\sum_{c_{ij}>0} c_{ij} \neq 0$. In cases in which the denominator is null, we substitute the whole fraction by 0.

FCR is the ratio of the contribution captured by the forward cogs to the total forward propensity found in C. RCR is analogous, and both these measures have a maximum value of 1. Magnitude of misplaced contributions between atoms from different clusters is indicated by GGR, which has a maximum value of 2.

We propose the following molecular cogs quality measure:

$$SCORE(FC, RC, C) := 0.5(FCR + RCR - GGR), \tag{3.8}$$

and SCORE := 0 for cases when any of the cogs consists of a single atom. The above scoring function was used in our genetic clustering procedure (described in the following section). See Figure 3.2 for an example of a clustering which maximizes the SCORE.

SCORE equals 1 if the corresponding molecular cogs cover the whole propensity of the system to move along a reaction coordinate, and no gear grinding occurs. We observed that a SCORE less than 0.5 often indicates that the subdivision of the system into molecular cogs is noisy.

Note that a trivial partitioning in which all atoms are assigned to one group, e.g. FC, indicates that the system as a whole has a tendency to move forward. We denote such trivial partitionings as "all FC" and "all RC". The maximal value of SCORE is then 0.5, and although this is unfavorable, in many cases a trivial partitioning had the highest SCORE. In such cases, there was no convenient clustering into two groups, which means that the gear grinding was high.

Genetic clustering algorithm

Genetic algorithms are widely used in optimization problems when potential solutions are readily assessed and codified. They emulate the process of natural selection, promoting solutions – referred to as *specimens* – with higher scores. The codification of a solution is treated as its chromosome, which allows for *mutation* of a solution, and *crossover* with other chromosomes, thus "spawning" new specimens. At the end of each iteration of a genetic algorithm, there is a stage called *selection*, during which low-scoring solutions have a lesser chance of "survival".

We used the genetic clustering algorithm to find molecular cogs with the highest SCORE, as defined in Equation (3.8) (see [14] for a good introduction to genetic clustering). The partitioning of a molecular system into molecular cogs was encoded by an array of numbers from the set $\{-1,0,1\}$, where the *i*th element of the array corresponded to the *i*th atom in a molecule. Values -1 and 1 translate into assignments to the forward and reverse cogs, respectively. An atom not belonging to any cluster was tagged by 0; this occurred when the atom did not cooperate with any other atom (see Figure 3.3).

To initialize the genetic clustering procedure, the AP algorithm was executed by applying five different methods of setting the diagonal elements in the affinity matrix (see *Supporting Information*). Next, it adds the "all FC" and "all RC" trivial solutions to the starting set. Following this, solutions are iteratively chosen, mutated and added to the set, until the starting population contained 200 candidate solutions.

Once the initial set is generated, the genetic procedure repeated the following steps:

- 1. Compose (randomly) 50 pairs of solutions to generate offspring using the crossover procedure.
- 2. Select (randomly) 20 solutions to generate offspring using the mutation procedure.

- 3. Calculate SCORE for the offspring and add it to the population.
- 4. Draw (randomly) 200 solutions from the population and discard the rest.
- 5. Choose the best scoring solution and check if there is any improvement in the SCORE. If there was none for 10 consecutive iterations, return the best solution. Otherwise, return to 1.

All random selections are done without repeats, so that a given solution with SCORE s is chosen with a probability proportional to e^{2s} . Although the number of parameters required in the genetic algorithm is daunting, changes to the majority of these parameters only influence the speed of arriving at the optimal solution (see the *Supporting Information* for a more detailed discussion of the parameter's influence on the outcome).

Figure 3.4 shows that the genetic clustering finds the optimal solution for a range of transient cooperation matrices for *nbd* (complete results can be found in the *Results* section). The optimal solutions were found by means of a brute-force search, i.e. by producing all possible partitionings and calculations of their SCOREs. It is worth noting that in all cases there was a singular solution with the highest SCORE.

Trapezoidal rule for integrating A

The free energy difference between ξ_X and ξ_Y can be expressed as an integral:

$$\Delta A = \int_{\xi_X}^{\xi_Y} A'(\xi^*) d\xi^*. \tag{3.9}$$

The free energy derivative, $A'(\xi^*)$, can be estimated at a given ξ^* from a cMD simulations via Equation (3.1). The integral in Equation (3.9) can then be calculated using the trapezoidal rule [42]:

$$\widehat{\Delta A} \approx \frac{\Delta \xi}{2} \left\{ \widehat{A'(\xi_1)} + 2\widehat{A'(\xi_{i+1})} + \dots + 2\widehat{A'(\xi_{M-1})} + \widehat{A'(\xi_M)} \right\}, \tag{3.10}$$

where $\widehat{A'(\xi_i)}$ denotes an estimate of the free energy derivative at ξ_i . This requires independent cMD simulations at M values of the collective variable, $\{\xi_i\}_{i=1}^M$, with the grid size of $\Delta \xi$.

Each estimate $\widehat{A'(\xi_i)}$ has a corresponding variance, $\sigma^2[\widehat{A'(\xi_i)}]$, which in turn propagates to the variance of the integral estimate, i.e. $\sigma^2[\widehat{\Delta A}]$. In the following section we explain how we estimated the variances $\sigma^2[\widehat{A'(\xi_i)}]$. Because estimates $\widehat{A'(\xi_i)}$ are independent, the variance of the whole integral follows from Equation (3.10) straightforwardly:

$$\sigma^{2}[\widehat{\Delta A}] = \frac{\Delta \xi^{2}}{4} \left\{ \sigma^{2}[A'(\xi_{1})] + 4\sigma^{2}[A'(\xi_{2})] + \dots + 4\sigma^{2}[A'(\xi_{M-1})] + \sigma^{2}[A'(\xi_{M})] \right\}$$
(3.11)

Details concerning estimation of the variances $\sigma^2[A'(\xi_1)]$ using a bootstrapping procedure can be found in the *Supporting Information*.

The same integration rule applies to all elements of the transient cooperation matrices. A matrix in which every element is a result of the above numerical integration is referred to as the *integrated cooperation matrix* or simply: cooperation matrix, and denoted by $C(\xi_X \to \xi_Y)$. The sum $\sum_{\alpha>\beta} [C(\xi_X \to \xi_Y)]_{\alpha\beta}$ is the energetic contribution to ΔA for the $\xi_X \to \xi_Y$ path. Molecular cogs found for this matrix are called global molecular cogs.

RESULTS

In the first part of this section we present detailed results for a small, 11-atom molecular model, to validate the concept of our theoretical approach. We attempted to indicate molecular cogs to verify whether the genetic algorithm finds the optimal clustering for the scoring function proposed in the *Objective function* section.

In the second part of the Results we show molecular cogs for the nbd, ele and vdw interactions for three other molecules. These case-studies allowed for testing of the transferability of the parameters used in the genetic clustering algorithm, but also uncovered certain subtleties characteristic to our approach.

The [NH3+]CC(I)I molecular model

We were interested in finding molecular cogs propelling a structural transition between two meta-stable states, separated by a high free energy barrier. For our first case-study we required a system with a fairly natural collective variable, in which all types of interactions are significant (conf as well as nbd). We used the 2,2-diiodoethan-1-aminium molecule, which in the SMILES format is encoded as: [NH3+]CC(I)I (we use the SMILES representation throughout this article because of its conciseness). The dihedral angle between atoms N1-C5-C6-I7 was our collective variable of choice (see Figure 3.5). We used the Generalized Amber Force Field (GAFF) to model interactions between atoms, with partial charges assigned using an empirical procedure, AM1-BCC (Table 3.1).

We focused on finding molecular cogs propelling the transition between the dihedral angles of $\xi_X := -172.5^{\circ}$ and $\xi_Y := -62.5^{\circ}$, which correspond to two PMF minima (Figure 3.5). The free energy barrier separating these minima is high, and interactions between the [NH3+] group and the iodine atoms provide strong *ele* and vdw contributions which influence this barrier.

Constrained MD simulations were carried out for fixed values of the ξ collective variable, each simulation with 10^5 1fs timesteps, at T=300K. We chose 239 points, $\{\xi_i\}_{i=1}^{239}$, equally separated by $\Delta\xi=0.5^{\circ}$, so that $\xi_1=-179.0^{\circ}$ and $\xi_{239}=-60.0^{\circ}$. This was done in order to encompass the $[\xi_X,\xi_Y]$ interval; note that $\xi_{14}=\xi_X$ and $\xi_{234}=\xi_Y$.

The length of the block in the block bootstrap estimation of averages and their corresponding variances was set to 10^2 , which led to 10^3 blocks for each simulation data set. We chose the length of the block with the assumption that the autocorrelation after 10^2 steps is negligible in the case of our simple system.

The cMD procedure was implemented in the Python programming language (ver. 2.7.2), using the Open Babel[39] package (ver. 2.3.2) to model the molecule (see *Supporting Information*). All simulations for the [NH3+]CC(I)I model took about 30h on a desktop computer.

We focused on the free energy differences and energetic contributions for the $\xi_X \to \xi_Y$ path. For these end-points we obtain the free energy difference $\Delta A \approx -16.4 \frac{\rm kcal}{\rm mol}$ which is close to the energetic contribution, $\Delta E \approx -16.7 \frac{\rm kcal}{\rm mol}$. The residual $0.3 \frac{\rm kcal}{\rm mol}$ is the entropic contribution, which was negligible for our model.

Overview

Results for the total, conf and nbd interactions are juxtaposed in Table 3.2, and for 2-body interactions (bond, ele, vdw) in Table 3.3. The first row contains optimal partitionings of the integrated cooperation matrices, i.e. global molecular cogs (see Figure 3.3 for explanation), along with a picture of the model colored according to the clustering. In the second row we placed the integrated cooperation matrices rearranged in accordance with the clusterings depicted in the previous row of the table (see Figure 3.2 for explanation). This representation visualizes the "density" of cooperativity within molecular cogs, and gear grinding. The third row shows SCOREs for transient molecular cogs and compares genetic clusterings with optimal, brute-force partitionings. In the next row we see an illustration presenting transient molecular cogs, which is a concise summary of how cooperation within the molecule changes with ξ . This plot is helpful in judging whether the global molecular cogs are similar to the transient molecular cogs, and in assessing the consistency of cooperation within the molecule. Finally, the last row contains a PMF-like contribution profile which we call the energetic contribution profile; this is the most important result of our analysis. The green line represents the total energetic contribution of a given interaction type, orange and blue lines show contributions of the reverse and forward cogs, respectively, and the purple line – the magnitude of gear grinding. Gear grinding is quantified as the sum of absolute values of all misplaced contributions, i.e. from pairs of atoms from different clusters. Forward and reverse cogs contributions are calculated as the sum of cooperation between atoms assigned to FC and RC, respectively.

Results for the [NH3+]CC(I)I molecular model

In this section we comment on the results presented in Tables 3.2 and 3.3. We explain the Tables row by row, highlighting important aspects of the analysis, starting with Table 3.2 and then proceeding with Table 3.3.

In the first row of Table 3.2, global molecular cogs for *total* and *conf* interactions are trivial ("all FC"), with very low SCOREs. Thus, no interesting cooperation was detected for these interaction types – the whole system has a general propensity for preferring the state around ξ_Y . On the other hand, global molecular cogs for the *nbd* interactions have a high SCORE. The clustering suggests, in particular, that non-bonded interactions between the nitrogen atom and the iodine atoms

hinder the transformation (as expected). A more surprising implication was that the hydrogen atoms from the [NH3+] group cooperate with the C6 atom.

Clustered cooperation matrices are placed in the second row of Table 3.2. For the *total* and *conf* interactions, for which the partitioning was trivial, the matrices kept their original form, whereas for the *nbd* interactions rows and colums were rearranged in accordance with the partitioning. Matrices for *total* and *conf* are almost indistinguishable, which suggests that the transition is mainly governed by the *conf* interactions. It is also clear that the cooperation matrix for *nbd* is less "dense", with little gear grinding.

In the next row we see that the partitionings into transient molecular cogs for nbd have consistently higher SCOREs than those found for total and conf. There is a drop in SCOREs corresponding to the crossing of the free energy maximum at about -115° dihedral angle. SCOREs for nbd molecular cogs vary significantly for different values of the collective variable, and the division into forward and reverse cogs becomes slightly more difficult after crossing the free energy maximum.

The transient molecular cogs depicted in graphs in the fourth row of the table show that the transient molecular cogs for *total* and *conf* exhibit no interesting structure and, in most cases, are of the "all FC" or "all RC" type. Transient molecular cogs for *nbd* are fairly consistent with global molecular cogs. Interestingly, the previously noted *nbd* cooperation between hydrogen atoms H2, H3, H4 and the C6 atom persists throughout the transition.

In the last row of the table we placed the energetic contribution profiles. Because the global molecular cogs for the *total* and *conf* interactions were of the "all FC" form, the only contributions come from the forward cogs. For the *nbd* interactions, the plot shows a beautiful separation of contributions coming from the forward $(-4.2\frac{\text{kcal}}{\text{mol}})$ and the reverse cogs $(3.7\frac{\text{kcal}}{\text{mol}})$ for non-bonded interactions, and low gear grinding for the $\xi_X \to \xi_Y$ transition.

Let us now look at the results in Table 3.3. In all three cases global molecular cogs were non-trivial, although the SCORE for bond is significantly lower than for ele and vdw. The partitioning for bond suggests that there is an impediment arising from interactions of the C6, I7 and I8 atoms, whereas the rest of the system favors the state around ξ_Y . Moving on to ele and vdw interactions we see that their global molecular cogs share a common pattern, although the SCORE for the latter is slightly lower. We can also see that the cooperation of the H2, H3, H4 and C6 atoms (indicated earlier for nbd) is caused by the ele interactions.

The next row of Table 3.3 shows graphical representations of the clustered cooperation matrices. Not surprisingly, the *bond* matrix is more sparse than any other matrix, however judging by the low SCORE for global molecular cogs, this was not sufficient to ensure a clear division into the forward and reverse cogs. Conversely, for *ele* and *vdw* there seems to be a higher degree of gear grinding, yet the SCOREs were higher. This is due to the fact that the cooperation within molecular cogs is much stronger than gear grinding between them.

SCOREs in the third row of the table show that the transient molecular cogs for the bond interactions were consistently low. We see the opposite for ele, and a completely different situation for the vdw transient molecular cogs. The case

of the vdw cooperation is particularly interesting because it shows that the global molecular cogs can have a high SCORE despite the fact that most of the transient molecular cogs have SCOREs below 0.5.

In the fourth row of the table we see that the transient molecular cogs for bond are consistent with their global molecular cogs for dihedral angles in the $[-150^{\circ}, -100^{\circ}]$ interval (i.e. around the free energy maximum at $\xi \approx -115^{\circ}$) For the ele interactions we see a stable cooperation between the H1, H2, H3 and C6 atoms, occasionally aided by atoms: H9 and I8. The reason behind the shape of the vdw SCORE plot becomes clearer once we see that the cooperation for this interaction type has two stages – before and after crossing the free energy maximum. Transient molecular cogs for vdw on the left side of the free energy maximum are mainly trivial ("all RC") with a SCORE of about 0.5. To the right of the maximum there is a change in cooperation; we see a steady partitioning into forward cogs composed of atoms: N1, I7, I8, H9, H10 and H11, and reverse cogs (atoms: H2, H3, H4 and C6).

In the last row of the table we see that the *bond* interactions lead to molecular cogs with high gear grinding, which is the cause of low SCOREs. The profile shows that these interactions lower the free energy gap in the $\xi_X \to \xi_Y$ transition by $-9.9 \frac{\text{kcal}}{\text{mol}}$. However, we can also see that the contribution from atoms C6, I7 and I8 alone increases this gap by $5 \frac{\text{kcal}}{\text{mol}}$. For the *ele* interactions the separation into molecular cogs was clean (low gear grinding), with a $-4.3 \frac{\text{kcal}}{\text{mol}}$ contribution from the forward cogs, and a $2.8 \frac{\text{kcal}}{\text{mol}}$ contribution from the reverse cogs due to the transition. For vdw, the separation has also led to low gear grinding, however the overall contribution from the forward cogs is much smaller than the one from the reverse cogs. This is an important observation which occurs again in the next section, where we analyze global molecular cogs determined for other model systems.

Results for related molecules

To better understand our approach, it is valuable to identify molecular cogs for other systems, related to the [NH3+]CC(I)I molecule scrutinized above. It was also valuable from the perspective of testing the transferability of the parameters used in the genetic clustering algorithm (as shown in the full results in the *Supporting Information*). We took into account molecules which instead of the [NH3+] group had the: NH2, CH3 and CH2Cl groups respectively, i.e. molecules: NCC(I)I, CCC(I) and CClCC(I)I. Partial charges assigned by the AM1-BCC procedure for these molecules are listed in Table 3.1. In this section we report and shortly discuss global molecular cogs for the *nbd*, *ele* and *vdw* interactions (see Table 3.4).

Because we analyzed closely related molecules, it was expected that molecular cogs for the nbd interactions would be comparable. All molecules presented here, except for CClCC(I)I, have partial charges of the same sign for corresponding atoms, therefore global molecular cogs for ele look similar. The more interesting outcome was related to discrepancies in molecular cogs identified for the vdw interactions.

As noted earlier, in the case of the [NH3+]CC(I)I molecule the overall vdw con-

tribution is almost entirely explained by cooperation within the reverse cogs. The forward cogs were identified merely because their contribution was non-positive. Nevertheless, this was a valuable insight – we have learned that the vdw steric effects are due to interactions of particular atoms: those which comprise the reverse cogs. On the other hand, the vdw molecular cogs for the NCC(I)I molecule are trivial ("all RC") and carry no such information. No non-trivial partitioning with a higher SCORE was found because there were no legitimate forward cogs, even as ineffective as those discovered in [NH3+]CC(I)I. This then suggests that we gained a simplified picture of vdw cooperation in the [NH3+]CC(I)I molecule simply because we were fortunate enough to have analyzed a system in which there had been at least a minimal contribution from the forward cogs. This is a consequence of the underlying assumption that a molecule's tendency to undergo a transition is a result of two opposing cooperations. Perhaps we should approach the problem differently for instances in which the molecule as a whole has the propensity to move forward/backward (as we also saw for the total and conf interactions).

Molecular cogs for vdw interactions for the CCC(I)I molecule are similar to those of [NH3+]CC(I)I. This might seem natural; the [NH3+] and CH3 groups share common properties. However, the energetic contribution profile for the vdw interactions (see $Supporting\ Information$) reveals that the forward cogs in CCC(I)I have a contribution comparable to that of the reverse cogs. Although vdw molecular cogs for CCC(I)I and [NH3+]CC(I)I are identical, the underlying mechanism is different.

The $\mathtt{CClCC}(\mathtt{I})\mathtt{I}$ model was designed to lower the ele barrier. However, the resulting molecular cogs for ele became trivial and, as in the case of vdw for $\mathtt{NCC}(\mathtt{I})\mathtt{I}$, we do not know which atoms are the main source of this effect. This again suggests that perhaps an alternative method of finding molecular cogs should be considered. In cases of trivial partitionings, such method should extract information about parts of the molecule which are the main source of the free energy difference. However, we leave investigation of properties and characteristics of alternative scoring functions for future studies.

DISCUSSION

Note that in all cases the genetic clustering algorithm gave clusterings with the best possible SCOREs. However, the efficiency of the genetic algorithm depends on the starting point, and without the help of the AP-generated initial population it took, on average, about 30 times longer, and the best result was not always achieved. To generate these initial solutions we adapted the AP clustering procedure (see Supporting Information).

The reason why conformational interactions lead to low-quality molecular cogs is that these are short-ranged interactions and our test molecular system is small. Partitioning of a graph into clusters is laden with a cost which depends on the weights of cross-cluster edges. But, as can be seen in Table 3.2, the integrated cooperation matrices for non-bonded interactions are more sparse than for confor-

mational interactions. Consequently, the cost of partitioning is greater for more "dense" matrices. But this cost decreases with increasing dimensionality of a matrix. It is therefore possible that, for more complex systems, molecular cogs for conformational interactions will have a higher SCORE.

The non-bonded interactions in the GAFF force field for atoms separated by three bonds of fewer are zero. This, and the fact that our model system is small, led to a sparse nbd cooperation matrix. Note, however, that a larger system may yield matrices that would be more "dense" for non-bonded interactions than for short-ranged conformational interactions. Whether a partitioning of these matrices would result in higher gear grinding remains an unanswered question.

We also anticipate that the entropic contribution in Equation (3.1) should play a more notable role for larger systems. Nonetheless, we have not yet considered clustering a graph with edges weighted by the entropic contributions of pairs of atoms. The reasons are twofold: practical (we want to avoid calculating Hessians of collective variables) and conceptual (the entropic cooperation of an atom with itself is non-zero). Entropic contributions need to be considered, but this should be the subject of future studies.

In the second part of the *Results* section we identified and analyzed molecular cogs for three additional molecules, related to the [NH3+]CC(I)I model. We discovered that trivial molecular cogs, which carry less information than any other partitioning, may occur whenever there is a lack of competition of cooperation within the molecule. This is a consequence of the proposed approach of dividing the system into two competing subsystems. In many cases in which global molecular cogs were trivial, we simply did not gain any insight into what is propelling the transformation. However, it seems that in such cases the analysis should be aimed at finding parts of a molecule that play the dominant role in its mobility.

Comparison with qualitative expectations

Results for the [NH3+]CC(I)I model, especially for the *ele* interactions, were useful in checking our intuitions against what was shown by the analysis. In this section we shortly discuss several sanity checks related to results for the *ele* interactions that were helpful in verifying whether our implementation had any critical errors, and whether the proposed approach leads to reasonable assessments.

From the plot depicting transient molecular cogs for ele (Table 3.3) we see that atoms N1 and I7 consistently hinder the transformation. These two atoms have negative partial charges (-0.85 and -0.08, respectively; see Table 3.1), and therefore repel each other throughout the transition. But the I8 atom (partial charge of -0.08) behaves differently, i.e. for ξ lower than -115° (free energy maximum) it impedes the process by repelling the N1 atom, and aids it for ξ larger than -115° . This effect was correctly captured by the above analysis, because the I8 atom finds itself in the reverse cogs in the first part of the transition, and in the forward cogs in the second part (as shown by the graph in the sixth row of Table 3.3). From the integrated contribution matrix for ele (second row in Table 3.3) we see, however, that the cumulative contribution from the N1-I8 atoms is positive, which results

in assigning them to one cluster.

The analysis revealed that atoms: H2, H3, H4 and C6 were consistently cooperating electrostatically, lowering the free energy barrier. This conclusion was more unexpected than the one concerning atoms: N1, I7 and I8, but was also compatible with our intuitions, taking into account the partial charges in Table 3.1.

Our method can suggest a qualitative interpretation of the cause behind a structural transition. However, it should be noted the most valuable information gained from this type of analysis is the quantitative description of the molecular $\cos via$ the energetic contribution profile.

CONCLUSIONS

The aim of this article was to introduce a new methodology of identifying molecular cogs – parts of a molecule that propel structural transitions in forward/backward directions along a collective variable. The current framework allowed us to track energetic contributions to the free energy, leaving the problem of including entropic terms for future developments. Results show that with the use of the genetic clustering algorithm we can successfully divide small molecules and identify forward and reverse molecular cogs associated with non-bonded interactions.

In particular, we proposed the approach of defining free energy contributions originating from pairs of atoms, and a method of dividing a molecule into molecular cogs. We showed that the proposed genetic clustering algorithm efficiently finds the optimal cogs, leading to high-quality partitionings for non-bonded interactions. However, we also found that conformational interactions lead to low-scoring molecular cogs, and that the system as a whole favored one meta-stable state over the other.

Currently, our method is based on cMD simulations for computing conditional averages (Equation (3.1)). Unfortunately, this solution has a critical drawback (see Supporting Information), but also, in order to determine the entropic contributions, requires computing second-order derivatives of the collective variable. To resolve this problem we should facilitate the Adaptive Biasing Force (ABF) scheme for calculating the PMF [15]. Specifically, our future work is aimed at reformulating the procedure as a plugin to the NAMD package (in which the ABF has already been implemented), to include entropic contributions and to assure scalable performance. Once this is done, numerous new opportunities will become available, some of which we mention below.

We constructed graphs in which nodes corresponded to atoms. Note, however, that they may also be assigned to amino acids (or more sizable objects) to construct graphs, which would lead to a more mosaic clustering. It could also be valuable to represent a whole ligand as a single object in a protein-ligand complex. One could also consider the role of functional groups comprising a ligand and amino acids in a binding pocket; it might then be helpful to represent the rest of the protein as a single node in the graph. Another example is a possible treatment of solvent molecules, for example: a particular group of interesting water molecules

could be transformed into separate nodes in the graph, while others reduced to a single node.

Our method may prove helpful in understanding why wild-type proteins perform better than mutants, or in explaining why certain drugs perform better than others, despite their structural similarity. It would also be interesting to consider a multistage induced fit docking, and to study cooperativity between amino acids and the ligand along a collective variable.

This research study demonstrates how to extract pair-wise energetic contributions to the free energy change along a reaction coordinate. In its present form¹, we could only use our method to analyze small, model molecules. However, we were able to verify the efficacy of the genetic clustering algorithm, and to learn what can be expected from the new notion of "molecular cogs". Our future aim is to analyze more complex systems, and to develop a publicly available implementation of our method for practical applications.

¹We made the source code available at: http://github.com/ponadto/molecular-cogs

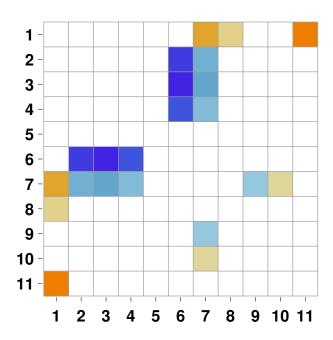


Figure 3.1: Graphical representation of a transient cooperation matrix for nbd interactions. Warm colors denote positive contributions, whereas blue – negative. White squares correspond to nil values.

[NH3+]CC(I)I		NCC(I)I		CCC(I)I		CClCC(I)I	
atom	partial	atom	partial	atom	partial	atom	partial
	charge		charge		charge		charge
N1	-0.85	N1	-0.92	C1	-0.10	C1	0.03
H2	0.47	H2	0.36	H2	0.04	CI2	-0.19
H3	0.47	Н3	0.36	H3	0.04	H3	0.08
H4	0.47			H4	0.04	H4	0.08
C5	0.13	C4	0.17	C5	-0.07	C5	-0.08
C6	0.08	C5	0.20	C6	0.21	C6	0.20
17	-0.08	16	-0.19	17	-0.19	17	-0.19
18	-0.08	17	-0.19	18	-0.19	18	-0.19
H9	0.13	H8	0.05	H9	0.06	H9	0.08
H10	0.13	H9	0.05	H10	0.06	H10	0.08
H11	0.13	H10	0.12	H11	0.10	H11	0.12

Table 3.1: Partial charges assigned by the AM1-BCC procedure.

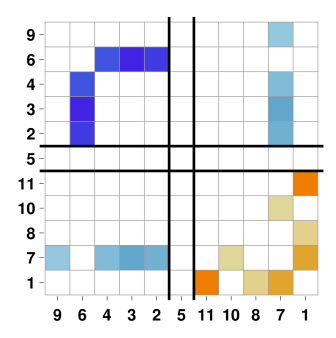


Figure 3.2: Graphical representation of a result of a clustering procedure. Rows and columns of the cooperation matrix in Figure 3.1 were reordered according to their assignments to: forward cogs, non-interacting atoms, and reverse cogs. In the upper left corner of this transformed matrix we have a block (square submatrix) with negative contributions, which come from the atoms 2, 3, 4 and 6 (from the forward cogs). In the lower right corner we have a block formed by pair contributions of the atoms in the reverse cogs. The gear grinding is small and results from contributions between the atom 7 and the atoms 2, 3, 4, 9.

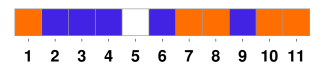


Figure 3.3: Graphical representation of an exemplary partitioning: $\{1, -1, -1, 0, -1, 1, 1, -1, 1, 1\}$. Orange squares correspond to 1 (reverse cogs), blue to -1 (forward cogs), and white to 0 (non-interacting). The cooperation matrix in Figure 3.1 was reordered according to this assignment, yielding the transformed cooperation matrix in Figure 3.2.

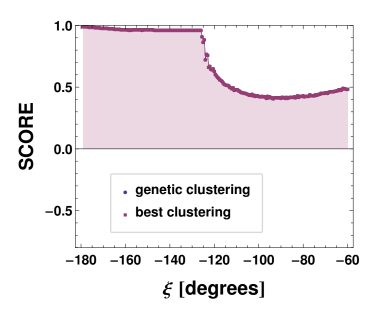


Figure 3.4: **SCORE plot for** *nbd* **interactions.** The plot shows that solutions found by the genetic clustering algorithm overlap perfectly with the highest scoring assignments to the transient molecular cogs.

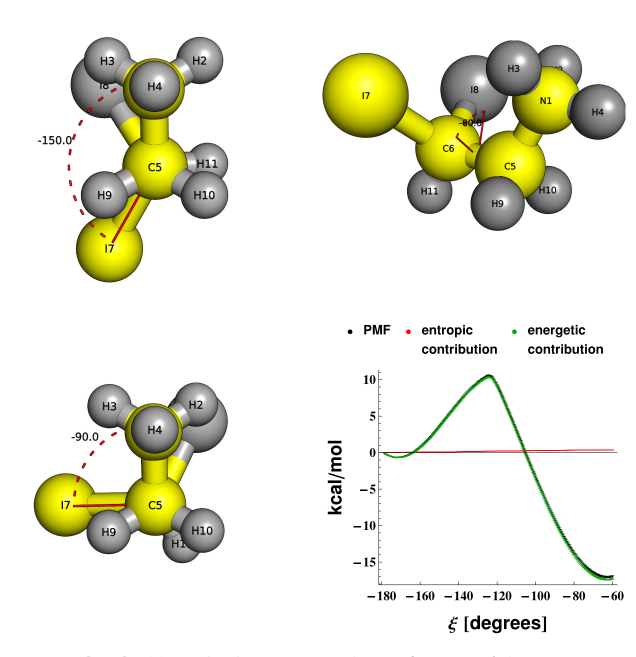


Figure 3.5: [NH3+]CC(I)I molecule summary. Three configurations of the molecule are shown, and the PMF with energetic and entropic contributions (for $T=300~\rm{K}$). Atoms N1-C5-C6-I7 (yellow) were used to define the dihedral angle, ξ .

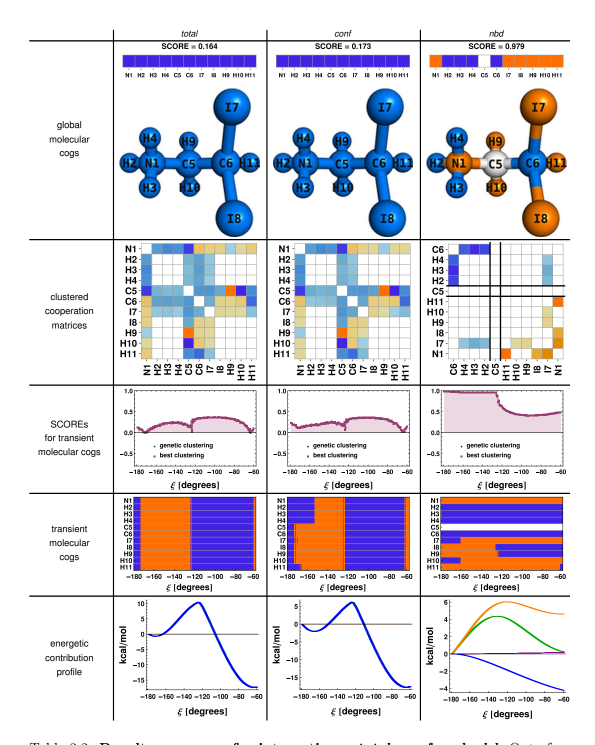


Table 3.2: Results summary for interactions: total, conf and nbd. Out of the triple: total, conf and nbd, only the last one leads to a high SCORE partitioning, yielding non-trivial molecular cogs. Colors: green, blue, orange and purple in the last row of the table denote contributions from the: whole system, forward cogs, reverse cogs and gear grinding, respectively (see the *Overview* section for details).

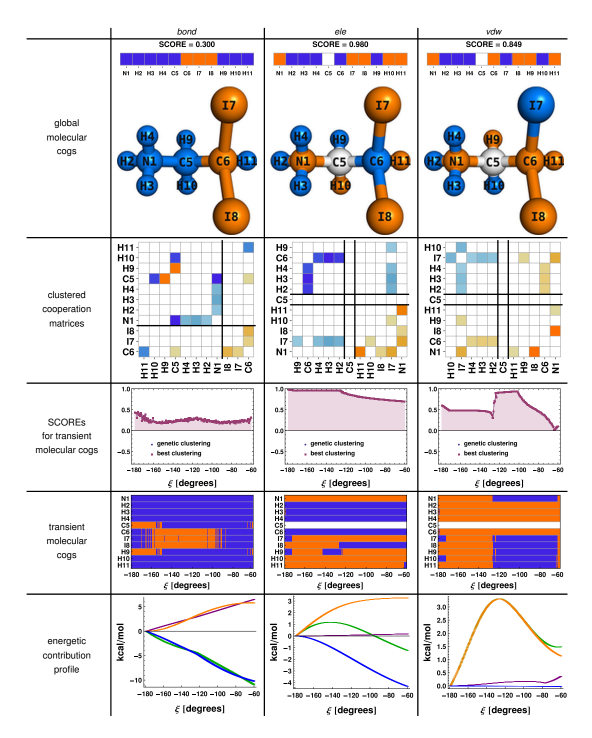


Table 3.3: Results summary for interactions: bond, ele and vdw. All three 2-atom interaction types lead to decompositions into non-trivial molecular cogs. However, the transient molecular cogs have high SCOREs only for the ele interactions. Colors: green, blue, orange and purple in the last row of the table denote contributions from the: whole system, forward cogs, reverse cogs and gear grinding, respectively (see the *Overview* section for details).

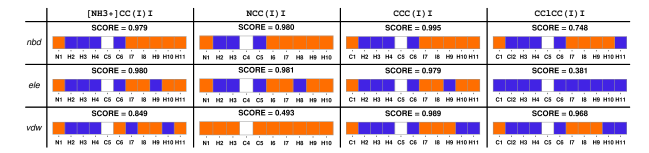


Table 3.4: Global molecular cogs for the following molecules: [NH3+]CC(I)I, NCC(I)I, CCC(I)I and CC1CC(I)I. We focused our discussion on partitionings into forward and reverse cogs for the *nbd*, *ele* and *vdw* interactions. Full results are detailed in the *Supporting Information*.

Chapter 4

Conclusions

In this closing chapter we re-iterate the most important findings of the methods presented in Chapters II and III, and propose several refinements to these methods.

4.1 Dynamic domains identification

ResiCon was built upon several ideas, some of which were essential for its high-quality results. The first one made use of the concept of a *contact* between residues (that had been proven to be advantageous in an earlier application) to define the *geometrical variability*. Thereupon, the second idea was to propose a specific *spectral clustering* algorithm, whose output were clusters corresponding to dynamic domains of a given protein structure. In its current implementation, ResiCon requires information about all pairs of configurations to calculate the geometrical variability.

To analyze MD simulations, ResiCon made use of the Principal Components Analysis (PCA) to select representative configurations. Applying PCA was necessary, because otherwise each pairs of configurations in the trajectory would have to be tracked for the similarity matrix to be constructed. However, the PCA relies on linearity assumptions, which in case of structural transitions of biomolecular systems may seem overly optimistic. It is possible to amend this obstacle by employing a non-linear algorithm (e.g., NLPCA, which stands for *Non-Linear PCA*), but at prohibitively high computational cost.

In order to truly adapt ResiCon to the MD trajectory set-up, one needs to amend the problem of analyzing each pair of configurations. In practice, only a few pairs contributed (or, more accurately: *could* have contributed) to the similarity matrix. It would certainly be worthwhile to test a sub-procedure that might substitute PCA by finding relevant pairs of configurations. In particular, the Locality-Sensitive Hashing (LSH) algorithm offers this kind of functionality.

4.2 Molecular cogs

The fundamental assumption of the molecular cogs methodology presented in Chapter III was that there are two, distinct groups of atoms, cooperating to propel the system forward (or backward) along a given collective variable. However, we concluded that the clustering procedure has its limitations in cases in which the molecular system as a whole has a general propensity to move in a certain direction. In such cases, the more valuable insight would be the knowledge about which group of atoms plays the dominant role in such an effect.

Instead of using the definitive approach of clustering, one might consider employing a more a general idea in which each element can be a member of a more than just one group. *Community detection* offers such functionality, but also allows for the identification of hierarchical structures. With the help from a community detection algorithm, we could, for example, search for groups of atoms with different degrees of contribution to the system's tendency to make a transition.

From a different stand point, the method presented in Chapter III employed constrained molecular dynamics (cMD) to estimate local changes in the Helmholtz free energy. However, as we noticed in Chapter III, the cMD has limitations, which would render it ineffective for large systems. An alternative sampling technique – most notably: the Adaptive Biasing Force (ABF) method – could remedy this impediment, but at the cost of a more complicated and time-demanding implementation. We left the ABF-driven improvement for a future molecular cogs identification scheme.

The ABF sub-procedure would also alleviate the problem of estimating entropic contributions of the molecular cogs. Nevertheless, we are not yet certain how the entropic contribution matrix should be partitioned. We need to re-evaluate the meaning of molecular cogs altogether, and decide on a strategy that would treat entropic and energetic contributions on an equal footing.

The molecular cogs setting in its current formulation suggests a more subtle analysis of the causality hidden in complex molecular transformations. We noticed that by modeling the contributions to the PMF we are close to modeling the kinetics of a transition, as well. That is, we can modify our approach to identify contributions of a given subsystem to the so-called *reaction rate constant*. For that, we would need to equip our analytical toolbox with yet another method called the Markov State Models (MSM). Proposed by Frank Noe, the MSM extracts kinetic properties of a system from the PMF. It was proven to give reliable estimates of the reaction rate constant for small enzymes, and seems to be an interesting future improvement to our molecular cogs methodology.

4.3 Final comments

Bibliography

- [1] DM van Aalten et al. "Engineering protein mechanics: inhibition of concerted motions of the cellular retinol binding protein by site-directed mutagenesis." In: *Protein engineering* 10.1 (1997), pp. 31–37.
- [2] Yelena A. Arnautova, Ruben A. Abagyan, and Maxim Totrov. "Development of a new physics-based internal coordinate mechanics force field and its application to protein loop modeling". In: *Proteins: Structure, Function, and Bioinformatics* 79.2 (2011), pp. 477–498.
- [3] Ivet Bahar, Ali Rana Atilgan, and Burak Erman. "Direct evaluation of thermal fluctuations in proteins using a single-parameter harmonic potential". In: Folding and Design 2.3 (1997), pp. 173–181.
- [4] G Bao. "Protein mechanics: a new frontier in biomechanics". In: Experimental mechanics 49.1 (2009), pp. 153–164.
- [5] Stefan Bernhard and Frank Noé. "Optimal identification of semi-rigid domains in macromolecules from molecular dynamics simulation". In: *PloS one* 5.5 (2010), e10491.
- [6] Stefan Boresch and Martin Karplus. "The meaning of component analysis: decomposition of the free energy in terms of specific interactions". In: *Journal of molecular biology* 254.5 (1995), pp. 801–807.
- [7] Peer Bork. "Shuffled domains in extracellular proteins". In: *FEBS letters* 286.1 (1991), pp. 47–54.
- [8] G Patrick Brady and Kim A Sharp. "Decomposition of interaction free energies in proteins and other complex systems". In: *Journal of molecular biology* 254.1 (1995), pp. 77–85.
- [9] ZIMEI Bu and DJ Callaway. "Proteins move! Protein dynamics and long-range allostery in cell signaling". In: *Adv Protein Chem Struct Biol* 83 (2011), pp. 163–221.
- [10] EA Carter et al. "Constrained reaction coordinate dynamics for the simulation of rare events". In: Chemical Physics Letters 156.5 (1989), pp. 472–477.

- [11] Harry Chiang et al. "Molecular mechanics and dynamics characterization of an in silico mutated protein: A stand-alone lab module or support activity for in vivo and in vitro analyses of targeted proteins". In: *Biochemistry and Molecular Biology Education* 41.6 (2013), pp. 402–408.
- [12] Christophe Chipot. "Frontiers in free-energy calculations of biological systems". In: Wiley Interdisciplinary Reviews: Computational Molecular Science 4.1 (2014), pp. 71–89.
- [13] Christophe Chipot and Andrew Pohorille. Free energy calculations: theory and applications in chemistry and biology. Vol. 86. Springer, 2007.
- [14] Rowena Marie Cole. Clustering with genetic algorithms. Citeseer, 1998.
- [15] Jeffrey Comer et al. "The Adaptive Biasing Force Method: Everything You Always Wanted To Know but Were Afraid To Ask". In: *The Journal of Physical Chemistry B* (2014).
- [16] Paweł Daniluk and Bogdan Lesyng. "A novel method to compare protein structures using local descriptors". In: *BMC bioinformatics* 12.1 (2011), p. 344.
- [17] Paweł Daniluk and Bogdan Lesyng. "Theoretical and Computational Aspects of Protein Structural Alignment". In: Computational Methods to Study the Structure and Dynamics of Biomolecules and Biomolecular Processes. Springer, 2014, pp. 557–598.
- [18] Ken A Dill. "Additivity principles in biochemistry". In: *Journal of Biological Chemistry* 272.2 (1997), pp. 701–704.
- [19] Bela Farago et al. "Activation of nanoscale allosteric protein domain motion revealed by neutron spin echo spectroscopy". In: *Biophysical journal* 99.10 (2010), pp. 3473–3482.
- [20] Darón I Freedberg et al. "Rapid structural fluctuations of the free HIV protease flaps in solution: relationship to crystal structures and comparison with predictions of dynamics calculations". In: Protein science 11.2 (2002), pp. 221–232.
- [21] Brendan J Frey and Delbert Dueck. "Clustering by passing messages between data points". In: *science* 315.5814 (2007), pp. 972–976.
- [22] Alessandro Genoni, Giulia Morra, and Giorgio Colombo. "Identification of domains in protein structures from the analysis of intramolecular interactions". In: *The Journal of Physical Chemistry B* 116.10 (2012), pp. 3331–3343.
- [23] A Gorecki, J Trylska, and B Lesyng. "Causality and correlation analyses of molecular dynamics simulation data". In: From Computational Biophysics to Systems Biology (CBSB07), volume NIC Series 36 (), pp. 25–30.
- [24] Adam Górecki et al. "RedMD—reduced molecular dynamics package". In: Journal of computational chemistry 30.14 (2009), pp. 2364–2373.
- [25] Barry J Grant et al. "Bio3d: an R package for the comparative analysis of protein structures". In: *Bioinformatics* 22.21 (2006), pp. 2695–2696.

- [26] James C Gumbart, Beno^t Roux, and Christophe Chipot. "Standard binding free energies from computer simulations: What is the best strategy?" In: Journal of chemical theory and computation 9.1 (2012), pp. 794–802.
- [27] Donald Hamelberg and J Andrew McCammon. "Fast peptidyl cis-trans isomerization within the flexible Gly-rich flaps of HIV-1 protease". In: *Journal of the American Chemical Society* 127.40 (2005), pp. 13778–13779.
- [28] Jiawei Han and Micheline Kamber. "Data mining: concepts and techniques". In: *United States of America: Morgan Kauff mann Publishers* (2001).
- [29] Steven Hayward and Herman JC Berendsen. "Systematic analysis of domain motions in proteins from conformational change: New results on citrate synthase and T 4 lysozyme". In: Proteins Structure Function and Genetics 30.2 (1998), pp. 144–154.
- [30] Steven Hayward, Akio Kitao, and Herman JC Berendsen. "Model-free methods of analyzing domain motions in proteins from simulation: a comparison of normal mode analysis and molecular dynamics simulation of lysozyme". In: *Proteins: Structure, Function, and Bioinformatics* 27.3 (1997), pp. 425–437.
- [31] Konrad Hinsen. "Analysis of domain motions by approximate normal mode calculations". In: *Proteins Structure Function and Genetics* 33.3 (1998), pp. 417–429.
- [32] Wolfgang Kabsch. "A solution for the best rotation to relate two sets of vectors". In: Acta Crystallographica Section A: Crystal Physics, Diffraction, Theoretical and General Crystallography 32.5 (1976), pp. 922–923.
- [33] Donata K Kirchner and Peter Güntert. "Objective identification of residue ranges for the superposition of protein structures". In: *BMC bioinformatics* 12.1 (2011), p. 170.
- [34] John G Kirkwood. "Statistical mechanics of fluid mixtures". In: *The Journal of Chemical Physics* 3.5 (1935), pp. 300–313.
- [35] Richard Lavery and Sophie Sacquin-Mora. "Protein mechanics: a route from structure to function". In: *Journal of biosciences* 32.1 (2007), pp. 891–898.
- [36] Richard A Lee, Moe Razaz, and Steven Hayward. "The DynDom database of protein domain motions". In: *Bioinformatics* 19.10 (2003), pp. 1290–1291.
- [37] Gary E Martin, Andrew S Zektzer, et al. Two Dimensional NMR Methods for Establishing Molecular Connectivity. VCH, 1988.
- [38] Marina Meilă. "Comparing clusterings—an information based distance". In: Journal of multivariate analysis 98.5 (2007), pp. 873–895.
- [39] Noel M O'Boyle et al. "Open Babel: An open chemical toolbox". In: *J Cheminf* 3 (2011), p. 33.
- [40] Xavier Periole et al. "Structural determinants of the supramolecular organization of G protein-coupled receptors in bilayers". In: *Journal of the American Chemical Society* 134.26 (2012), pp. 10959–10965.

- [41] Raffaello Potestio, Francesco Pontiggia, and Cristian Micheletti. "Coarse-grained description of protein internal dynamics: an optimal strategy for decomposing proteins in rigid subunits". In: *Biophysical journal* 96.12 (2009), pp. 4993–5002.
- [42] William H Press. Numerical recipes 3rd edition: The art of scientific computing. Cambridge university press, 2007.
- [43] Jane S Richardson. The anatomy and taxonomy of protein structure. Vol. 34. Academic Press, 1981.
- [44] Julia Romanowska, Krzysztof S Nowinski, and Joanna Trylska. "Determining geometrically stable domains in molecular conformation sets". In: *Journal of Chemical Theory and Computation* 8.8 (2012), pp. 2588–2599.
- [45] Sophie Sacquin-Mora. "Motions and mechanics: investigating conformational transitions in multi-domain proteins with coarse-grain simulations". In: *Molecular Simulation* 40.1-3 (2014), pp. 229–236.
- [46] S Kashif Sadiq and Gianni De Fabritiis. "Explicit solvent dynamics and energetics of HIV-1 protease flap opening and closing". In: *Proteins: Structure, Function, and Bioinformatics* 78.14 (2010), pp. 2873–2885.
- [47] Christian Seifert and Frauke Gräter. "Protein mechanics: How force regulates molecular function". In: *Biochimica et Biophysica Acta (BBA)-General Subjects* 1830.10 (2013), pp. 4762–4768.
- [48] Anton V Sinitskiy, Marissa G Saunders, and Gregory A Voth. "Optimal number of coarse-grained sites in different components of large biomolecular complexes". In: *The Journal of Physical Chemistry B* 116.29 (2012), pp. 8363–8374.
- [49] David A Snyder and Gaetano T Montelione. "Clustering algorithms for identifying core atom sets and for assessing the precision of protein structure ensembles". In: *Proteins: Structure, Function, and Bioinformatics* 59.4 (2005), pp. 673–686.
- [50] Daniel Taylor, Gavin Cawley, and Steven Hayward. "Classification of domain movements in proteins using dynamic contact graphs". In: (2013).
- [51] Glenn M Torrie and John P Valleau. "Nonphysical sampling distributions in Monte Carlo free-energy estimation: Umbrella sampling". In: *Journal of Computational Physics* 23.2 (1977), pp. 187–199.
- [52] Jiri Vondrasek and Alexander Wlodawer. "HIVdb: a database of the structures of human immunodeficiency virus protease". In: *Proteins: Structure*, Function, and Bioinformatics 49.4 (2002), pp. 429–431.
- [53] Marcus Weber, Wasinee Rungsarityotin, and Alexander Schliep. *Perron cluster analysis and its connection to graph partitioning for noisy data*. Konrad-Zuse-Zentrum für Informationstechnik Berlin, 2004.

- [54] Hyung-June Woo and Beno[†]t Roux. "Calculation of absolute protein–ligand binding free energy from computer simulations". In: *Proceedings of the national academy of sciences of the united states of america* 102.19 (2005), pp. 6825–6830.
- [55] Willy Wriggers and Klaus Schulten. "Protein domain movements: detection of rigid domains and visualization of hinges in comparisons of atomic coordinates". In: *Proteins Structure Function and Genetics* 29.1 (1997), pp. 1–14.
- [56] Yuzhen Ye and Adam Godzik. "Flexible structure alignment by chaining aligned fragment pairs allowing twists". In: *Bioinformatics* 19.suppl 2 (2003), pp. ii246–ii255.
- [57] Semen O Yesylevskyy, Valery N Kharkyanen, and Alexander P Demchenko. "Dynamic protein domains: identification, interdependence, and stability". In: *Biophysical journal* 91.2 (2006), pp. 670–685.
- [58] Zhiyong Zhang et al. "Defining coarse-grained representations of large biomolecules and biomolecular complexes from elastic network models". In: *Biophysical journal* 97.8 (2009), pp. 2327–2337.