Computational Analysis of Single Molecule Force Spectroscopy Experiments on Native Membranes

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Chapter 1

Introduction

Every living cell is surrounded by a biological membrane that separates the cell’s inner content from the outside world. The basic structural unit of every membrane is the lipid bilayer, which is built from phospholipids, sphingolipids and sterols. Phospholipids and sphingolipids are amphiphatic molecules, which means that their molecules have hydrophilic (water-loving) head-groups and hydrophobic (water-fearing) tails. The specific chemical properties of these lipid molecules determine the characteristic structure of the lipid bilayer: the polar head-groups face the outer environment of the cell on both sides, while the hydrophobic tails form the interior of the bilayer (see Figure 2.1).

The main functional components of the cell membrane are the membrane proteins, which are embedded in the lipid bilayer. Membrane proteins are active participants in the most important biological processes in the cell membrane, like active transport of ions and molecules across the membrane, signal transduction and cell-cell communication. Nearly 30% of all proteins in eukaryotic cells are membrane proteins. They are the targets of more than 50% of modern medicinal drugs and many diseases are found to be related to membrane proteins misfolding [1]. Despite their significance, the number of three-dimensional membrane protein structures remains small. Currently, there are 827 unique structures of membrane proteins available [2]. The reason is that the standard experimental techniques used for studying the structure of soluble proteins (X-ray crystallography, NMR spectroscopy and cryo-electron microscopy (cryo-EM)) experience difficulties when it comes to membrane proteins. The main problems are associated with membrane proteins extraction, solubilization, and purification. For example, the solubilization and purification of a specific membrane protein require the selection of a suitable detergent and that step is critical [3].

Atomic force microscopy (AFM) turned out to be a promising alternative to the conventional experimental methods used for studying proteins. Developed initially to image surfaces with atomic resolution, the AFM method went far beyond the expectations. The atomic force microscope is a scanning probe microscope, namely a device that uses a physical probe to scan the sample surface producing a topographical image of the surface. The probe in this case is the tip, which is the most
important component of the microscope because it is the only component which physically touches the sample. The AFM tip is mounted on the end of a flexible cantilever (see Figure 2.8a). Once brought in contact with the sample, the tip raster scans the surface and experiences the attraction forces acting between its sharp edge and the sample surface. Accordingly, the spring-like cantilever bends. A laser beam pointed to the cantilever bounces off its back onto a position-sensitive photo detector and the deflection of the light is measured with high precision. The roughness of the sample surface induces changes in the cantilever deflection and its measurement and processing shapes the final topographical image. In addition to this, the interaction forces between the tip and the sample can be estimated from the deflections through Hooke’s law. AFM is used in a wide range of disciplines, from solid state physics to medicine, from the study of quantum dots to the imaging of living cells. What makes this method extremely suitable for biological samples is the fact that AFM can operate in water solution, in physiological conditions and at body temperature. This allowed using AFM for imaging of native cell membranes [4].

Apart from the imaging mode, AFM quickly became one of the most common techniques used for single molecule force spectroscopy (SMFS) experiments. In this kind of experiments a single molecule, like the double-helix of DNA, a protein or a polysaccharide, is directly manipulated with a probe, usually of microscopic size [5], obtaining information about its structural and dynamical properties in real time and at the single molecule level. In AFM-based SMFS, the microscope is used as a pico-Newton force measuring device [6]. The protein molecule is on one side bound to the surface, while on the other side it gets picked by the AFM tip and stretched as the tip retracts from the surface (see Figure 2.9a). The molecular force response versus the distance between the tip and the sample surface is recorded and presented in the form of a force-extension (F-x) curve (see Figure 2.9b). F-x curves, known also as traces, are direct probes of the protein’s unfolding pathways. These curves store the information about the unfolding of the protein domains, their mechanical stability, the order in which they unfold, etc. This information has been used to understand better protein folding[7]. These studies highlighted a very important fact: that the force pattern contained in these curves is a fingerprint of the protein under examination. For example, the mechanical unfolding of the multidomain protein titin [8], responsible for the passive elasticity of muscle cells, results in F-x curves bearing a characteristic sawtooth pattern in which the number of force peaks is equal to the number of immunoglobulin domains (see Figure 2.10). The order in which the individual domains unfold is governed by the strength of the interactions which hold them, thus the weakest domains unfold first. This example is a clear demonstration of the enormous potentialities of this technique.

These potentialities are exploited at best for studying membrane proteins, where, as we have seen, other methods often fail. AFM allows both imaging of the cell membrane and manipulating single membrane proteins embedded in it. All of this is accomplished under physiological conditions. The number of membrane proteins unfolded in AFM-SMFS experiments is rapidly growing. The list includes the light-sensitive receptor proteins bacteriorhodopsin and rhodopsin [9, 10, 11, 12], the
CNGA1 channel [13, 12], the Na+/K+ antiporter [14], the leucine-binding protein [15] and many others. The first membrane protein studied with AFM-SMFS was bacteriorhodopsin (bR). This was done in a work by Muller et al. [9] dealing with the unfolding pathways of bR. In this study, the AFM was used at the same time for imaging the membrane and unfolding of the bR present in it (see Figure 2.11). The obtained F-x patterns revealed pairwise unfolding of the transmembrane helices of bR. However, rather surprisingly, in this experiment not all the F-x curves looked similar. In some of them, two of the helices were unfolding one after the other, which indicates different unfolding pathways. This illustrates that AFM can also be used to detect different unfolding pathways of the same protein.

This technique has not yet shown all its potentialities, and new avenues are opened almost every year. Recent advances in AFM-SMFS enabled the acquisition of a huge amount of data in reasonable time [16, 17]. Moreover, this has been achieved with experiments performed directly in native cell membranes under physiological conditions [18]. The obtained data are highly heterogeneous, the amount of high-quality traces is very low and they come from the unfolding of different proteins with different size. Between 1,000 and 10,000 F-x curves are usually generated in a single experiment. The availability of this huge amount of experimental data encouraged the development of novel automatic tools promoting its processing with the least possible manual intervention. Several tools aimed at this scope have already been proposed [19, 20, 21, 22]. These approaches work very well for membrane proteins of the same type or for samples with well-known protein composition. However, the same approaches have some serious limitations when it comes to data sets with unknown protein composition like those collected in native membranes. The main issue with AFM-SMFS data is that in < 1% of the cases membrane proteins are completely unfolded [21]. This means that in the large bulk of data, only a tiny portion of traces is worth looking at. Not only this, but the majority of the traces do not contain any unfolding events. This makes the careful preprocessing of the raw data a crucially important step. All the available methods are based on knowing the approximate length of the fully-stretched protein under investigation. This information allows the selection of F-x curves having their last force peak at extension values comparable with the protein length. A solution of this kind is useful in AFM experiments performed in controlled environments where a certain protein of interest is unfolded. However, if we want to analyze data coming from experiments in native cell membranes, this approach is incapable of spanning all the possibilities simply because the membrane for sure contains proteins unidentified so far. Ideally, an appropriate method for analysis AFM-SMFS data coming from native cell membranes should be able to select in an unsupervised manner all traces which contain successful unfolding events and to group those that are similar to each other in separate clusters. In this way, all the meaningful data will be extracted and presented in a human-readable manner, easier to interpret.

This thesis was motivated by developing tools for interpreting AFM-SMFS experiments on membrane proteins performed in native cell membranes. In particular, we address two problems. The
first one deals with the effects of lipid composition of different membranes on the mechanical sta-
bility and unfolding pathway of membrane proteins. The second one deals with the analysis of
the huge amount of data generated by AFM-SMFS experiments in native cell membranes.

In Chapter 2 we provide a general overview on cell membranes and a brief description of the
main components they contain. Our focus is on membrane proteins embedded in the cell membrane,
known also as integral membrane proteins, since they are the molecular objects of this study. We
describe the two basic secondary motifs by which they are built: the α-helix and the β-barrel. We
discuss two classifications of membrane proteins: topology-based and function-based. In subsection
2.0.3 we shortly explain how the AFM imaging mode works, while in subsection 2.0.4 we comment
its performance in the SMFS mode, providing a description of the experiments. After that, we
discuss molecular models used to describe these experiments. Finally, we make an overview of some
of the currently available procedures for analysis of AFM-SMFS data.

The work presented in Chapter 3 was motivated by AFM-SMFS experiments performed in the
discs and the plasma membrane of the rod OS, where rhodopsin is the dominant protein. The
obtained F-x curves from pulling rhodopsin from the discs and from the plasma membrane re-
vealed two strikingly different unfolding patterns [12]. Moreover, rhodopsin in the discs initiates
the phototransduction cascade, while rhodopsin in the plasma membrane does not. The discs and the
plasma membrane have different lipid composition [23]. The plasma membrane has higher choles-
terol concentration. This suggests that the different lipid environment affects both the rhodopsin
mechanical properties and its function. In order to test this hypothesis we used a simple topology-
based coarse-grained model of the protein-membrane system. The protein molecule was described
with a coarse-grained Go-like model initially developed by Cieplak et al. [24] to study the mechanical
unfolding of soluble proteins [25, 26, 27]. This model successfully reproduced the experimental F-x
curves and provided an important insight in the interpretation of these spectra. A modified version
of this model was applied also to membrane proteins, to bacteriorhodopsin in particular [28]. When
it comes to the unfolding of membrane proteins, the proper modeling of the system becomes much
more demanding in comparison to soluble proteins. Membrane proteins do not only get stretched
but they also get extracted out of the membrane. This necessarily requires taking the membrane
into account in the model. Cieplak et al. [28] did this explicitly representing the membrane with a
lipid bilayer, modeled in a coarse-grained manner. When the protein is pulled the membrane is kept
frozen: as a portion of the polypeptide chain gets extracted, the space it was occupying remains
empty, leaving a hole in the membrane. In simple terms, the lipid bilayer does not adjust to the
new configuration of the system. This approximation is in our opinion unrealistic. To model more
accurately the unfolding of a membrane protein we developed a new approach in which we model
the effect of the membrane by adding to the original Go model of Cieplak et al. [24] an extra poten-
tial energy term $V_{MEMBR}$. The potential operates in a different manner on the different residues
depending on their hydrophobicity and the contact they form with the membrane in the native
configuration. When a hydrophobic residue forms a native contact with the membrane and it is positioned inside the membrane, $V_{MEMBR}$ is 0. If that same residue is in a region corresponding to the region occupied by the polar head-groups of the lipids, it gets moderately unfavored. If the residue is outside the membrane and is water-exposed, it gets a full penalty, which is defined by the only important parameter of the model, $\epsilon_{MEMBR}$. This parameter determines the strength of the membrane potential. Furthermore, by varying $\epsilon_{MEMBR}$ one can model different lipid compositions of the membrane. For example, larger values of $\epsilon_{MEMBR}$ specify a more hydrophobic membrane like a cholesterol-rich membrane.

Figure 1.1: (a,c) Simulated force-distance traces for bovine rhodopsin (PDB code: 1U19) pulled by the C-terminal at $k_BT = 0.52\epsilon$ for two different values of the parameter $\epsilon_{MEMBR}$ as indicated ((a) 4.03$\epsilon$ and (c) 10$\epsilon$). Each plot represents the superposition of 10 traces obtained from 10 independent simulations. (b,d) Cartoon representations of the order in which the transmembrane helices unfold in the simulations of the left panels, as derived by a visual inspection of the trajectories. The color map is the same as for the traces. The numbers on top of each peak correspond to the length of the stretch that is unfolded up to the time step when the force drops (expressed in number of amino acids, n). The traces presented in panel a reproduce the experimental traces obtained for the unfolding of rhodopsin in the discs of the rod outer segment (OS). The traces in panel c reproduce the experimental traces obtained for the unfolding of rhodopsin in the plasma membrane of the rod OS.
Changing the values of $\epsilon_{MEMBR}$ in our model is like changing the hydrophobicity of the membrane arising from its different lipid composition. We performed stretching MD simulations with four different values of $\epsilon_{MEMBR}$. With our simple model, we were able to qualitatively reproduce the difference in the experimental curves obtained from unfolding of rhodopsin in the discs and in the plasma membrane. In Figure 1.1 we show F-x curves predicted by our model together with cartoon representations of rhodopsin colored in correspondence with the order of unfolding of the single units. The superimposition of simulated traces depicted in Figure 1.1a is in good agreement with the experimental curves from unfolding of rhodopsin from the discs, while the superimposition of simulated traces in Figure 1.1c corresponds to the experimental curves from unfolding of rhodopsin from the plasma membrane. These results support the hypothesis that the reason we observe different unfolding patterns of rhodopsin from the discs and from the plasma membrane is in the different lipid composition implying different membrane hydrophobicity captured in our model by the value of $\epsilon_{MEMBR}$.

In Chapter 3 we also attempt to model and understand the inactivation of rhodopsin in the plasma membrane. Since the main difference between the plasma membrane and the discs is the higher cholesterol concentration of the former, we decided to check if the conformation with a larger membrane-exposed hydrophobic area would be favored in a more hydrophobic membrane. Our coarse-grained model is a Go-model, and therefore cannot be used to describe conformational changes. Therefore, it is not appropriate for addressing this problem. Instead, we used all-atom MD simulation of rhodopsin embedded in a DPPC bilayer in two different conformations relevant to the light-harvesting cycle. To evaluate the effect of cholesterol on rhodopsin’s flexibility we used free energy perturbation theory. The final results suggest that the higher cholesterol concentration of the plasma membrane favors the inactive rhodopsin conformation hence preventing rhodopsin from activation. These results confirm the utmost influence of membrane composition on the mechanical properties of membrane proteins.

Chapter 4 describes the main contribution of this Thesis. We there describe a fully-automatic procedure for the analysis of F-x curves coming from experiments performed in native cellular membranes. This work is motivated from the recent development of techniques for performing AFM-SMFS experiments on native membrane patches [18]. The obtained data is highly heterogeneous and presents some serious challenges to data analysis. In the available methods [19, 20, 21, 22] the analysis is guided by knowledge on the protein sample composition. In native membrane patches this information is simply not available. Another issue is determined by the extremely small amount of high-quality traces in these data sets. Here by high-quality traces, we mean F-x curves associated with meaningful unfolding events.

To address these issues, we developed an automatic procedure that does not require knowledge on the protein sample composition and is able to extract high-quality traces from the data bulk in an unsupervised manner. The idea is to be able to distinguish different proteins based on the
Figure 1.2: Block diagram of the algorithm developed by us.
characteristic unfolding patterns observed in their F-x curves without knowing who these proteins are. In Figure 1.2 we show a block diagram representing the procedure. Initially, we process all traces removing the noisy parts of the signal, aiming at obtaining only this part of the F-x curve that contains the unfolding pattern. This step corresponds to the "Cutting & Filtering" block in Figure 1.2. The nature of the data requires alignment of the F-x curves to overcome horizontal shifts due to the different tip-protein attachment positions. For this scope, we use dynamic programming alignment, which gives an alignment score that provides a measure of the similarity between two traces. A pair of similar traces is characterized by a high alignment score. This alignment approach has been already applied to SMFS data and described in the literature by Marsico et al. [20]. In order to extract high-quality traces from the large data set, we assigned a quality score to each trace and used that for selecting the meaningful traces. The quality score measures the consistency of each trace with a polymer physics model called the worm-like chain (WLC) model (see subsection 2.0.4). The WLC model has proved to provide a reliable interpretation of the experimental F-x curves and moreover, a quantitative one. The WLC consistency score evaluates the quality of a trace. If the score is high, the trace is good; if the score is low, the trace is bad. Once we have selected good traces, based on their quality score we would like to group them into clusters based on the unfolding patterns they contain. In order to do this, we use density-peak clustering [29], a clustering approach which robustly recognizes groups of data points belonging to separate peaks in the probability distribution. The crucial ingredient of this approach is the distance between two data points, in our case two AFM-SMFS traces. We defined a distance which combines the alignment score and the quality score. According to this distance, high-quality traces similar to each other have a small distance. Instead, low-quality traces, even if they are similar to each other, have a larger distance. This guarantees that the clusters we obtain contain traces which are not only similar to each other (which is the case in the work of Marsico et al. [20]) but which are also both of a high quality, in terms of consistency with the WLC model. We will show how important it is using a distance with these properties.

We tested our method on a data set containing ∼ 100 manually selected traces corresponding to the unfolding of the CNGA1 channel and ∼ 4,000 traces of unidentified origin and quality. Our method was able to detect the CNG traces and to put them in a separate cluster. Remarkably, the method was also able to find other CNG traces that escaped the manual selection and to assign them to the CNG cluster. In addition, we obtained other clusters whose molecular origin we are currently unable to identify.

We also applied our procedure on a data set containing ∼ 400,000 traces from pulling experiments in the plasma membrane of the rod outer segment (OS). This large data set presents a challenge for every currently available tool for SMFS data analysis, including our method. The time it took to our program to analyze this huge amount of traces on a workstation with 16 CPUs is ∼ 90 minutes. Not surprisingly, only ∼ 5 % of all traces passed the selection procedure. The plasma membrane
of the rod OS hosts primarily rhodopsin and CNG channels, thus we expect to obtain clusters corresponding to the unfolding these two proteins. Two of the clusters contain decent candidates for the unfolding of rhodopsin, but we couldn’t find a cluster corresponding to the CNG channel. We assume that the reason for this is that the number of good CNG traces that are similar to each other is insufficient to form a cluster. To test this hypothesis we added the manually selected CNG traces from the previous data set to this data set and applied our procedure. As a final result we obtained an additional cluster containing the manually selected CNG traces supporting our hypothesis. We looked also at the other clusters and indeed, the number of cluster members very similar to the cluster center is very small suggesting that not only the CNG traces similar to each other are very little but the number of high-quality traces similar to each other is very low. This seriously troubles the clusters identification. We need more data sets from native membranes in order to further validate this hypothesis.
Chapter 2

AFM-SMFS of membrane proteins: an overview

2.0.1 Cell Membranes

Every living cell is surrounded by a biological membrane that separates the cell’s inner content from the outside world. This biological membrane is not an inactive barrier but a protection sheath through which transport of nutrients and waste products is accomplished. The cell membrane is composed of lipids, proteins and carbohydrates. Lipids are represented by phospholipids, sphingolipids and sterols. Phospholipids and sphingolipids are amphipathic molecules: they have a hydrophilic (water-loving) group attached to a hydrophobic (water-fearing) chain. The hydrophilic heads of the lipid molecules face the outer environment on each side of the membrane, while the hydrophobic tails point to each other in the membrane interior escaping the water environment. This molecular orientation provides the formation of the lipid bilayer - the basic structure of the cell membrane (Figure 2.1). The lipid bilayer is fluid with viscosity similar to that of olive oil [30]. The most common sterol molecule in animal cell membranes is cholesterol. Cholesterol is randomly distributed in the bilayer and the fluidity of the membrane depends on its presence. Cholesterol helps the phospholipid molecules to stay together, not moving too far from each other or packing too tightly. This is important because if the phospholipids are separated at a great distance, unwanted toxic substances might easily enter the cell; if the phospholipids are too close, the passive transport of ions and small molecules through the membrane might be hindered.
The membrane hosts a large number of proteins. Early studies suggested that the protein molecules are located outside the membrane [32, 33]. In 1972 Singer and Nicholson published the fluid mosaic model [31] which proved that this is wrong. If we look at the cell membrane through a microscope we would notice that it looks like a mosaic built from different molecules. The components of this mosaic are not static, they are in constant motion mainly in the lateral directions. According to the fluid mosaic model, the lipid bilayer acts as a solvent for the embedded proteins which float in it like icebergs in the ocean (Figure 2.2). Experimental evidence has demonstrated that this qualitative picture is correct. The driving forces for this particular molecular arrangement are determined by the amphipathic properties of the molecules involved.

![Figure 2.1: A schematic representation of the cross-section of a membrane lipid bilayer. The black circles correspond to the hydrophilic heads of the lipid molecules. The curved lines correspond to the hydrophobic tails. From ref. [31]](image)

![Figure 2.2: Graphical representation of the fluid mosaic model. The solid bodies embedded in the lipid bilayer represent the integral membrane proteins. From ref. [31]](image)
CHAPTER 2. AFM-SMFS OF MEMBRANE PROTEINS: AN OVERVIEW

The protein molecules in the membrane can be divided into two groups: integral and peripheral proteins. The integral membrane proteins are embedded in the membrane and their extraction requires usage of detergents, non-polar solvents and denaturing agents. Transmembrane proteins play an active role in transport, in signal transduction, in cell-cell communication, among the most crucial biological processes in the cell. The peripheral proteins are located on the membrane surface, bounded through electrostatic or hydrogen bonds with the polar groups of the lipid molecules or the integral proteins. They can be easily extracted through changes in the external conditions, for example the pH of the environment.

Now that we briefly introduced the composition and the structure of the cell membrane we focus on the object of this study: integral membrane proteins. Later on when we use 'membrane proteins' in the text we refer to integral membrane proteins.

2.0.2 Integral Membrane Proteins.

Most of the protein structures we know, have been determined by X-ray crystallography. Membrane proteins turned out to be harder to crystallize compared to soluble proteins and in the very beginning only few structures were available. Recent advances in crystallography brought high-resolution X-ray structures for a larger number of membrane proteins. Protein structures can be determined also with NMR spectroscopy and electron microscopy or through a combination of the mentioned techniques. This has led to an increase in the number of solved three-dimensional structures of membrane proteins as depicted in Figure 2.3. Anyway, compared to soluble proteins, the number of available three-dimensional structures of membrane proteins remains small.

The folded state of a membrane protein does not depend only on the protein itself, like in the case of soluble proteins. The folded state strongly depends on the presence of the lipid bilayer. A simple comparison between the properties of the plasma membrane and the cytosol reveals more differences regarding membrane proteins and soluble proteins. Unlike cytosol, the plasma membrane is a heterogeneous and anisotropic environment with a very low dielectric constant. In most membranes gradients of pH, electric field, pressure, dielectric constant, and redox potential are present [30]. As a consequence, the lipid bilayer restricts the conformational space of membrane proteins.

Integral membrane proteins can be divided in two groups depending on their structural characteristics: α-helical and β-barrel proteins. α-helical membrane proteins are far more common. They are present in all types of biological membranes. It has been estimated that 27 % of all proteins in humans are α-helical membrane proteins. This can be explained by the underlying physics and chemistry of the interactions which stabilize α-helices in membranes (Figure 2.4).
Without knowing the structure of a membrane protein, by simply looking at its amino acid sequence, one can notice that it often contains regions of more than 20 consecutive hydrophobic residues. In practice, every integral membrane protein has one or more hydrophobic regions, corresponding to the transmembrane segments. Hydrophobic amino acids have side chains which orientate towards the lipid bilayer interior, forming favorable hydrophobic interactions. On the other hand, the protein backbone is hydrophilic because the peptide bonds connecting the amino acids are polar. Due to the absence of water in the protein interior, the backbone atoms tend to form hydrogen bonds with each other. The outstanding stability of the $\alpha$-helix is determined by the large number of hydrogen bonds formed along the protein backbone in this conformation. Since the lipid bilayer thickness is around 3 nm and the relative length of an $\alpha$-helix is 0.15 nm per amino acid residue, a sequence of 20 to 25 residues is sufficient to span the bilayer [34]. Depending on the number of transmembrane segments and their topology, we distinguish four types of integral membrane proteins as illustrated in Figure 2.5. If the protein has a single transmembrane segment, it is called biotopic protein; if the protein contains multiple transmembrane segments, it is called polytopic protein. If
the N-terminus of a biotopic protein is outside the membrane, the protein is classified as type I; if the C-terminus is outside the membrane the protein is classified as type II. When the multiple transmembrane segments of a polytopic protein are connected by loops the protein is assigned to type III. Biotopic proteins can oligomerize and form integral membrane proteins (type IV).

Figure 2.4: Schematic representation of a transmembrane alpha-helix. From ref. [30]

![Schematic representation of a transmembrane alpha-helix](image)

Figure 2.5: Topology-based membrane proteins classification. From ref. [30]

![Topology-based membrane proteins classification](image)

One of the most widely studied membrane proteins is bacteriorhodopsin (bR), a photosynthetic protein found in the plasma membrane of the archaeon *Halobacterium salinarium*. bR became the
paradigm for α-helical integral membrane proteins and ion transporters. bR is a small protein (~250 a.a.) with seven tightly-packed transmembrane α-helices connected by short loops. The N-terminus of the protein is outside the cell, while the C-terminus is in the cytoplasmic side. Together with specialized lipids, BR molecules form trimers arranged in a two-dimensional hexagonal lattice, known as the purple membrane (see Figure 2.11a). The purple membrane is a good example of a membrane with a complex composition. It contains 75 % proteins and 25 % lipids. The light-absorbing molecule retinal is bound to a lysine residue in bR’s helix G and has a deep purple color. When a photon gets absorbed by retinal, the molecule undergoes a conformational change, which leads to series of small conformational changes in bR. Due to these changes a proton gets transferred from the retinal outside the cell. Polar amino acids take specific positions along the protein and assist the proton transport. bR restores its initial conformation after the retinal recovers by accepting a proton from the cytosol. Then the cycle can be repeated again [35]. When this cycle is repeated multiple times by thousands of bR molecules, a large electrochemical proton gradient across the membrane is generated. In this sense, bR acts as a light-driven proton pump that moves protons out of the cell, generating a proton gradient. This proton gradient is used by the cell to synthesize adenosine triphosphate (ATP) molecules.

The second widely distributed conformation in membrane proteins is the β-barrel. β-barrel membrane proteins are present in the outer membranes of gram-negative bacteria, in the cell walls of gram-positive bacteria and in the outer membranes of mitochondria and chloroplasts in eukaryotes. The basic structural motif in β-barrels is the β-sheet. In comparison with an α-helix, the polypeptide chain in a β-sheet is more extended. If the sheet is properly oriented, only 7 residues are sufficient to span the bilayer. Another important difference between an α-helix and a β-sheet is that in the latter the side chains of alternating amino acids are found above and below the β-sheet. In practice this means that every second amino acid in the transmembrane segments of the barrel is hydrophobic, with its side chain placed in contact with the lipid bilayer. It is not mandatory for the rest of the residues to be hydrophilic. As a consequence, the transmembrane segments in β-barrels cannot be that easily detected from the amino acid sequence. The number of β-strands in a β-barrel may vary between 8 and 22. In many proteins they are tilted from the perpendicular to the bilayer plane by ~45° but in some cases the angle can be smaller. This orientation of the β-sheets leads to the formation of a cylinder known as the β-barrel (Figure 2.6a). The same factors that govern the formation of helical bundles, govern the formation of β-barrels. The conformation with maximum intrachain hydrogen bonds between the backbone atoms is the most favorable. The intrachain hydrogen bonds in β-barrels make them rigid and very stable. β-barrel integral proteins often form homotrimers but they can also be monomers.

The paradigm for the structure of β-barrel integral proteins are porins (Figure 2.6). Porins are transmembrane proteins which form wide pores in the outer membranes of Gram-negative bacteria, mitochondria and chloroplasts. The pores provide pathway for the passive diffusion of small polar
molecules across the outer membrane. Each $\beta$-barrel is composed from 16 to 18 $\beta$-strands with antiparallel orientation between each other (Figure 2.6a). The strands are connected by short loops on the periplasmic side, and long irregular loops on the extracellular side. There is an internal loop that faces the barrel interior and keeps the structure more compact. These architectures are remarkably stable and denaturation is the only way to disassemble them [36].

Figure 2.6: A schematic representation of the transmembrane $\beta$-barrel protein OmpF porin from *E. coli*. In panel a: side view; in panel b: top view. From ref. [36]

Most of the functions of the cell membrane are performed by certain types of membrane proteins. Based on their functions they can be divided into transporters, anchors, receptors and enzymes (Figure 2.7). Transporters carry nutrients, metabolites and ions across the membrane. The solutes can be distinguished by their size and charge (in ion channels) or according to their ability of fitting into the binding site of the protein. Anchors bind macromolecules on both sides of the membrane. For example, integrins bind fibronectin outside the membrane and they are linked to the cytoskeleton inside the membrane. Integrins facilitate the cell-extracellular matrix (ECM) adhesion. Receptors generate intracellular signals through binding specific extracellular molecules. In this manner communication between the outside environment and the cell is accomplished and in response a particular action is performed by the cell. Platelet-derived growth factor (PDGF) receptor binds PDGF and causes the cell to grow and divide. Membrane proteins which act as enzymes catalyze different reactions. For example, adenylyl cyclase catalyzes the production of cyclic AMP inside the cell.
The role of membrane proteins in living organisms is so essential, that understanding their structure and function has triggered an enormous amount of research. Since membrane proteins pose a serious challenge to conventional experimental techniques for studying proteins, new experimental approaches, like atomic-force microscopy (AFM), offer a promising alternative.

### 2.0.3 Probing the Structure of Membrane Proteins by Atomic Force Microscopy (AFM).

Atomic force microscopy is a novel technique that examines the surface structure and topography of different samples, with application in wide range of disciplines, from solid state physics to molecular biology and medicine. The method is also successfully applied to biological membranes. The AFM is a scanning probe microscope (SPM). What SPMs have in common is the probe (tip) that scans the sample surface in a different manner in the different methods. In AFM, the tip is brought in physical contact with the sample surface and raster scans it, "sensing" the surface through the tip-sample interaction forces. As a result, a three-dimensional topographical image of the sample with submolecular resolution is obtained.

Scanning probe microscopy was developed in the early 80s by Binnig and Rohrer [37] and only four years after its discovery they were awarded with the Nobel Prize in Physics. The main difference between SPM and optical and scanning electron microscopy is that the image obtained with SPM is three-dimensional; indeed, the height is revealed in SPM images. The scanning tunneling microscope (STM) is a SPM but it has the disadvantage that it can be used only to study conductive surfaces. The AFM was developed to overcome this disadvantage and can be applied to all kinds of surfaces. Furthermore, the AFM can operate in physiological solutions at temperature 37° which makes it extremely useful for studying biomolecules in their native environment. The main component of the AFM is a sharp tip mounted on the end of a flexible cantilever (Figure 2.8a). As the tip approaches the surface attractive forces between the two surfaces (the tip surface and the sample surface) cause the spring-like cantilever to bend. Laser light pointed to the cantilever bounces off its
back onto a position-sensitive photo detector. As the cantilever bends, the position of the reflected laser beam on the photo diode shifts and the deflection of the light is measured with high precision. Since protrusions and indentations in the sample surface lead to changes in the cantilever deflection, measuring and processing this deflection leads to the topographical image of the examined surface. Furthermore, Hooke’s law can be applied to the cantilever and knowing the cantilever’s spring constant, the deflections can be transformed into forces. In this manner, the interaction forces acting between the AFM tip and the sample are measured.

Figure 2.8: Schematic representation of an atomic-force microscope (AFM). a. The AFM cantilever. b. The basic components in a standard AFM setup. From ref. [38, 39]

The basic components of an AFM are: the tip and the cantilever, the laser, the photodiode detector, the piezoelectric scanner and the feedback electronics (Figure 2.8b). The tip is the most important component since it is the part of the microscope that physically touches the sample. The AFM tip comes in different sizes and shapes, but the typical tip radius is between 5 and 20 nm. The resolution of the image depends strongly on the sharpness of the tip. The sharper the tip, the higher the resolution. But we should keep in mind that due to the physical interactions with the sample, the tip changes during the experiment and the operator should take account for these changes. Working with the sharpest tip does not guarantee the optimal experiment outcome. The AFM cantilever is 100-200 $\mu m$ long and what really matters is its spring constant. The smaller the spring constant, the smaller the forces "sensed" by the cantilever. The photo detector used in commercial AFMs is usually a quadrant photo diode (QPD). Once a photo diode gets hit by light, a voltage is generated. In a QPD the voltage magnitude is position-dependent. For example, if the
light beam falls in the centre of the diode, the voltage is zero; in any other place, a finite voltage is generated. In many instruments, a piezoelectric scanner governs the motion of the sample stage. Piezo is a material that changes its dimensions due to applied voltage. For example, if a positive voltage is applied, the material elongates; if a negative voltage is applied, the material becomes shorter and thicker. These properties of the piezo allow control on the distance between the AFM tip and the sample. In other AFMs, the piezoelectric scanner is implemented in the cantilever holder and controls the cantilever motion. In the most frequently used AFM operational modes constant contact force between the tip and the sample during scanning is maintained through a feedback electronics loop. The feedback electronics maintains constant contact force by adjusting the tip-sample distance with the piezoelectric scanner. In practice, the cantilever deflection is monitored and kept at a user-predefined value.

2.0.4 AFM based Single Molecule Force Spectroscopy (SMFS).

Single molecule force spectroscopy techniques enable the manipulation of single molecules one at a time as opposed to bulk experimental approaches. Studying the properties of single molecules becomes extremely important at the cellular level where the concentrations of biopolymers, such as DNA and proteins, can be on the nanomolar scale. Single molecule force spectroscopy (SMFS) techniques allow to study the mechanical properties of single biomolecules through measurement of the inter- and intramolecular forces acting in these molecules. The AFM is one of the most commonly used techniques for SMFS. The physical contact between the tip and the sample allows direct manipulation. The forces acting between the tip and the sample can be measured straightforward and in addition, external forces can be exerted on the sample. These operations can be performed under native conditions, which is without any doubt the major advantage of the method.

AFM-based SMFS has been used to study protein unfolding [8, 9], antigen-antibody binding [40], protein-ligand interactions [41], polysaccharides elasticity [42] etc. Here we will focus on the application of AFM-SMFS in protein unfolding, membrane proteins unfolding in particular. In these experiments, referred to as pulling experiments, the protein molecule is on one side bound to the surface in solution. For example, if the surface is made of gold, the sulfhydryl group of a cysteine residue added to one of the protein ends, can bind covalently to the surface. On the other side, the molecule gets picked by the AFM tip and stretched as the tip retracts from the surface. The molecular force response versus the distance between the tip and the sample surface is recorded and presented in the form of a force-extension (F-x) curve (Figure 2.9). The red line in Figure 2.9b represents the approach of the tip towards the surface, while the black line represents the retraction cycle.
Figure 2.9: Schematic representation of the process of obtaining a force-extension (F-x) curve. (a) Step 0: The AFM tip is brought in contact with the sample and the retraction cycle can begin. Step I: A polyprotein got attached to the tip. As the tip moves away from the surface, the distance between the two increases and the protein elongates. Step II: Further extension of the polyprotein leads to the unfolding of the domain in the middle. The protein contour length increases, the force acting on the cantilever suddenly decreases due to the released tension and we observe a force drop in the F-x curve. Step III: The unfolded protein domain is stretched until it is fully elongated and the contour length of the polypeptide chain grows accordingly. Step IV: steps I, II and III repeat until the remaining protein domains are unfolded and step V is reached. Step V: The entire polypeptide chain is unfolded and fully-elongated. This corresponds to the detachment peak in the F-x curve. Step VI: the protein gets detached from the AFM tip. (b) The F-x curve corresponding to the unfolding events presented in panel a. The red line represents the approach of the cantilever to the sample surface (0). The initial forces are negative because once the cantilever is in contact with the sample surface it gets deflected upwards. The black line represents the retraction cycle of the cantilever at constant velocity. As the distance between the tip and the surface grows, the protein exerts a resisting force and the cantilever bends correspondingly until a certain force is reached (I). At this force the protein domain gets unfolded and it no longer exerts resistance during elongation. As a result the force drops (II). This scenario repeats for the remaining domains (III, IV) until all of them are unfolded and the polypeptide chain is fully elongated (V). After this, the protein detaches from the tip (VI). From ref. [43]
What happens in pulling experiments is the following. When the AFM tip is brought in contact with the sample surface a single protein can attach to the tip. The physics behind this interaction is still unclear. A possible explanation includes non-specific physioadsorption and electrostatic interactions [43]. The results show that the adsorption is increased when a large force (∼ 800-3,000 pN) is applied to the surface once the tip is in contact [43]. The AFM tip is then retracted from the surface. During retraction, the protein chain, tethered between the tip and the surface, elongates. The polypeptide chain resists to this elongation. The resistance forces are entropy-driven. The fully-stretched conformation is unfavorable. The preferred conformation of an unfolded protein is a random coil, which maximizes the entropy. As the tip moves away from the surface, it experiences the resistance force of the protein and the cantilever bends. The cantilever deflection is proportional to the force acting on the cantilever. Once the protein is unfolded and fully-stretched the tension is released and the force drops. In the case of a multi-domain protein, like the one in Figure 2.9a, the scenario repeats until all protein domains are unfolded. The resulting force-extension curve (Figure 2.9b) typically bears a sawtooth pattern. Each force peak in the curve corresponds to the unfolding of a single protein domain. The last peak is known as the detachment peak and is assumed to correspond to a point in which the polypeptide chain is fully stretched and detaches from the tip.

Polymer elasticity models can be used to describe the mechanical behavior of proteins during unfolding. Some of these models turned out to be very useful in the analysis of force-extension curves. The standard model used for the analysis of F-x curves is the worm-like chain (WLC) model. Bustamante et al. were the first to derive an interpolation formula for F-x curve and to apply it to their DNA unfolding experiments [44]. The resulting formula, now known as the WLC equation, is:

\[ F(x) = \frac{k_B T}{l_p} \left( \frac{1}{4} \left( 1 - \frac{x}{L_c} \right)^{-2} + \frac{x}{L_c} - \frac{1}{4} \right) \]  

where \( F \) is force, \( x \) is extension, \( k_B \) is Boltzmann’s constant, \( T \) is temperature, \( l_p \) is persistence length and \( L_c \) is contour length. Eq. 2.1 enables computation of the entropic restoring force \( F \) exerted by the polymer at different extension values. This computation requires two parameters: the persistence length, \( l_p \), and the contour length, \( L_c \). The persistence length is a measure of the stiffness of the polymer; at length above \( l_p \) the polymer behaves like an elastic rod. The contour length, \( L_c \), is defined as the length of a polymer chain at maximum physically possible extension [45]. Fitting the experimental data with the WLC model at fixed persistence length, gives the portion of unfolded protein in terms of \( L_c \). When the fit is performed for each peak in the spectra, we obtain information about the length of the unfolded segments. Failure of the model at large forces due to overstretching of the bonds has been reported [46]. In a first approximation, the persistence length is assumed to be the same for the folded and unfolded states and to be independent of the extension. In the literature, different \( l_p \) values for proteins have been reported [47, 8, 9], 0.4 nm is accepted to be the standard for membrane proteins [9].
CHAPTER 2. AFM-SMFS OF MEMBRANE PROTEINS: AN OVERVIEW

To illustrate better the pulling experiments with AFM, let’s look more closely at some real experiments. AFM-based SMFS was used to study the mechanical properties of the globular protein titin [8]. Titin is a multidomain protein composed by 244 domains (immunoglobulin (Ig) and fibronectin domains) connected by unstructured loops. The length of each domain is around 89-100 amino acids. The length of this protein is greater than 1 µm [48]. Titin is the largest known protein [49]. The mechanical properties of this giant molecule are responsible for the passive elasticity of muscles. By measuring these properties we can understand better the structure-function relation in titin. In a work by Gaub et al. [8] AFM pulling experiments on titin were performed. Native titin solution in phosphate-buffer saline (PBS) with concentration 10-100 µg/ml was applied on gold surface and left to adsorb for 10 min. It was then rinsed with PBS and sampled in the fluid cell of the microscope. The AFM tip was brought in contact with the gold surface and F-x curves were recorded through retraction of the tip from the surface. The resulting spectra are shown in Figure 2.10. The part in the spectra that is hard to interpret is the initial part. It contains high force peaks suggesting multiple molecular interactions between the tip and the gold surface. What is characteristic about all curves is the presence of a sawtooth pattern of equally spaced peaks with spacing around 25 nm and maximum unfolding forces in the range 150-300 pN. The length of a stretched Ig domain is expected to be 31 nm. This lead to the working hypothesis: the repetitive peaks in each spectrum represent the successive unfolding of the individual domains in the titin molecule. To test this hypothesis, the authors constructed two recombinant titin fragments. The first one consisted of four Ig domains, Ig4, and the second one consisted of eight Ig domains, Ig8. The fragments were anchored to the gold surface by two cysteines added to the C-terminal. The pulling experiments were performed and the resulting curves supported the working hypothesis. All F-x curves displayed the sawtooth pattern with varying number of equally spaced peaks at distance 25 nm with forces between 150 and 300 pN. In the case of the Ig4 the maximal number of peaks observed was 4, while in the case of the Ig8 - 8. The variability of the total number of peaks is a consequence of the randomness with which the AFM tip attaches to a certain position in the polypeptide chain. In general, the AFM tip can be functionalized in such a way that the attachment site along the polypeptide chain is known. For example, one can pick the protein with a gold-coated tip through a cysteine residue added in the C-terminal of the protein. If this step is skipped, the maximum extension and the shape of the resulting force-extension curves vary as a consequence of the different attachment sites between the tip and the polypeptide chain. In Figure 2.10b, a force-extension curve representing the unfolding of Ig8 is shown. The WLC fit with persistence length 0.4 nm is depicted. The increase in the contour length is 28-29 nm per domain, which is closer to the expected length for a fully-stretched domain, 31 nm. The force peaks magnitude increases with the extension. This is a sign that the weakest domains unfold first and the unfolding events are ordered by force not by the domains positions along the chain. The authors performed also unfold-refold experiments in which they showed that once unfolded, after relaxation, the protein gets refolded again. The protein’s
ability to refold is assumed to be important for its biological role. The example of the unfolding of single titin molecules clearly shows the power of AFM-SMFS in the investigation of the mechanics of globular proteins.

Figure 2.10: a. Force-extension curves obtained by stretching the soluble multidomain protein titin. Three representative curves are shown. They all contain the characteristic sawtooth pattern with periodicity in the force peak positions. The spacing between two successive peaks is between 25 and 28 nm and is consistent with the sequential unfolding of individual titin domains. The randomness in the protein-tip attachment positions can be seen in the patterns appearance at different initial extension values. b. Force-extension curve obtained from the unfolding of a single titin fragment containing eight immunoglobulin domains (Ig8). The number of force peaks observed in the spectrum is 7. Each peak is fitted with the WLC model with a persistence length 0.4 nm. The WLC model predictions are consistent with the explanation that a single peak corresponds to the unfolding of a single immunoglobulin domain. According to the model, the contour length of the polypeptide chain increases by 28 to 29 nm each time an individual domain unfolds. This is very close to the contour length predicted from the amino acid sequence of a single immunoglobulin domain, which is 30 nm. From ref. [8]

Unlike other experimental techniques, AFM-SMFS turned out to be a promising tool for examination of the structure of membrane proteins. As we commented in subsection 2.0.2, there are differences between globular and membrane proteins, mainly coming from the hydrophobic nature of the latter and the presence of the lipid bilayer with its specific properties. The lipid bilayer creates obstacles to the traditional methods used to determine protein structure. On the other side, neglecting the membrane and its effect on protein structure and function is not possible. AFM allows imaging and manipulation of membrane proteins in their natural environment, embedded inside the lipid bilayer. The number of SMFS experiments on membrane proteins reported in the literature is growing fast. For example, bacteriorhodopsin and rhodopsin [9, 10, 11, 12], CNGA1 channel [13, 12], Na+/K+ antiporter [14], leucine-binding protein [15] etc have been already studied
Figure 2.11: a. High-resolution AFM topographical image of the cytoplasmic surface of a wild-type purple membrane of *H. salinarium*. The hexagonal lattice built from bacteriorhodopsin (bR) trimers is captured. b. AFM topographical image of the same membrane patch after the AFM tip was positioned above a single protein (marked with a white circle in both panels a and b) and brought in contact with it. Contact force of 1nN was applied on the sample for 1 second to facilitate the protein adsorption to the tip. The image reveals the gap appearing in the position on which the AFM tip was pushed and retracted (white circle). Thus these images show the extraction of a single bR in AFM-SMFS experiments. c. Superimposition of 13 F-x curves representing the unfolding of bR from the purple membrane. The three major force peaks are fitted with the worm-like chain (WLC) model with persistence length 0.8 nm. The contour length values predicted by the model are written above the fitted lines. d. Schematic representation of the bR structure, relating the contour length values given in panel c with the protein transmembrane helices and the loops connecting them. From ref. [9] with AFM-SMFS.

Bacteriorhodopsin (bR) was the first membrane protein unfolded with SMFS. The unfolding pathways of bR in native purple membranes were first described by Muller et al. [9]. In this study, AFM was used both for imaging the purple membrane and pulling and unfolding a single bR out of it. The native purple membrane of *Halobacterium salinarium* was adsorbed on a mica surface and scanned with the AFM. The obtained topographical images revealed the characteristic purple membrane structure: a hexagonal lattice composed of bR trimers (Figure 2.11a). The topography of the purple membrane has been already examined with electron microscopy and x-ray crystallography but not in aqueous solution. After the pulling experiments were performed, the sample surface was imaged again showing a vacancy in the position where the extracted protein was located (Figure 2.11b). The randomness in the tip attachment site along the protein chain resulted in force-extension
curves of different lengths and shapes. To overcome this the authors selected only the spectra in which the position of the detachment peak is between 60 and 80 nm, which corresponds to the length of a fully-extended bR molecule. Figure 2.11c shows the superimposition of the force spectra obtained showing also their fit with the WLC model with persistence length 0.8 nm. The selected curves have four major peaks with contour length values matching perfectly a scenario in which each peak corresponds to the pairwise extraction and unfolding of the bR helices. bR has 7 transmembrane α− helices, labeled as A, B, C, D, E, F, G and connected with each other through non-structured loops (Figure 2.11d). The WLC fit to the second force peak revealed a contour length of 88 a.a., which matches the number of a.a. in the C-terminal, helices G and F, the F-G loop and the E-F loop. Following the same logic, the third peak has been assigned to the unfolding of helices E and D and the last peak - to the unfolding of helices C and B followed by the extraction of helix A. The first peak is assumed to reflect the extraction of helices G and F. Another interesting observation is that the peaks tend to order in a descending manner with respect to the force magnitude. We saw that in the titin example, it was the opposite, indicating that the weakest domains unfold first. In the case of bR the order of the unfolding events depends on the position of the transmembrane domains not on the strength of the interactions that stabilize them. To validate the results, same experiments were repeated with a gold-coated tip and a cysteine residue added in the bR C-terminus. The results were consistent with the previous experiments. The authors performed also another set of experiments in which they cleaved the E-F loop in the protein. The obtained curves were shorter than 45 nm with three major peaks. The peak that was corresponding to the unfolding of helices G and F was missing as it can be expected. The spectra were analyzed and interpreted following the same logic as before. Since helices G and F remained in the membrane, the authors speculate that their presence in the lipid bilayer additionally stabilized helices C and B, which lead to the gradual unfolding of helix C.

The F-x curves obtained with AFM-SMFS experiments performed on membrane proteins provide new information about the structure of these proteins, about the inter- and intramolecular interactions which stabilize them. Furthermore, they provide new information, unavailable before, about the protein-membrane interactions and the role of the membrane in the proteins functional cycles. The interpretation of the experimental data can be assisted by computational methods offering deeper understanding of the data at the molecular level. This is the topic of the next subsection.

2.0.5 Molecular modeling.

Theoretical and computational approaches provide a complementary information related to proteins structure, dynamics and function and are particularly useful for membrane proteins, where experimental information is still relatively scarce.

Molecular dynamics (MD) is widely used in studying all kinds of biomolecules. The standard MD setup uses all-atom molecular models, in which a molecule (the protein, DNA, water, the
phospholipids of the membrane, etc.) is represented explicitly by all of its atoms. This approach is accurate and realistic from both the physical and the chemical point of view but it is computationally expensive. Atomistic MD simulations have been successfully used to investigate the conformational changes of membrane proteins in lipid environments [50, 51]. Even though advances have been made, still there are limitations regarding the time scales accessible to these simulations. It is known that the transition time between the functional states of membrane proteins is much longer than the time accessible with conventional MD simulations [52]. The time it takes to perform AFM pulling experiments is of the order of magnitude of 0.1 s, $10^5$ times more that the time that can be simulated on systems of this complexity with ordinary resources. Indeed, a MD simulation of an AFM experiment requires simulating a system containing thousands of solvent molecules, hundreds of lipid molecules and the membrane protein itself. Given that the number of calculations per molecule scales linearly with the number of particles in the molecule, the larger the system, the bigger the simulations length.

Despite the limitations of conventional MD, Kappel et al. [53] investigated the mechanical unfolding of bacteriorhodopsin (bR) using all-atom MD simulations. The membrane was included explicitly in the model, represented with a POPC (1-palmitoyl-2-oleoyl-sn-glycero-3-phosphocholine) lipid bilayer. Four bR trimers were embedded in the bilayer reproducing the characteristic purple membrane structure, in which bR trimers are organized into two-dimensional hexagonal lattice (see subsection 2.0.2). Therefore, the simulation box contained 12 bR monomers. When a protein is pulled, it elongates accordingly. This demands the simulation box to provide enough space to host the extracted polypeptide chain. A water layer with thickness 10 nm was added to the simulation box in order to provide such space. With this, the total number of atoms in the simulation box became 236,124. The mechanical stretching of the protein was achieved by applying a harmonic potential to the $C_\alpha$ atom at the terminus, which was pulled in the $z$-direction at a constant velocity moving away from the membrane. Anyway, the size of the simulation box still did not provide enough space to comprise the fully elongated polypeptide chain of the bR monomer. The $z$-dimension of the box was 15.32 nm, while the approximate length of a single bR monomer is $\sim 92.4$ nm. A novel computational protocol was introduced to deal with this issue. The unfolded parts of the protein at a certain distance from the upper wall of the simulation box and from the lipid bilayer border, were repeatedly removed. The holes left by the missing residues were filled with water, the energy of the system was minimized and the system was equilibrated. A new terminal $C_\alpha$ atom was defined and subjected to the pulling potential. These steps were iterated until complete protein unfolding occurred. The authors performed MD simulations using different pulling velocities. The smallest was 1 m/s, which is $\sim 10^7$ times larger than the typical values used in real experiments. In this field, the gap between simulation conditions and experimental conditions is so large that the improvement of computer hardware is not likely to fill it in the near future.

To overcome the main limitations of all-atom MD simulations two main approaches can be
used: one is to simplify the molecular model of the protein-membrane system, and the second is to use enhanced sampling techniques. A combination of the two can also be attempted. Here we are going to describe only the first approach. We will describe the different coarse-grained (CG) techniques specifically developed for membrane proteins and we are going to see how the method of MD combined with CG models has been used to reproduce the characteristic F-x curves obtained with AFM-SMFS.

**Implicit solvation models.**

In these schemes, the protein is modeled explicitly in an atomistic manner, while the solvent and the membrane are included implicitly with a mean-field model, for example the generalized Born (GB) model [54]. Water is modeled as a continuous environment with dielectric constant $\epsilon = 80$. A solvation energy term is added to the molecular mechanics potential energy function to account for the solvent-solute interactions. The conventional lipid bilayer is replaced by a low-dielectric slab of a certain thickness, placed in the high-dielectric environment induced by the water molecules and the lipids polar head-groups. The dielectric constant in the protein interior is typically set to 1.

In Figure 2.12 we illustrate graphically two implicit membrane models currently in use. In the GBSW model [55] (Figure 2.12a), the membrane area occupied by the hydrophobic lipid tails is modeled as a slab having the same dielectric constant of the protein, $\epsilon = 1$. A smoothing function is included in the model acting on the two dielectric borders: between the hydrophobic tails and the water, and between the water and the protein. The GBSW model, in combination with advanced computational sampling methods, was applied to three membrane proteins: melittin from bee venom, the transmembrane domain of the M2 protein from Influenza A (M2-TMP), and the transmembrane domain of glycophorin A (GpA), investigating the membrane effects on conformational changes, the helix-to-helix interactions in membranes, etc [55]. The model was successfully used to fold and assemble helical membrane proteins [55].

In the HDGB model [56] (Figure 2.12b), the membrane slab has two layers described with two different dielectric constants. The first layer is associated with the membrane hydrophobic core and has a dielectric constant $\epsilon = 2$ which is slightly different with respect to the protein. The second layer is associated with the polar lipid head-groups in the membrane and it has dielectric constant $\epsilon = 7$. The bacteriorhodopsin monomer and trimer were simulated with the HDGB model [57] and the obtained trajectories were in excellent agreement with explicit membrane simulations.
Implicit solvent and membrane models have been successfully applied for simulating AFM-SMFS experiments. Seeber et al. [58] modeled the forced unfolding of bacterioopsin, which is the retinal-free form of bacteriorhodopsin, using atomistic description of the protein and implicit model for the membrane and the solvent. The results from these simulations revealed details at the atomistic level about the sequential unfolding of the individual protein helices, suggesting that the characteristic F-x curves and the sequence of unfolding events are altered by the up-and-down topology of the seven-helix bundle. Yamada et al. [59] also used implicit membrane and solvent models but combined with coarse-grained model of the bR protein. In this study, the key features of the experimental F-x curves were successfully reproduced, including the peak positions, suggesting that the peak positions are determined exclusively by the residue-lipid and the intrahelix interactions.

**Full coarse-graining: the MARTINI force field.**

In this approach the protein, the membrane and the solvent are included explicitly but not with all their atoms. Groups of atoms get replaced by a single bead with certain properties, so that the molecules get literally "coarse-grained". The MARTINI model developed by Marrink et al. [60] is the most popular model of this kind.

In the MARTINI model [60], on average four heavy atoms together with the corresponding hydrogen atoms form a single coarse-grained (CG) bead. This is called four-to-one mapping. In the water solvent case, four water molecules are modeled with a CG water bead. The number
four was chosen to reach a compromise between the computational efficiency and the realism of the chemical description. For certain types of molecular fragments, like aromatic rings, a single CG particle contains only two heavy atoms for a more proper characterization. Practically, the initial all-atom molecular structure is mapped onto CG particles connected to each other in such a way that the overall topology of the molecules is preserved. The MARTINI particles are divided into types and subtypes based on polarity and hydrogen-bonding capacity. In this manner, 18 particle types are obtained, which are called ‘building blocks’. The main assumption in MARTINI is that the parametrization of the building blocks is transferable to different molecules and the model does not need to be reparametrized in each case. The parametrization is performed using an extensive calibration of chemical building blocks towards thermodynamic data, mainly oil/water partition coefficients. The MARTINI force field currently contains parameters for lipids, sterols, proteins, sugars etc.

The MARTINI CG model has been successfully applied to MD studies of the self-aggregation of rhodopsin monomers in different explicit CG lipid membrane environments [61]. In a study by Xu et al. [62] the effect of different cholesterol content in the membrane on the aggregation of the toxic peptide amylin was investigated. They performed simulations in a mixed dipalmitoylphosphatidylcholine (DPPC) and dipalmitoylphosphatidylserine (DPPS) bilayers with different cholesterol concentrations and without cholesterol. It was shown that in the absence of cholesterol, the amylin aggregates are located inside the bilayer, while in the presence of cholesterol, the amylin aggregates are positioned outside the bilayer, on the bilayer-water interface. These results are consistent with cholesterol hindering formation of the toxic amylin oligomers inside the cell membrane. To the best of our knowledge the MARTINI force field has never been used to simulate AFM-SMFS experiments.

Go-like models.

An even simpler approach for simulating AFM-SMFS experiments was developed by Cieplak et al [25]. Since in our work we use the Cieplak model as a starting point for the investigation of the mechanical unfolding of rhodopsin (see Chapter 3), we are going to describe it in more details.

The model introduced by Cieplak et al. [24] is a Go-type model in which the native protein conformation is determined by the experimental structure at room temperature. The protein is represented as a chain of beads centered at the \( C_\alpha \) atoms in the protein backbone. A harmonic backbone potential acts between consecutive beads, tethering them at the peptide bond length, \( r_0 = 3.8 \, \text{Å} \):

\[
V_{BB} = \sum_{i=1}^{N-1} \frac{1}{2} k_{BB} (r_{i,i+1} - r_0)^2
\]  

(2.2)

where \( r_{i,i+1} = r_i - r_{i+1} \) is the distance between two adjacent beads and \( k_{BB} = 100 \epsilon / \text{Å}^2 \) is the spring constant.

In order to preserve the native chirality, a potential \( V_{CHIR} \) was introduced in the model. The
The chirality potential has the form:

\[ V^{CHIR} = \sum_{i=2}^{N-2} k_{CHIR} C_i^2 \Theta(-C_i C_i^{NAT}) \]  

(2.3)

where the chirality of the residue \( i \) is given by

\[ C_i = \left( \frac{(v_{i-1} \times v_i) \cdot v_{i+1}}{r_0^3} \right) \]  

(2.4)

with \( v_i = r_{i+1} - r_i \) and \( r_0 \) the peptide bond length. \( C_i \) can take values between -1 and 1. The positive values of \( C_i \) correspond to a right-handed helix, the negative values - to a left-handed helix. \( C_i^{NAT} \) is the chirality of residue \( i \) in the native conformation. \( \Theta \) is a step function that returns 1 for positive arguments and 0 otherwise. In practice, if the chirality sign for a given residue matches the native chirality sign for the same residue, \( V^{CHIR} \) is 0. If there is a mismatch in the two signs, that conformation is punished with the square of \( C_i \). \( k_{CHIR} \) is set to be equal to \( \epsilon \).

The non-bonded interactions are divided into native and non-native interactions. This is done following the procedure developed by Tsai et al. [63]. The distances between all heavy atoms in the native conformation are computed and if a distance is smaller than the sum of the vdw radii of the two atoms multiplied by a factor of 1.244, the contact between these two atoms is considered native; otherwise the contact between the two atoms is considered non-native.

The interaction between residues that form a native contact is described by the Lennard-Jones potential:

\[ V^{NAT} = \sum_{i<j}^{NAT} 4\epsilon \left[ \left( \frac{\sigma_{ij}}{r_{ij}} \right)^{12} - \left( \frac{\sigma_{ij}}{r_{ij}} \right)^6 \right] \]  

(2.5)

Here, \( r_{ij} \) is the distance between the \( C^\alpha \) atoms of residues \( i \) and \( j \). The \( \sigma_{ij} \) values are computed by multiplying the experimental distances by \( 2^{-\frac{1}{6}} \), which adjusts the potential minima to the native state. The parameter, \( \epsilon \), when divided by \( k_B \), has value 900 K, which proved to be appropriate in previous simulations of folding and unfolding, yielding the correct force peak magnitudes for titin [25] and ubiquitin [27] at room temperature.

The non-native interactions are assumed to be purely repulsive for distances shorter than the equilibrium distance corresponding to \( \sigma_0 = 5\text{Å}, r_{cut} = 5.61\text{Å} \):

\[ V^{NON} = \sum_{i<j}^{NON} V_{ij} \]  

(2.6)

\[ V_{ij}^{NON} = \begin{cases} 4\epsilon \left[ \left( \frac{\sigma_0}{r_{ij}} \right)^{12} - \left( \frac{\sigma_0}{r_{ij}} \right)^6 \right] + \epsilon, & \text{if } r_{ij} < r_{cut} \\ 0, & \text{otherwise} \end{cases} \]  

(2.7)
The model described so far was successfully applied to soluble proteins like titin [25], calmodulin [26] and ubiquitin [27]. The simulated F-x curves preserved the characteristic features of the experimental curves. Moreover, the simplicity and the computational efficiency of this model enabled the evaluation of the effects of varying temperature and pulling velocity.

Afterwards, Cieplak et al. extended the model to simulate the unfolding of membrane proteins, in particular bacteriorhodopsin (bR) [28], where the membrane effects are essential for the final outcome from the simulations. They approached the problem in the following way: the membrane was included in the model explicitly with a coarse-grained representation similar to the one used for the protein. In order to arrive to an adequate initial configuration of the protein-membrane system, all-atom MD simulation of bR embedded in a POPC lipid bilayer with water solvent was performed with computational protocol described in ref. [28]. The resulting conformation was used as a starting point for the stretching simulations. In the coarse-grained model of the membrane, the lipids were represented by merely all of their C atoms. The contacts of these carbon atoms with the protein were determined using the same procedure by Tsai et al. [63] used to derive the native contacts in the protein. The protein-membrane interactions were modeled with the Lennard-Jones potential applying the same strength used for the native protein interactions, namely $\epsilon$. The main limitation of this approach is that during the stretching simulations, the membrane was held frozen. Therefore, when an unfolded part of the protein is extracted from the membrane it leaves a hole in it. Clearly, such description is not very realistic, since in a real membrane the lipid molecules fill the voids left by the extracted protein. In Chapter 3 we describe a model capable of overcoming this limitation.

2.0.6 Analysis tools for AFM-SMFS experiments.

SMFS experiments need to be performed multiple times on the same molecule in order to obtain statistically relevant information. Recent improvements allow the collection of large amount of data [16] but careful preprocessing is required because not every F-x curve contains meaningful unfolding events. Most of the traces result from non-specific interactions, or contain partial unfolding events, or simply noise. Typically membrane proteins are completely unfolded in $< 1\%$ of the cases [21]. Furthermore, there is a certain variability in the peak patterns of F-x curves of the same protein reflecting different unfolding pathways. What experimentalists often do is manual notation by visual inspection of the traces. If the data set contains thousands of traces, handling the problem manually is far from optimal, not only because it is time consuming but also because the result depends on the user’s expertise. Recording thousands of force-extension curves in a reasonable time is now possible. This huge amount of data calls for an automatic procedure for classification and statistical data analysis. Many attempts of developing such a procedure have been made. Next, we will describe some of these methods in a chronological order.
The analysis approach by Kuhn et al. (2005)

One of the earliest attempts for the development of an automatic SMFS data analysis tool was published in 2005 by Kuhn et al. [19]. The software they developed allows classification and statistical analysis of large amount of force spectra. The procedure performs fit with the WLC model, automatic alignment of F-x curves and peak patterns classification for detection of different unfolding pathways. The data they analyzed comes from unfolding of two membrane proteins: bacteriorhodopsin (bR) from *Halobacterium salinarium* [9, 64] and the sodium-proton antiporter NhaA from *Escherichia coli* [65]. The algorithm is based on two steps: first, individual processing and analysis of each trace; and second, pairwise alignment of all traces, followed by hierarchical clustering and peak classification.

As a first step, each F-x curve undergoes determination of a zero-force baseline and contact point detection. The zero-force baseline was defined using a linear fit to the non-contact part. The contact point corresponds to the point at which the undeflected cantilever is in contact with the sample surface [19]. Next, the detachment peak is identified in the following way: the standard deviation from a linear fit to the last 5 nm of the trace was estimated and the first point, moving in the direction of the trace origin, in which the standard deviation increased by a factor of 1.5 was the detachment peak. Next, the peaks in each F-x curve were detected. In order to ease the peak detection step, each F-x curve was smoothed. The force peaks were detected based on the differences between local minima and local maxima. If a peak with height above 30 pN is surrounded by force minima with force difference of at least twice the standard deviation of the baseline to the maximum, it is recognized as a force peak by the program. All peaks were fitted with the WLC model except the ones in the first 15 nm which are assumed to come from non-specific interactions.

In general, the analysis of F-x curves requires an alignment step. The reason is that different attachment sites between the AFM tip and the protein might introduce horizontal shifts in the curves [21]. The authors tested three different approaches for alignment. First, they tried to align the force spectra using the contour length values of each peak. This attempt was unsuccessful because the variability in the tip-sample attachment sites leads to high variability in the absolute and relative contour length values. The second trial was based on alignment of pixel images of the F-x curves represented as two-dimensional histograms. In this procedure, two pixel images are shifted with respect to each other and an alignment score is computed. The results were encouraging but the alignment quality was affected by variations of the data points density, for example in the base of the peaks. In the third approach, each F-x curve was represented as a sequence of equally-spaced normal distributions of the force. The alignment problem was reduced to shifting two vectors with respect to each other and estimating their alignment score.

The last step of the procedure is a hierarchical clustering classification. The distance measure used for clustering is the negative of the alignment score. Small distances correspond to similar
traces, while large distances correspond to incompatible traces. This distance measure showed to be appropriate to distinguish deviant spectra, e.g. spectra with no attachment or erratic peaks. The first step in hierarchical clustering is connecting the two traces with maximal alignment score in a cluster. The other traces either joined the existing cluster or formed a new one. This was repeated until all traces have been assigned and in the end one cluster containing all traces was obtained. In the process, the extension offsets corresponding to the alignment scores were applied to the traces. If two clusters needed to be merged, the extension offset between the two closest cluster members was used.

As a benchmark, F-x curves from unfolding of bR and NhaA were mixed together. Consistently with the ground-truth, the generated hierarchical tree had two main roots: a bR cluster and a NhaA cluster. This result demonstrated that the distance measure or the alignment score is good enough to distinguish between two different groups of force spectra. The cut of the single-linkage tree was made on the level of the average alignment score plus one standard deviation of the score.

The described procedure includes also a peak classification scheme used for detection of different unfolding pathways. Different unfolding pathways are usually distinguished by the presence or the absence of some peaks; these peaks can be minor side peaks of the same major peak. The peak classification is based on the contour length differences between peaks in different traces. At first, the two peaks with closest $L_c$ values were united into a peak class. The contour length value representing that class was the average between the $L_c$ values of the two peaks. Then the next two closest peaks were selected and either created a new class, or one of them or both joined the existing class. Peaks coming from the same trace were not allowed to enter into the same peak class. Therefore, a peak class contains only peaks from different traces. With this procedure, the F-x curves in the bR cluster were divided into five subgroups. Each subgroup contained a specific combination of peaks corresponding to different unfolding pathways. The so-found five subgroups support the results from a different study on bR [64].

As a general weakness of the method, the susceptibility to large numbers of deviant spectra was noted. Deviant spectra were easily detected in the hierarchical tree but their large number affects the overall quality of the alignment. It was also mentioned that the alignment quality is reduced by F-x curves with short non-contact parts. In that case, the problem is related to a slightly wrong slope of the baseline fit to such short non-contact parts causing a shift in the alignment.

**The analysis approach by Marsico et al. (2006)**

In the study by Marsico et al. [20], the alignment problem was handled using dynamic programming, an approach which we are also going to follow. The algorithm has three major steps: noise reduction in the force spectra, pairwise distances computation by dynamic programming alignment and hierarchical clustering. The procedure was tested on 135 F-x curves coming from unfolding experiments of P50A bacteriorhodopsin (bR) mutant. 61 of these curves were manually selected as
good traces that correspond to the complete unfolding of the protein.

We already mentioned that before any statistical analysis, SMFS data requires careful preprocessing. What we didn’t mention is that spurious curves create obstacles and introduce errors in automatic procedures and hinder the results interpretation. The work by Marsico et al. is a first attempt to address this problem. In the beginning of the procedure, spurious curves were automatically detected and removed. Good traces had to meet two requirements: (1) to have at least one peak with force magnitude higher than two times the standard deviation of the noise; (2) the position of their last peak to correspond to the length of the protein under consideration when fully-stretched. The second requirement implies that the selected traces represent the complete unfolding of the protein and, more importantly, that the identity of the protein is known. The zero-force baseline and the contact point were determined in each F-x curve following the same protocol as Kuhn et al. [19](see above). The noise coming after the detachment peak was removed.

As a next step, noise reduction was performed. The main source of noise in AFM-SMFS experiments are the thermal fluctuations of the cantilever. In fact, the spring constant of the cantilever sets the standard deviation of the noise [66], which is usually between 10 and 40 pN. The noise in each trace was reduced using a dimension reduction with singular value decomposition (SVD). In the bR curves, the standard deviation of the noise after this reduction was 8 pN, in comparison with 14 pN obtained with moving average.

The alignment of the curves was done using global sequence alignment with dynamic programming. In dynamic programming algorithms, each pair of F-x curves is transformed into a pair of force sequences, \(a\) and \(b\). The alignment is then built iteratively by a sequence of moves. There are three possible moves, each one associated with a score: match/mismatch, adding a gap in sequence \(a\) or adding a gap in sequence \(b\). Once the three scores are computed for a pair of points, the move associated with the maximum score is accepted. As a result, the points remain aligned with each other if the match/mismatch score is the highest, or a gap is introduced in one of the sequences correspondingly. The match/mismatch score is proportional to the absolute value of the force magnitude difference divided by the average of the maximum force in the two traces. The match score favors the alignment. The gaps are unfavorable and associated to a penalty. If you have to introduce too many gaps in order to align two sequences, this is an indication that the sequences are very different. The gap opening penalty was 0.002 in the first 10 nm of the trace and 0.8 for the rest of the trace. This choice reflects the presence of non-specific interactions in the very beginning of each curve up to extension 10 nm. The gaps concept suits very well SMFS data since the position of a peak might vary by up to six residues and some peaks might be absent [20]. Once the final pair of points is reached, the final alignment score is computed as the maximum score between the match/mismatch score and the two gaps accordingly. The similarity score between two traces is simply the final alignment score. In Figure 2.13 the power of dynamic programming alignment is demonstrated. In Figure 2.13a all peaks, except the first one, are misaligned but introducing only a
single gap in the first peak (Figure 2.13b) results in optimal alignment between all peaks. In order to cluster the traces, the method of hierarchical clustering was applied. The distance measure was one minus the similarity score. Pairs of traces with a similarity score lower than 0.65 were considered as outliers and were excluded from the clustering.

In order to benchmark the method, force peaks were manually detected and fitted with the WLC model. This information was used for the interpretation of the clustering results. All F-x curves had three main peaks associated with the pairwise unfolding of the protein’s transmembrane helices. In addition to the main peaks, the curves contained minor peaks or side peaks, which varied between traces and might be related to different unfolding pathways. In the case of bR, the side peaks tell us that the helices do not unfold always in pairs [9]. Five subclusters associated with different unfolding pathways of P50A bR mutant were found. The interpretation of the curves is in agreement with the one made by Muller et al. [9] for bR and described in 2.0.4. The success rate of the method in distinguishing between good and bad traces was estimated to 81 % and the success rate of classifying unfolding events was estimated to 76 %.

The analysis approach by Bosshart et al. (2012)

Another automatic procedure for SMFS data analysis specifically designed for high-throughput experiments was developed by Bosshart et al. [21] in 2012. The procedure consists of filtering,
reference-free (RF) alignment and cross-correlation-based classification. This method can process large data sets which contain F-x curves with a force pattern corresponding to the same protein plus noisy traces of heterogeneous nature. The latter and the speed of the method were pointed out as major advantages of the method. The procedure was applied on data sets containing F-x curves coming from unfolding of bR and the soluble multidomain $(Ig27)_8$ protein [67], for a total of $\sim 450,000$ curves.

The first step of the method is a coarse-filtering procedure developed in another study by Bosshart et al. [17]. The traces are filtered according to four requirements:

1. The standard deviation of the force, $\sigma_F$, is computed for the last 5% of data points in each F-x curve. $\sigma_F$ describes the flatness of the non-contact part. Only traces with flat non-contact parts are included in the analysis. This was accomplished by setting a threshold of 20 pN on the maximum allowed $\sigma_F$.

2. F-x curves that have unfolding events at extensions larger than a contour length, $L_{c,\text{max}}$, were omitted. $L_{c,\text{max}}$ corresponds to the contour length of the fully unfolded protein $(Ig27)_8$ in the case of [67]. This criterion ensures that only curves that correspond to the complete unfolding of the protein under investigation are analyzed. Clearly this criterion requires the knowledge on the sample protein composition. A maximum allowed extension value, $x_{\text{max}}$, is defined and chosen slightly below $L_{c,\text{max}}$ because the final unfolding event in a SMFS experiments occurs before the protein is fully-stretched.

3. The different attachment sites between the tip and the protein generate horizontal shifts in F-x curves. That means that the last unfolding event in different spectra can appear in an interval of extension values. The parameters $x_{\text{min}}$ and $x_{\text{max}}$ are set to define this interval. The last unfolding event must occur at extension values between $x_{\text{min}}$ and $x_{\text{max}}$ for a F-x curve to be accepted.

4. F-x curves with negative force peaks are excluded from the analysis. A check on the presence of negative force peaks (below a certain threshold) between extensions $x_{\text{low}}$ and $x_{\text{max}}$ is performed. The value of $x_{\text{low}}$ is set to 5 nm accounting for the non-specific interactions in the beginning of a F-x curve.

As a result of this filtering procedure, the number of bR traces jumped off from 450,000 to 1,534 ($\sim 0.3\%$). In the $(Ig27)_8$ data set, 1,074 traces remained after filtering.

As a next step, the contour length ($L_c$) histograms for the remaining traces were estimated. The WLC equation (Eq.2.1) was solved for every data point using a persistence length of 0.4 nm. Points with force values below 45 pN and above 500 pN were excluded because the WLC model is not valid for these force values. The histogram bin size is 1 nm, which roughly corresponds to 3 a.a. per bin. Additional filtering based on the properties of the auto-correlation function (ACF) of
the Lc histograms was applied. The position and the amplitude of the first side peak in the ACF were used as a filter for detection of characteristic unfolding events in the F-x curves. Their values were obtained form the ACF of the average Lc histogram for an entire data set. For example, the position of the first side peak reflects the periodicity of the pattern. In the case of the multidomain protein (Ig27)$_8$, the first side peak is positioned at 27.6 nm, which is an agreement with the interpeak distances published in the literature [67, 8]. We remark that also this procedure requires the previous knowledge of the identity of the protein that is analyzed. Next, the peaks in each Lc histogram were automatically detected and their positions were stored in a matrix. To avoid overestimation of the number of peaks, the Lc histograms were smoothed. The traces were divided into peak-number groups depending on the number of peaks found in their Lc histograms. In general, the larger the number of peaks in the Lc histograms, the more the details and the side peaks that could be found in the F-x curves.

The reference-free alignment algorithm [68] was then performed separately in each peak-number group. The algorithm can be divided in four blocks.

1. Two randomly selected Lc histograms are cross-correlated and aligned based on the correlation maximum. An arithmetic average of the two histograms is then computed. A third Lc histogram is randomly chosen, aligned and then added to the arithmetic average. This is repeated until all histograms belonging to that peak-number group are included in the arithmetic average. A Lc histogram is counted only once. The final arithmetic average is the global average representing that particular group.

2. Each Lc histogram in the group gets aligned to the global average on the basis of their correlation maximum.

3. Each Lc histogram is subtracted from the global average and a temporary average is computed. The histogram is then aligned to that temporary average and a new arithmetic average is computed. This is repeated for all histograms.

4. The global averages of all peak-number groups get aligned to each other. This makes the comparison between different peak-number groups easier and overcomes the different offsets which might turn out to be problematic. In the end, all histograms in a peak-number group get aligned to the final global average.

The goal of this procedure is determining the unfolding fingerprint of a given protein. The power of alignment in solving this problem is illustrated in Figure 2.14 for the (Ig27)$_8$. It becomes clear that the characteristic unfolding pattern of the protein is hardly seen in the superimposition of the unaligned spectra, while obvious in the aligned superimposition.
In the next step, the $L_c$ histograms in each peak-number group are classified using cross-correlation by a procedure similar to K-means clustering\cite{69}. The aim is to distinguish between curves belonging to the same peak-number group and group them into classes based on minor details. In this approach, a small number histograms from the same group are randomly selected. Each of these histograms starts its own class. Then each class is filled with a user-defined number of randomly chosen histograms coming from the same group. For each class, the initial class average is computed as the arithmetic sum-normalized average of the histograms in that class. A certain $L_c$ histogram can contribute to the initial class average only once. One by one, all histograms belonging to that peak-number group get aligned to the initial class averages and assigned to the class with which they have the highest correlation. Once all histograms are assigned to a class, the global class averages are recomputed. Then all histograms get aligned to the updated class averages and this refinement cycle is repeated. After each refinement cycle a series of statistical descriptors are computed to monitor the convergence process. The number of refinement cycles is another parameter in this procedure. The obtained classes were dominated by the high probability densities of the main peaks. It is pointed out that cross-correlation based classification can be useful in spotting contaminating spectra in a peak-number group so that they can be excluded from the subsequent analysis.

Next, the method of principal component analysis (PCA) followed by standard K-means clustering was used to subclassify the classes obtained with cross-correlation. The PCA method is applied to the peak positions of the aligned $L_c$ histograms. Even if all F-x curves share the same number of peaks in their $L_c$ histograms that doesn’t mean they represent the same unfolding pathway. Indeed
the peaks might have different positions. This is the reason why the PCA was applied to the peak positions within a class. K-means clustering is then applied in the space of the principal components; the number of clusters, \( n_{cl} \) is a user-defined parameter. It is normally set to a value slightly higher than the number of expected clusters. The number of expected clusters in this approach is roughly estimated from the PCA factor map. The number of estimated clusters in the largest class in the bR peak-number group with six peaks, was 3. In all of them, four main peaks were present, while the differences between the clusters arose from the densities of the side peaks. The combination of PCA and K-means clustering did not work in all cases. For example, in the largest class of the peak-number group with seven peaks, it was not possible to identify any clusters in the PCA factor map and the K-means clustering was not executed. In addition, the described method was successfully used to localize the unfolding barriers in the bR structure. All peaks in the average \( L_c \) histogram were related to a position in the bR helices.

The analysis approach by Galvanetto et al. (2018)

Recently, Galvanetto et al. developed an open-source software for analysis of F-x curves, named Fodis (for Force-distance software) [22]. The software operates in MATLAB and offers a variety of tools for the manipulation and statistical analysis of F-x curves. It was tested on SMFS data from unfolding of the cyclic nucleotide-gated CNGA1 channel overexpressed in oocytes membranes [13].

Fodis implement a filtering procedure, also based on the prior knowledge of the protein that is analyzed. Before filtering the \( L_c \) histograms of all traces need to be estimated. The persistence length used for the \( L_c \) transformation was 0.4 nm. Only \( L_c \) values corresponding to forces larger than 30 pN were included in the histograms. The filtering criterion in the first step is based on the length of the F-x curves and adopts the fully-stretched condition. The fully-stretched condition implies that F-x curves longer or shorter than the \( L_c \) value corresponding to the fully-stretched protein can be discarded. The trace length is determined by the position of the last peak in the \( L_c \) histogram. The authors noted that the computed contour length strongly depends on the persistence length value, which can vary between 0.3 and 0.8 nm [9]. For this reason, instead of fixing a \( L_c \) threshold, a window of values centered around the expected length was defined. The size of the window is recommended to be at least 30 % of the protein expected length. In the CNGA1 example, the window was between 210 and 360 nm. We remark that this procedure is robust and accurate, but requires the knowledge of the identity of the protein.

In the second filtering step, traces with high non-specific adhesion (F>150 pN) in the very beginning of the force spectra (up to 70 nm) were discarded.

The third filtering step is based on cross-correlation and finds similar F-x curves. The similarity between two traces is given by the cross-correlation computed for their \( L_c \) histograms. A similarity matrix containing the correlation values for all pairs of traces is generated. The symmetric approximate minimum permutation algorithm [11] is applied to the matrix in order to detect clusters of
similar traces. The obtained clusters can be further inspected and analyzed.

The reference-free alignment algorithm [21] described above was implemented in Fodis with some modifications. The first modification is related to the alignment of the force spectra, which is performed by adding a Gaussian curve centered at zero extension to the cross-correlation function for each pair of histograms. In this manner, the alignment in which the zero points of the two curves match each other is favored. The second modification is related to the division of force spectra in peak-number groups. The global histogram generated from all $L_e$ histogram is manually divided into peak intervals for the purposes of the analysis. As an alternative to manual intervention, the software offers an option in which the peak intervals are automatically determined and extracted from the global histogram of maxima. This plot shows the $L_e$ peaks with highest probability, or the most abundant $L_e$ peaks in the entire data set. The global histogram of maxima reveals the main unfolding pattern in the particular group of force spectra. Each trace is then transformed into a binary string of size equal to the total number of peak intervals. A value of 0 is assigned if no peak is present in the corresponding interval; a value of 1 is assigned if there is a peak in that interval. From the binary representation, two sequences are generated: $N$ and $P_n$. The sequence $N$ contains information for the interval position occupied by the peak; if the peak is in the fourth interval, the value of this peak in $N$ will be 4. The sequence $P_n$ keeps track on the order in which the peak appears along the trace; if it is the third peak in the trace but is positioned in the fourth peak interval, the value it gets will be 3 not 4. The plot sequence $N$ versus sequence $P_n$ for all traces of interest is quite informative. It enables the identification of different unfolding pathways and offers a graphical summary of the number and the position of occurrence of unfolding events. In Figure 2.15 the power of this algorithm is illustrated for 106 CNGA1 curves. The thickness of the lines reflects the abundance of traces following that path.
Figure 2.15: Path plot for a subset of 106 CNGA1 F-x curves. a. The path plot was generated using 6 manually selected $L_c$ intervals. Each color represents a possible unfolding pathway. The thickness of the line represents the curves population; their number is shown in the legend bar. b. Global histograms of four identified clusters and the superimposition of sets of traces belonging to each cluster. From ref. [22]
Chapter 3

Molecular modeling of single molecule force spectroscopy.

The phototransduction process, in which light undergoes a biochemical transformation to electric signals, takes place in the rod outer segment (rod OS) of photoreceptor cells. The rod OS is a cylindrical biological structure of highly ordered membranes. It hosts discrete membranous discs enclosed by the plasma membrane of the cell [70]. The disc membranes and the plasma membrane are two membranes with a different lipid and protein composition and independent functions. The plasma membrane is enriched in cholesterol and saturated fatty acids compared to the discs [23]. Rhodopsin is the only membrane protein present both in the discs and in the plasma membrane [23]. Moreover, rhodopsin is the dominant protein by mass in both membranes [23]. Strikingly, rhodopsin is active (i.e. initiates the phototransduction cascade) only in the discs.

The starting point of our investigations were experiments in which rhodopsin was unfolded using AFM-SMFS from the discs and from the plasma membrane of *Xenopus laevis* retinas [12]. Surprisingly, the obtained unfolding patterns were quite different from each other. Normally, the unfolding pattern of a protein in F-x curves is considered an unique fingerprint of that protein. However, in AFM-SMFS experiments on membrane proteins, the protein is not only stretched but it is also extracted out of the membrane. Therefore, the obtained unfolding pattern may depend on the composition of the membrane in which the protein is embedded and with which it interacts. Following this line of reasoning, we formulated the hypothesis that the different cholesterol concentration of the discs and the plasma membrane introduces changes in the mechanical stability of rhodopsin embedded in the two lipid environments.

To test this hypothesis, we designed a coarse-grained model of the rhodopsin-membrane system in which the membrane is modeled implicitly with an additional potential energy term, $V_{MEMBR}$,
which favors the native contacts between the transmembrane hydrophobic residues and the membrane. Then, we performed molecular dynamics (MD) simulations of the pulling AFM experiments to evaluate the mechanical stability of rhodopsin. The only free parameter in the model is $\epsilon_{\text{MEMBR}}$ which defines the strength of the membrane potential, thus reflecting the different membrane compositions.

The activation of rhodopsin in the phototransduction cascade is associated with a series of conformational changes. Rhodopsin (PDB: 1U19) is known to be the inactive conformation and metarhodopsin II (PDB: 3PXO) is known to be the active conformation. We also investigated the effect of the higher cholesterol concentration in the plasma membrane on the flexibility of rhodopsin. The assumption we made is that a cholesterol-rich membrane, like the plasma membrane, might stabilize the inactive conformation of rhodopsin, hindering its transition to the active metarhodopsin II. We used all-atom MD simulations in a combination with free energy perturbation theory to address this issue.

3.1 Experimental results.

Our work is motivated by the experiments performed by Maity et al. [12] in the rod OS of *Xenopus laevis* retinas. They used the AFM both as an imaging technique and as a SMFS tool. The experiments were made in the discs and in the plasma membrane of the rod OS. The obtained F-x curves revealed two strikingly different patterns describing the unfolding of rhodopsin in the disks and in the plasma membrane. These observations together with the fact that rhodopsin is active only in the discs were calling for a molecular explanation.

3.1.1 AFM imaging.

Rod OSs from dark-adapted *Xenopus laevis* retinas were isolated and purified plasma membrane and discs were extracted as described in ref. [12]. The AFM (JPK NanoWizard 3) was used in tapping mode to image 500 x 500 nm$^2$ patches of the cytoplasmic side of the discs and the rod OS plasma membrane. The cantilever spring constant was $\sim 0.08$ N/m. The topography images revealed two types of protrusions in the plasma membrane: one with relative height 1.5 ± 0.7 nm and another with height 4.9 ± 0.9 nm (Figure 3.1a). In the discs only the low protrusions with height around 1.5 nm were present. Subsequently, SMFS experiments were performed to identify the molecular origin of the protrusions.
CHAPTER 3. MOLECULAR MODELING OF RHODOPSIN.

Figure 3.1: (a) AFM topography image of the rod outer segment (OS) plasma membrane. The numbered circles point to protrusions in the lipid bilayer with different height: 1,2,3 point out high protrusions with $4.9 \pm 0.9$ nm height; 4, 5 point out low protrusions with $1.5 \pm 0.7$ nm height. (b) F-x curves obtained after pulling the selected protrusions 1,2,3,4,5. Protrusions 1,2,3 were identified as native CNG channels. Protrusions 4,5 were identified as rhodopsin. From ref. [12]

3.1.2 SMFS experiments.

The same AFM tip used for imaging was also used for the SMFS experiments. The cantilever was calibrated before each experiment. The AFM tip was pushed to the surface with a contact force of 1 nN for 0.5 s and then retracted at a constant speed 500 nm/s. The high and the low protrusions in the plasma membrane were localized and pulled out of the membrane, obtaining longer and shorter F-x curves respectively (Figure 3.1). The proteins in the high protrusions in the plasma membrane were identified as native CNG channels after validation towards the F-x curves obtained from unfolding of the CNGA1 channel [13]. The curves obtained from the unfolding of the proteins in the lower protrusions were analyzed as described in ref. [13] and assigned to the unfolding of rhodopsin.

In Figure 3.2, F-x curves resulting from the unfolding of rhodopsin in the discs and in the plasma membrane are plotted. The unfolding of rhodopsin from the discs is characterized by 5 major force peaks, while the unfolding from the plasma membrane - by 7 major force peaks. Furthermore, the forces, required to unfold rhodopsin from the plasma membrane are larger compared to the discs. The average force to unfold rhodopsin from the plasma membrane is $136 \pm 135$ pN, compared to $74 \pm 140$ pN from the discs.

One can notice that the $L_c$ of the last peak in both membranes corresponds to roughly 240 a.a.. At the same time, rhodopsin from the rod OS of *Xenopus laevis* retinas contains 354 a.a.. This is explained by the fact that the functional form of rhodopsin is known to contain a disulfide bridge
between Cys110 and Cys187 [71]. The bridge connects two of the transmembrane helices, C and D. For this reason the final $L_c$ is smaller than the one corresponding to the total number of amino acids.

### 3.2 Theoretical approach.

The starting point of our investigation is that the unfolding pattern of the same protein, rhodopsin, in the discs and in the plasma membrane is very different. The simplest explanation for this is in the composition of the two lipid environments. The plasma membrane is known to contain more cholesterol compared to the disc membranes [23]. In addition, cholesterol is known to contribute to the membrane’s stiffness making it more rigid. Furthermore, cholesterol can participate in hydrophobic interactions with the transmembrane hydrophobic residues exposed to the membrane stabilizing the rhodopsin molecule. All of this possibly alters the mechanical stability of rhodopsin like the experimental data suggests.

Another difference between rhodopsin in the discs and rhodopsin in the plasma membrane is...
related to the protein activation. It is well known that the transition of rhodopsin to its active form, metarhodopsin II, is accompanied by a series of conformational changes. Rhodopsin is functional only in the discs. A possible explanation for this is that the higher cholesterol concentration in the plasma membrane favors the inactive rhodopsin conformation impeding the phototransduction process.

In order to address these issues, we first developed a coarse-grained model of the rhodopsin-membrane system. We started from a Go-like model introduced by Cieplak et al. [24] (see subsection 2.0.5). We modeled the presence of the membrane with an additional potential energy term accounting for the protein-membrane interactions. As far as we know, this way of modeling the membrane hasn’t been reported before in the literature. Our model has one free parameter, $\epsilon_{\text{MEMBR}}$, which determines the strength of the rhodopsin-membrane interactions. The main goal of this model is to reproduce the characteristic F-x curves obtained experimentally when rhodopsin was unfolded from the discs and from the plasma membrane.

In the second part, we examined the effect of the higher cholesterol content in the plasma membrane on the rhodopsin flexibility with all-atom MD simulations in a lipid bilayer. We used a statistical mechanics approach to extract qualitative description of the rhodopsin behavior in a cholesterol-rich membrane from the atomistic simulations performed in a cholesterol-free lipid bilayer.

### 3.2.1 The molecular model of the rhodopsin-membrane system.

We represented the rhodopsin polypeptide chain in a simplified manner as a chain of beads. We implemented this using the coarse-grained Go-like model developed by Cieplak et al. [24]. The detailed description of the model is provided in subsection 2.0.5. Briefly, the amino acids are represented as beads centered around the $C_\alpha$ atoms. The beads are tethered together at the typical peptide bond length, 3.8Å, by a harmonic potential. The native chirality of the protein is conserved by a chirality potential. The non-covalent interactions are divided into native and non-native depending on the contact type in the native state. The contact type is determined from the distances between all heavy atoms in the atomistic representation of the native state using the procedure of Tsai et al. [63]. According to this procedure, if the distance between two atoms is larger than the sum of their van der Waals radii multiplied by 1.244, the contact is non-native. If the distance is smaller - the contact is native. The potential used to describe the interactions between native contacts is the Lennard-Jones potential with a minimum corresponding to the distance between the $C_\alpha$ atoms in the native state. The potential used to describe the interactions between non-native contacts is purely repulsive. It has the form of the repulsive part of the Lennard-Jones potential corresponding to a minimum at $\sigma = 5\text{Å}$. This potential is shifted upward with zero energy at $\sigma = 5\text{Å}$ and vanishes at larger equilibrium distances. The protein system evolves in time according to the Langevin equation (see below, Eq. 3.2). Stretching is implemented in the following manner: one of
CHAPTER 3. MOLECULAR MODELING OF RHODOPSIN.

the two protein ends is kept fixed, while the other, attached to a harmonic spring, is being pulled in the z-direction. The separation of the moving end from its origin corresponds to the cantilever displacement in the AFM experiment. This model has been successfully used to describe the elastic properties of globular proteins like titin [25], calmodulin [26] and ubiquitin [27].

Cieplak et al. used the same model in combination with all-atom MD simulations to describe the unfolding of the membrane protein bacteriorhodopsin [28]. The atomistic simulations were necessary to determine the initial state of the rhodopsin-membrane system. The membrane was modeled by a POPC lipid bilayer. Rhodopsin was embedded in the lipid bilayer and the system was equilibrated. Afterwards, the equilibrated rhodopsin-membrane system was coarse-grained as follows: rhodopsin was modeled in the way we just described and the POPC lipids were represented by their carbon atoms (see subsection 2.0.5). Native and non-native contacts between the protein and the lipids were determined again following the Tsai et al. [63] protocol. The potentials describing the native and non-native interactions were of the same functional forms. During the stretching, the coarse-grained lipid bilayer was kept rigid and not allowed to adjust to the new configuration of the system. In this manner, when the protein is pulled out, the space it was filling in the membrane remains empty, leaving a sort of a hole. This is unrealistic, since in a real membrane the phospholipids are able to fill the space left empty by the protein.

In order to circumvent this approximation, we model the effect of the membrane with an extra potential energy term, \( V_{MEMBR} \), applied only along the z-axis and which acts in a different manner on the different residues based on their chemical nature and on their native contact with the membrane. Initially, all amino acids are divided into hydrophobic, hydrophilic and unspecified according to the Kyte and Doolittle hydrophobicity scale [72]. We used the program GetArea [73] to compute the solvent accessible surface area (SASA) and to determine which amino acids in the experimental structure are exposed to the solvent, with the membrane being the solvent in our case. If a residue \( i \) is membrane exposed and hydrophobic, it contributes to \( V_{MEMBR} \) as follows:

\[
V_{MEMBR}(z_i) = \begin{cases} 
0, & \text{if } |z_i| < \frac{l}{2} \\
\epsilon_{MEMBR}, & \text{if } |z_i| < \frac{l}{2} + 3\AA \\
\epsilon_{MEMBR}\left(\frac{|z_i| - \frac{l}{2}}{3}\right), & \text{otherwise}
\end{cases}
\]  

(3.1)

where \( z_i \) is the z-coordinate of residue \( i \) and \( l \) is the membrane thickness in Å. Here we take \( l = 33\AA \). \( \epsilon_{MEMBR} \) is the most important parameter in our model. It is a measure of the strength of the membrane potential: a larger value of \( \epsilon_{MEMBR} \) defines a highly hydrophobic membrane such as a cholesterol-rich membrane. The same potential multiplied by -1 acts on the hydrophilic residues exposed to the membrane.

A scheme representation of the membrane potential is presented in Figure 3.3. When a hydrophobic residue in native contact with the membrane has a z-coordinate corresponding to the hydrophobic core of the membrane formed by the lipid tails, \( V_{MEMBR} = 0 \). If the z-coordinate
value corresponds to the $\sim 3 \text{ Å}$ thick layer occupied by the lipids head groups [74], the potential is disfavoring this conformation moderately. If the residue $z$-coordinate value falls in the fully hydrated layer, the penalty is larger, equal to $\epsilon_{MEMBR}$. The role of the membrane potential is to keep the hydrophobic residues in contact with the membrane, inside the membrane.

Figure 3.3: (a) Schematic representation of the cross section of a lipid bilayer including the fully hydrated hydrophilic heads (blue), the fully dehydrated hydrophobic tails (dark green) and a short intermediate region on the border between the hydrophilic heads and the hydrophobic tails with partial hydration. (From ref. [74]). (b) A scheme representing the membrane potential. The colors of the lines match the colors used in panel (a) to illustrate different regions in lipid bilayers depending on their accessibility to water molecules.

3.2.2 Coarse-grained MD simulations setup.

The initial conformation is set by the coordinates of all atoms in the bovine rhodopsin crystallographic structure (PDB code: 1U19) (Figure 3.4). This initial conformation is considered to be the rhodopsin’s native conformation in this model. There is a small difference between rhodopsin from *Xenopus laevis* retina examined in the experiments and bovine rhodopsin used in our model. The latter is 6 a.a. shorter. We note that the disulfide bridge between Cys110 and Cys187 is present in the PDB structure.
Figure 3.4: (a) Rhodopsin (PDB: 1U19) all-atom graphical representation colored by the amino acids names. (b) Graphical representation of coarse-grained rhodopsin. Each bead corresponds to the $C^\alpha$ atom of the residue.
The only parameter we changed in comparison with Cieplak et al. [28] is the value of the spring constant in the harmonic potential, $k_{BB}$ (Eq. 2.2). We used $k_{BB} = 33.34 \frac{\epsilon}{\AA^2}$, while Cieplak et al. used $k_{BB} = 100 \frac{\epsilon}{\AA^2}$. The reason we did this is that our system was unstable with the recommended value and was exploding from time to time. $k_{BB} = 33.34 \frac{\epsilon}{\AA^2}$ is the smallest value that allowed us to use a large timestep of 15 fs in the MD simulations.

The protein-membrane CG system evolves in time through the Langevin equation:

$$m\ddot{r}_i = -\gamma \dot{r}_i + F_{c,i} + \sqrt{2\gamma k_B T} \xi$$

(3.2)

where $r_i$ is the position of $i$-th bead of the chain, $F_{c,i}$ is the force acting on bead $i$, $T$ is the temperature and $\xi$ is a Gaussian noise term. The friction coefficient $\gamma$ is equal to $8.14 \times 10^{-4}$. The equations of motion were solved with the Velocity Verlet integration scheme with timestep $\Delta t = 15$ fs.

For the pulling simulations, an additional harmonic spring attached to the C-terminal of rhodopsin was introduced, while the N-terminal was kept fixed. The outer end of the spring was pulled at constant velocity $v_{pull}$ in the $z$-direction like in Cieplak et al [28]. The extensions measured in an AFM experiment, here correspond to the deviations of the pulled end from its origin.

In order to fix the effective temperature of the system, we performed MD simulations of our model of rhodopsin for a set of preselected temperatures in two cases. First, the membrane potential was set to 0 and no pulling force was applied. The average root mean square deviation (RMSD) was computed for different temperatures. Then we did the same applying the membrane potential with $\epsilon_{MEMBR} = 10\epsilon$, the largest value we considered. The results are plotted in Figure 3.5. The jump in the RMSD indicates that the melting temperature has been reached. The melting temperatures in the two cases are quite similar. The melting temperature for $\epsilon_{MEMBR} = 0\epsilon$ is 0.65$\epsilon$, while for $\epsilon_{MEMBR} = 10\epsilon$ it is 0.7$\epsilon$. This means that the membrane potential changes the melting temperature only slightly. Knowing that usually the melting temperatures of membrane proteins are 10-20% above room temperature, we took $k_B T = 0.52\epsilon$ as the room temperature in our simulations.
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Figure 3.5: Average RMSD computed from MD simulations of the coarse-grained rhodopsin model at different temperatures. The RMSD is plotted with errorbars. The blue line represents the results from the simulations with $\epsilon_{MEMBR} = 0\epsilon$. The red line represents the results from the simulations with $\epsilon_{MEMBR} = 10\epsilon$. No pulling force was applied in these simulations.

3.2.3 Validation of the membrane potential.

In order to validate the capability of the membrane potential to capture the hydrophobic effect of the membrane we performed four additional simulations with our model using the same set of MD parameters, $\epsilon_{MEMBR} = 10\epsilon$ and $v_{pull} = 0$, (no pulling force). The number of hydrophobic residues was artificially changed in each simulation. We took the amino acid LEU which is the most abundant transmembrane hydrophobic residue in rhodopsin, and mutated it to an unspecified hydrophilic residue. We performed 1, 4, 11 and 23 mutations, where 23 is the total number of transmembrane LEUs in rhodopsin. For each configuration we computed the average potential energy from the MD run and the SASA occupied by the hydrophobic residues, $A_{PHOB}$. In Figure 3.6, the energy is plotted as a function of $A_{PHOB}$. As $A_{PHOB}$ is increasing the energy is decreasing due to the hydrophobicity effects included in the membrane potential. This indicates that the functional form we use is able to capture, at least qualitatively, the effect of transferring a hydrophobic moiety into an hydrophobic environment: the larger the area of the moiety, the lower the average energy.

Here we also verified that the exact choice of the thickness of the intermediate membrane layer does not affect the qualitative behavior of $V_{MEMBR}$. We performed the same set of four MD simulations described above for thickness 1 and 5 Å. In Figure 3.6 we show that the trend of the average energy as a function of the hydrophobic area remains qualitatively similar even if the thickness parameter is changed rather dramatically.
3.2.4 All-atom MD simulations.

We used all-atom MD simulations to attempt explaining why rhodopsin in the discs is active, while it remains inactive in the plasma membrane. Since the activation of rhodopsin is related to conformational changes in the protein, our coarse-grained model no longer provides a physically meaningful description. Indeed, in this model the native structure is strongly favored energetically. This assumption is the basic limitation of Go-like models. Practically, the protein is incapable of moving away from its native conformation because if it attempts to do so, it has to pay a very high energy cost. This condition automatically excludes the possibility that the model may occupy an energy minimum different from the one corresponding to its native state. That is not the case in all-atom MD simulations, where the protein is allowed to freely explore the conformational space. The possibility of observing different stable conformers is necessary in order to carry on the analysis described in the following subsection 3.2.5.

We therefore performed all-atom molecular dynamics with the software Gromacs 4.6.7 [75]. Rhodopsin (PDB code: 1U19) is oriented following the OPM [76] database model and embedded in a pre-equilibrated lipid bilayer with 128 DPPC and 3,655 water molecules [77] using the g_membed [78] tool of Gromacs. The protein charge, -3, is neutralized with 3 sodium (Na+) ions. The box is enlarged in the z-direction to yield the dimensions 6.88x6.91x17.33 nm, resulting in a
system of 1 protein, 118 DPPC and 20,826 water molecules and 3 Na+. Periodic boundary conditions are applied. The force field GROMOS96 53a6 [79] with included Berger lipids parameters [80] for the DPPC molecules is used. All bonds are constrained to their equilibrium lengths with the LINCS algorithm [81]. The non-bonded interactions are described by Lennard-Jones potential with a cutoff at 1.2 nm. The electrostatic interactions are estimated by the particle mesh Ewald (PME) method [82] with cutoff of 1.2 nm. The equations of motion are integrated with the leap-frog algorithm, using a time step of 2 fs. The equilibration is performed in three steps: (1) The system was first heated to a temperature of 323 K in 1 ns keeping the protein backbone fixed; (2) then a 7 ns run in a NPT ensemble is performed, with a surface tension equal to 28 mN/m (corresponding to a -55.56 bar pressure) in the x and y directions and 1 bar in the z direction with a semi isotropic Parrinello-Rahman barostat [83]; temperature is kept fixed at 323 K with Berendsen thermostat [84]; (3) Finally, a 70 ns production run is performed.

### 3.2.5 Estimating the effect of membrane hydrophobicity on rhodopsin flexibility.

Here we derive a set of equations which will allow us to evaluate the effect of the membrane hydrophobicity on rhodopsin flexibility from the atomistic MD simulation of rhodopsin in the cholesterol-free DPPC bilayer.

The partition coefficient per unit area of a hydrophobic molecule between a cholesterol-free membrane and a membrane with a given cholesterol content can be used to measure the effect of a change in the cholesterol content in the membrane. This quantity can be in principle measured experimentally, and it has been estimated for a few compounds by atomistic simulations. We denote this partition coefficient by $\gamma$. We model the effect of the change in the membrane hydrophobicity by adding to the ordinary potential energy function, an extra term of the form:

$$V_\gamma(x) = -\gamma A(x)$$

(3.3)

where $A(x)$ is the transmembrane hydrophobic SASA in configuration $x$. This term favors configurations with large $A$ and vanishes if $\gamma = 0$. The functional form of $V_\gamma$ is consistent with the mesoscopic definition of the hydration free energy, which should be proportional to the surface of the molecule. It is also consistent with the apolar contribution to the free energy of solvation used in several implicit solvation models [85, 86].

The probability distribution function, $P_\gamma(x)$, is given by:

$$P_\gamma(x) = c e^{-\frac{V_0(x)}{kT} - \gamma A(x)} = c' e^{\frac{\gamma A(x)}{kT}} P_0(x)$$

(3.4)

where $c$ and $c'$ are normalization constants, $V_0(x)$ is the potential energy function and $P_0(x)$ is the
canonical probability distribution of the system when $\gamma = 0$.

Then, we estimate the change in the probability distribution as a function of a geometrical observable, the angle $\alpha$, due to the additional term. The joint probability distribution as a function of $\alpha$ and of the hydrophobic SASA, $A$, for $\gamma = 0$ is given by:

$$P_0(\alpha, A) = \int dx \delta(\alpha(x) - \alpha)\delta(A(x) - A)P_0(x)$$  \hspace{1cm} (3.5)

If $\gamma \neq 0$, the joint probability distribution is:

$$P_\gamma(\alpha, A) = \int dx \delta(\alpha(x) - \alpha)\delta(A(x) - A)P_\gamma(x) =$$

$$= \int dx \delta(\alpha(x) - \alpha)\delta(A(x) - A)e^{\gamma A(x)/k_B T}P_0(x) = ce^{\gamma A / k_B T}P_0(\alpha, A)$$  \hspace{1cm} (3.6)

where $c$ is a normalization constant. This equation allows estimating the probability distribution for any value of $\gamma$ from the probability distribution measured in a reference condition, for example in a cholesterol-free membrane. Finally, the probability distribution as a function of $\alpha$ alone is given by:

$$P_\gamma(\alpha) = \int dA P_\gamma(\alpha, A) = c \int dA e^{\gamma A / k_B T}P_0(\alpha, A)$$  \hspace{1cm} (3.7)

The probability distribution $P_0(\alpha, A)$ entering in this equation is estimated from the atomistic MD trajectory on a 100x100 regular grid ranging between 2.58 and 2.83 rad in $\alpha$, between 4,009.88 and 4,737.92 in $A_{HPHOB}$ using a Gaussian kernel estimator [87] with Gaussian variance equal to 4 grid spacing in both directions. The protein residues considered as hydrophobic transmembrane are listed in Table 3.1.
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3.3 Results.

3.3.1 Coarse grained MD simulations of the unfolding experiments.

In order to reproduce the experimental curves obtained from unfolding of rhodopsin in the discs and the plasma membrane, we designed a model of the rhodopsin-membrane system in which the membrane is accounted for by an extra potential energy term, $V_{MEMBR}$, with strength determined by $\epsilon_{MEMBR}$. The different values of $\epsilon_{MEMBR}$ can be associated with membranes with different hydrophobicity as a result of their different lipid composition. The larger the $\epsilon_{MEMBR}$ value, the larger the membrane hydrophobicity effect felt by the protein. Since no other effects are included in this model, changes in the simulated F-x curves should arise only due to variations in the $\epsilon_{MEMBR}$ values.

We performed four series of molecular dynamics (MD) simulations of unfolding of rhodopsin using four different $\epsilon_{MEMBR}$ values: 4.03$\epsilon$, 5.64$\epsilon$, 7.25$\epsilon$, and 10$\epsilon$ (Figure 3.7a-h) By increasing the value of $\epsilon_{MEMBR}$, the forces required to unfold the protein become larger and larger, the simulated F-x curves change their shape and more force peaks start appearing.

<table>
<thead>
<tr>
<th>HYDROPHOBIC RESIDUES</th>
<th>Number</th>
</tr>
</thead>
<tbody>
<tr>
<td>ILE</td>
<td>48, 54, 75, 123, 133, 154, 205, 213, 214, 217, 219, 255, 256, 259, 263, 275, 286, 290, 305, 307</td>
</tr>
<tr>
<td>LEU</td>
<td>40, 46, 47, 50, 57, 59, 72, 76, 77, 79, 84, 95, 99, 112, 119, 125, 128, 131, 165, 172, 216, 262, 266</td>
</tr>
<tr>
<td>VAL</td>
<td>81, 87, 129, 130, 157, 162, 173, 204, 209, 210, 218, 254, 258, 271, 300, 304</td>
</tr>
<tr>
<td>PHE</td>
<td>37, 45, 52, 56, 85, 88, 91, 115, 116, 159, 203, 208, 212, 220, 221, 261, 273, 276, 287, 293, 294</td>
</tr>
<tr>
<td>ALA</td>
<td>41, 42, 80, 82, 117, 124, 132, 153, 158, 164, 166, 168, 169, 260, 269, 272, 292, 295, 299</td>
</tr>
</tbody>
</table>

Table 3.1: Hydrophobic transmembrane residues and their numbers in rhodopsin (PDB:1U19).
Figure 3.7: (a,c,e,g) Simulated force-distance traces for bovine rhodopsin (PDB code: 1U19) pulled by the C-terminal at $k_B T = 0.52 \epsilon$ for different values of the parameter $\epsilon_{MEMBR}$ as indicated ((a) 4.03\,\epsilon, (c) 5.64\,\epsilon, (e) 7.25\,\epsilon, (g) 10\,\epsilon). Each plot represents the superimposition of 10 traces obtained from 10 independent simulations. (b,d,f,h) Cartoon representations of the order in which the transmembrane helices unfold in the simulations of the left panels, as derived by a visual inspection of the trajectories. The color map is the same as for the traces. The numbers on top of each peak correspond to the length of the stretch that is unfolded up to the time step when the force drops (expressed in number of amino acids, n).
For $\epsilon_{\text{MEMBR}} = 4.03\epsilon$, the obtained F-x curves contain three major force peaks with force amplitudes between 150 and 200 $\frac{\epsilon}{nm}$ (Figure 3.7a and b). The visual inspection of the trajectories revealed pairwise unfolding of helices E and F, and A and B. Helices C and D remain bound together by the disulfide bond. Helix G unfolds separately including the stable structural domain H located inside the cell and the C-terminus.

When the $\epsilon_{\text{MEMBR}}$ value was changed to 5.64$\epsilon$, the observed unfolding pattern also started changing. The major force peaks became four and their force magnitude got up to 300 $\frac{\epsilon}{nm}$ (Figure 3.7c and d). Transmembrane helices G, F, and E unfolded separately, while A and B unfolded pairwise. These qualitative features are consistent with the interpretation of the experimental curves from pulling rhodopsin in the discs [12].

With $\epsilon_{\text{MEMBR}} = 7.25\epsilon$, the major force peaks are still four, but the third major peak got split in two, illustrating the stepwise unfolding of helix E (Figure 3.7e and f). The peaks’ force magnitudes went above 300 $\frac{\epsilon}{nm}$. Helices G and F unfolded separately and helices A and B were unfolded together.

When the value of $\epsilon_{\text{MEMBR}}$ was further increased to 10$\epsilon$, the total number of peaks increased to seven and the forces went up to 500 $\frac{\epsilon}{nm}$ (Figure 3.7g and h). All $\alpha$-helices were unfolded sequentially, except for helices D and C, linked together by the disulfide bond, and $\alpha$-helices F and E which unfolded in two steps. These results resemble the experimental ones when rhodopsin was unfolded from the plasma membrane [12].

The number of a.a. unfolded at each step obtained from the unfolding simulation is in very good agreement with the number of a.a derived from the experimentally estimated value of $L_c$ (Figure 3.8). This applies for both, SMFS experiments from discs compared to simulations with $\epsilon_{\text{MEMBR}} = 5.64\epsilon$, and SMFS experiments from the plasma membrane compared to simulations with $\epsilon_{\text{MEMBR}} = 10\epsilon$.

The number of unfolded a.a. in the simulation was determined by careful visual inspection of the generated trajectories. Some points (the two red points in panel (a) and the two blue points in panel (b)) can not be uniquely assigned to a single experimental peak, since the single experimental peak can be associated with two theoretical peak values. A possible explanation for this is that in real experimental data the initial parts of the F-x curves are highly dominated by non-specific interactions.

These results may explain the differences in the F-x curves obtained when rhodopsin is unfolded from the discs and from the plasma membrane, whose significant hydrophobicity is caused by its higher cholesterol content. Our model suggests that the different F-x curves are only due to the different values of the hydrophobicity. Given that the model is built on the basis of essential inter- and intramolecular interactions only, its success in reproducing the experimental data supports the hypothesis we made in the beginning of this chapter, namely that the different cholesterol concentration in the discs and in the plasma membrane affects the mechanical stability of rhodopsin, and therefore the F-x curves.
Figure 3.8: (a, b): Correlation between the experimental values of $n$, the number of unfolded amino acids, as deduced by the values of $L_c$ and the theoretical values of $n$, observed in the pulling trajectories. The panel (a) corresponds to the simulation with $\epsilon_{MEMBR} = 5.64\epsilon$ (Figure 3.7c), compared with the experimental values for the discs [12]. The panel (b) corresponds to the simulation with $\epsilon_{MEMBR} = 10\epsilon$ (Figure 3.7g), compared with the experimental values for the plasma membrane [12]. The color map in panel a is the same as the one used in Figure 3.7c. The color map in panel b is the same as the one used in Figure 3.7g. The points with the same coloring (two red points in panel (a) and two blue points in panel (b)) correspond to ambiguous cases, in which, for instance, a single experimental peak may be associated with two theoretical peaks (and vice versa).

3.3.2 Rhodopsin’s activation is affected by the membrane hydrophobicity.

The transition of rhodopsin to its active state, metarhodopsin II [88, 89] requires a series of conformational changes. The effect of membrane composition, and in particular cholesterol concentration, on similar conformational changes has been studied in detail, primarily for G-protein coupled receptors [90, 91]. By using all-atom MD simulations we argue that cholesterol also alters the flexibility of rhodopsin and the related equilibrium between its active and inactive forms. If the membrane is more hydrophobic, the configuration with more exposed hydrophobic residues will be favored reducing also the extent of conformational fluctuations and the ability of rhodopsin to become active.

To test this hypothesis, we measured the angle $\alpha$ between the transmembrane helices D and E (Figure 3.9a), thought to be a key descriptor for the G-protein activation [92]. We also estimated the solvent accessible surface area (SASA) of the hydrophobic transmembrane residues, $A_{HPHOB}$, and the SASA of the hydrophilic transmembrane residues, $A_{HPHIL}$, using the program Free SASA [93]. The probability distribution of $\alpha$ (the red line in Figure 3.9b) is characterized by a broad maximum at about 2.72 rad. Remarkably, $A_{HPHOB}$ and $A_{HPHIL}$ have a maximum and a minimum - respectively - approximately in correspondence with the most probable value of $\alpha$ (Figure 3.9c). This implies that in the most likely conformations rhodopsin exposes to the membrane the largest number of hydrophobic residues allowed by its fold and the minimum possible number of hydrophilic residues.
A membrane with a higher hydrophobicity will hamper large fluctuations, therefore favoring the native conformation.

To quantify this effect, we exploited the estimate from ref. [94] of the partition coefficient of a triethylamine, a hydrophobic molecule, from a pure membrane to a membrane containing various concentrations of cholesterol. Using this estimate, we applied Eq. 3.6 and Eq. 3.7 to determine the probability distribution of the values of $\alpha$ induced by cholesterol in the membrane (the blue dotted line in Figure 3.9b). High cholesterol content shifts the distribution to the right and makes it slightly narrower, favoring the conformations with more hydrophobic residues exposed towards the membrane, and hindering fluctuations of $\alpha$.

We noted that larger distortions of the probability distribution would imply a change in the tertiary packing of the helices. This would be inconsistent with the experimental observations, and indeed we did not observe it in our simulations. A change of approximately 0.18 radians in $\alpha$ corresponds to a change of approximately 9.5 Å in the distance between $C_\alpha$ 159 and 222. Therefore, a tiny change in the distribution of the angle $\alpha$, leads to a significant difference in the structural ensemble.

Based on the same model, we estimate the change in the relative population of the metarhodopsin and rhodopsin as a function of the cholesterol concentration. We provide only a rough estimate, based on the value of the SASA measured in the crystal structures of metarhodopsin II (PDB:
CHAPTER 3. MOLECULAR MODELING OF RHODOPSIN.

3PXO) and rhodopsin (PDB: 1U19). The probability distribution as a function of the cholesterol concentration is:

\[ P(c) = e^{-\frac{V_0}{k_B T} e^{\frac{\gamma(c)A}{k_B T}}} = e^{\frac{\gamma(c)A}{k_B T}} P(0) \]  

(3.8)

where \( P(0) \) corresponds to the probability distribution in a cholesterol-free membrane.

Denoting by \( \frac{P_{\text{METARHOD}}}{P_{\text{RHOD}}}(c) \) the ratio between the population of metarhodopsin and rhodopsin in a membrane characterized by a concentration \( c \) of cholesterol and applying Eq.3.8, we obtain

\[ \frac{P_{\text{METARHOD}}}{P_{\text{RHOD}}}(c) = \exp\left[\frac{\gamma(c)}{k_B T}(A_{\text{METARHOD}} - A_{\text{RHOD}})\right] \frac{P_{\text{METARHOD}}(0)}{P_{\text{RHOD}}(0)} \]  

(3.9)

where \( A_{\text{METARHOD}} \) and \( A_{\text{RHOD}} \) are the SASA of the hydrophobic residues in the two conformations and \( \gamma(c) \) is the partition coefficient per unit area of a hydrophobic molecule between a cholesterol-free membrane and a membrane characterized by a cholesterol concentration \( c \). \( \gamma(c) \) is estimated from a work by Zocher et al, [94] using the value of triethylamine (TEA) in membranes composed of DOPC with cholesterol concentrations 0, 20, 30, 40 and 50 mol %. To compute the values of the partition coefficient we used the relations:

\[ P_{\text{TEA}}(c) = \exp\left[ \frac{A_{\text{TEA}}}{k_B T} \gamma(c) \right] P_{\text{TEA}}(0) \]  

(3.10)

\[ \gamma(c) = \frac{(\ln P_{\text{TEA}}(c) - \ln P_{\text{TEA}}(0))}{A_{\text{TEA}}} k_B T \]  

(3.11)

where \( P_{\text{TEA}}(c) \) corresponds to the probability distribution of TEA as a function of the cholesterol concentration, \( c \) (values extracted from Figure 4B in ref. [94]), \( \gamma(c) \) is the partition coefficient between the two lipid environments and \( A_{\text{TEA}} \) the SASA of a single triethylamine molecule. The values for \( \gamma \) are presented in Table 3.2. In this definition \( \gamma \) estimates the surface tension per unit area of a hydrophobic moiety. The results are largely insensitive to the choice of the molecule used to compute the surface tension per unit area, if this is hydrophobic. For example, for pyridine in membranes composed of DOPC with cholesterol concentration 40 mol % we have \( \frac{\gamma_{\text{PYR}}}{A_{\text{PYR}}} = 0.0018 \), a value similar to the one observed for TEA, \( \frac{\gamma_{\text{TEA}}}{A_{\text{TEA}}} = 0.0014 \) (values extracted from Figure 6C and D in ref. [94]).

Due to the hydrophobicity effect, the cholesterol concentration changes the relative ratio between metarhodopsin II and rhodopsin (Figure 3.9d) and therefore the rate of rhodopsin activation. At a concentration of 50% the ratio between metarhodopsin II and rhodopsin is approximately 0.17 of the ratio observed in the absence of cholesterol, but even at a lower concentration the ratio of the two populations is affected.
Table 3.2: Partition coefficient values for triethylamine (TEA) estimated at different cholesterol concentrations in DOPC lipid membrane with cholesterol concentration 0 mol % as reference value.

<table>
<thead>
<tr>
<th>Cholesterol concentration, c [mol %]</th>
<th>Partition coefficient, $\gamma$ [kcal $\cdot$ mol$^{-1}$Å$^{-2}$]</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>20</td>
<td>7.061 $\cdot$ 10$^{-4}$</td>
</tr>
<tr>
<td>30</td>
<td>2.889 $\cdot$ 10$^{-4}$</td>
</tr>
<tr>
<td>40</td>
<td>1.027 $\cdot$ 10$^{-3}$</td>
</tr>
<tr>
<td>50</td>
<td>8.987 $\cdot$ 10$^{-3}$</td>
</tr>
</tbody>
</table>

3.4 Conclusions.

AFM-SMFS experiments allow the studies of membrane proteins in their natural lipid environment - the cell membrane. Cell membranes have different lipid composition and many studies suggest that the lipid composition, in particular the cholesterol content, alters the stability and modulates the activity of integral membrane proteins[95, 23]. In this chapter we presented a simple coarse-grained model of the rhodopsin-membrane system, which can be used to perform "cheap" molecular dynamics (MD) simulations of the pulling AFM experiments. This model, albeit simple, is able to reproduce the differences between the unfolding pattern of rhodopsin in the discs and in the plasma membrane of the rod outer segment (OS) as it was observed experimentally [12].

Our molecular model is an extension of the model developed by Cieplak et al. and used in a MD study of the unfolding of bacteriorhodopsin (bR) [28]. The main drawback in Cieplak’s model is that the lipid bilayer used to represent the membrane is kept frozen during the pulling of the protein. The space left by the extracted polypeptide chain remains empty, leaving a hole in the bilayer. Such behaviour is quite unrealistic and possibly altering the stability of the intermediate states of the unfolding. In order to circumvent this problem, we modeled the membrane with an extra potential energy term, $V^{MEMBR}$, acting only in the z-direction. $V^{MEMBR}$ favors the native protein-lipid interactions and in this manner mimics the hydrophobic effect of the membrane. The only free parameter in this approach is the strength of the membrane potential: $\epsilon^{MEMBR}$. The value of $\epsilon^{MEMBR}$ can be changed in order to model different membrane hydrophobicities. Larger values of $\epsilon^{MEMBR}$ correspond to more hydrophobic membranes like cholesterol-rich membranes.

We performed MD simulations using different values of $\epsilon^{MEMBR}$, which according to our model correspond to pulling experiments in membranes with different hydrophobicity. When we increase the $\epsilon^{MEMBR}$ value, more force peaks appear in the simulated curves and the force required to unfold the protein becomes larger. This is consistent with the presence of stronger interactions between the protein and the membrane, which increases the mechanical stability of the protein. Furthermore, we obtained two groups of theoretical curves in very good agreement with the experimental curves from the plasma membrane and the discs by simply using two different $\epsilon^{MEMBR}$ values. These results
suggest that the differences in the F-x curves of rhodopsin in the discs and in the plasma membrane are due to the higher hydrophobicity of the plasma membrane, induced by its higher cholesterol concentration.

Our findings were supported by additional experiments performed in the plasma membrane of the rod OS treated with cyclodextrin [12]. Cyclodextrin is an agent that binds cholesterol and lowers its concentration. As a result, in many traces the number of force peaks decreased from 7 to 5, which is the number of force peaks present in the traces from the discs.

In this chapter, we addressed also another problem concerned with the inactivation of rhodopsin in the plasma membrane. We argue that the cholesterol-rich plasma membrane reduces the flexibility of rhodopsin and alters its ability to change conformation. This is in agreement with the larger mechanical stability of rhodopsin observed experimentally in the plasma membrane and supported by our CG simulations. If this is true, rhodopsin can not switch to its active conformation metarhodopsin II and the phototransduction cascade can not be initialized. In order to test this hypothesis, we performed all-atom MD simulations of the inactive conformation of rhodopsin embedded in a DPPC bilayer without cholesterol. We used a statistical mechanics apparatus in combination with theoretical data on partition coefficients in membranes with different cholesterol concentrations, to evaluate the cholesterol effects on rhodopsin’s flexibility. Our results suggest that the higher cholesterol content might shift the equilibrium ratio between rhodopsin and metarhodopsin towards the inactive state, which may explain the inactivation of rhodopsin in the plasma membrane.

The coarse-grained MD approach described in this chapter is not restricted to rhodopsin. It can be applied to any other membrane protein of known structure. However, our results indicate that the F-x curves of membrane proteins depend on many factors, including not only the lipid composition of the membrane but possibly its protein composition too. We saw that by simply changing a single parameter in the model, $\epsilon_{MEMBR}$, the features of the F-x curves change tremendously. This makes the identification of membrane proteins based on the unfolding pattern observed in their F-x curves difficult, because clearly this pattern is not uniquely defined like for soluble proteins. Therefore, even more than for soluble proteins, for membrane proteins experimental data need to be provided in order to verify the results coming from the model.

In the next chapter, we describe an automatic tool we developed for the analysis of AFM-SMFS data coming from experiments in native cell membranes. With this procedure we are able to find clusters of traces sharing the same features, possibly describing the unfolding of same membrane protein. A key missing ingredient, which should be addressed in the future, is that these clusters should be assigned to a specific membrane proteins. A possible manner of achieving this goal is using our MD tool to simulate the F-x curves for many proteins of known structure with different $\epsilon_{MEMBR}$ values and look for a match with the experimental curves included in the cluster.
Chapter 4

Automatic classification of AFM traces from native membranes.

Membrane proteins perform a variety of functions in the cell. They are key mediators in the processes of signaling, cell-cell recognition and transport of ions and molecules across the membrane. Nearly 30% of all proteins in eukaryotic cells are membrane proteins. They are the targets of more than 50% of modern medicinal drugs and many diseases are found to be related to membrane proteins misfolding [1]. However, studying membrane proteins is non-trivial, mainly because they are difficult to purify and crystallize. Studying membrane proteins in their native environment is even more difficult due to the heterogeneity of the membrane and its specific chemical properties. Moreover, most of the modern experimental techniques have been designed to study soluble proteins.

A way out, as we will show in this chapter, seems to be offered by AFM-based SMFS. This method allows localizing and quantifying key inter- and intramolecular interactions, estimating the effect of environmental factors on the unfolding process, probing structural and conformational properties of membrane proteins. The interpretation of the obtained experimental data reveals important insights in the membrane proteins structure-function relations in the presence of the membrane. Recent developments in the AFM-SMFS experimental approach enabled the accumulation of hundreds of thousands of traces in a reasonable time [16, 17]. Strikingly, these experiments can nowadays be performed in native cell membranes under physiological conditions: in section 1.1 we will describe an experimental technique which allows reaching this goal.

The availability of this huge amount of data calls for the development of specific theoretical tools allowing analysis and interpretation. However, these data require careful preprocessing since most of the F-x curves do not contain meaningful unfolding events. It has been estimated that membrane proteins get completely unfolded by SMFS in < 1% of the cases [21].

To the best of our knowledge, there is no automatic procedure which allows the classification of
the SMFS data in an unsupervised manner, especially in the case of data coming from experiments performed in native cell membranes. Indeed, there are several automatic procedures for AFM-SMFS data analysis reported in the literature [19, 20, 68] but all of them require an approximate knowledge on the sample composition. The reason is that in these methods the preprocessing of the raw data is often based on the selection of force-displacement curves with length values in the range corresponding to the fully stretched protein under investigation. In simple terms, if you know that in your experiments you are unfolding bacteriorhodopsin and your bacteriorhodopsin contains 248 a.a., given that the peptide bond length on average is 0.4 nm, in your subsequent analysis you are going to include only traces with contour length around 100 nm. In the case of experiments in which the protein sample composition is well known, this approach can be extremely useful. But in the case of pulling experiments performed in the native cell membranes, where there are plenty of membrane proteins, some of which unknown, this approach has clear limitations.

In this chapter we describe a procedure for the automatic classification and analysis of highly heterogeneous SMFS datasets in which the protein sample composition is unknown, just like in native cell membranes. Accordingly, our method does not include a filtering step based on the length of the fully stretched protein under investigation. Instead, we developed a filtering procedure based on the quality of each trace evaluated by a carefully defined quality score. The next step aims at detecting different unfolding patterns in the data, arising from the unfolding of different proteins in the membrane. For this purpose, we use a modified version of the density-peak clustering [29], in which a key ingredient is defining a suitable distance between two traces. The distance measure we propose, combines the dynamic programming alignment score, introduced in ref. [20] and the traces’ quality score, in such a way that if two traces are well-aligned but are low quality their distance is large, while if two well-aligned traces are high quality, their distance is small.

The procedure we developed is fully-automatic and unsupervised. In addition, it allows the processing of large amount of data in a reasonable computational time.

This chapter is organized as follows. We begin with a description of a recently developed experimental technique which allows the performance of AFM-SMFS directly in the native membranes of different types of cells and motivates the development of our procedure. Next, we provide a detailed description of our algorithm and we discuss briefly its relation with other approaches reported in the literature. Then, we introduce the data sets used to benchmark and test the performance of our method. Finally, we report the results and end with conclusions.

4.1 High throughput AFM in native membranes.

For many years, the imaging of native cell membranes remained a challenge. Significant progress was made in the 70s, when experimentalists started developing the so-called 'cell unroofing' techniques. 'Cell unroofing' straightforward means 'breaking the cell'. First, the cells get attached to
a surface through a gluing substance (polylysine or Alcian blue). Then, they have to be broken so that pieces of the membrane are extracted. There are three main strategies to accomplish this: (1) break the cells with a strong lateral flux of medium, leaving pieces of membrane attached to the substrate [96]; (2) squash the cells between two coverslips, freeze them and fracture them by separation of the coverslips [97]; (3) use sonic waves to break the cells leaving pieces of the membrane on the substrate [98]. The scanning electron microscope is then used to image the membrane patches.

These strategies are difficult to implement. The sonic waves, for example, affect the entire cell culture in an unconstrained manner. Moreover, the AFM might be more appropriate for direct imaging of cell membranes since it can operate in buffer solutions under physiological conditions.

Recently, a new methodology designed by Galvanetto et al. [18] made possible the investigation of the native membranes of a variety of single cells in a simplified and handy manner. In this work a new ‘cell unroofing’ strategy is developed in which a single cell is ruptured with a sharp triangular piece of glass, called the arrow. A piece of the membrane remains on the arrow and gets imaged with the AFM. A characteristic feature of this approach is that the AFM is used also in the sample preparation.

The sample preparation requires three main ingredients: the cell culture coverslip, the cell culture holder and the arrow. The cell culture coverslip is simply a glass round coverslip (12 mm in diameter, 200 $\mu m$ thick), plasma cleaned and coated with poly-D-lysine. The coating step enhances the adhesion of the cells and the substrate, which is necessary because otherwise the entire cells might get adsorbed on the arrow. The cell culture holder acts as a motor with micrometer precision used to bring in contact the cell culture coverslip and the arrow. The motor role is performed by the AFM. The arrow was prepared by breaking a round coverslip (24 mm in diameter, 200 $\mu m$ thick) in four with hands. It was also immersed in poly-D-lysine but for a shorter time period.

The AFM was mounted on an inverted optical microscope. The cell culture coverslip was attached to the AFM holder. The coverslip was moved down with the AFM, towards the arrow. A single cell was brought in contact with the apex of the arrow and squeezed for $\sim 3$ minutes. Afterwards, the cell culture coverslip was rapidly moved away from the arrow. As a result, a piece of the membrane remained on the arrow as shown in Figure 4.1. The AFM images were obtained with a NanoWizard 3 JPK system in the intermittent contact mode. The cantilever spring constant was 0.08 N/m.

The method was tested on five different cell types, among which human brain cancer cells and primary hippocampal neurons of rats. The success of the experiment depends on the adhesion of the cell types to the substrate. If this effect is not strong enough, the entire cell gets adsorbed on the arrow.

Once the membrane patches were isolated, their topology and mechanics were examined. Furthermore, using the same technique, SMFS experiments were performed and a huge amount of F-x curves obtained.
The next challenge is to detect the characteristic unfolding fingerprints in such large heterogeneous data sets and to relate them to the unfolding of certain membrane proteins.

4.2 The algorithm.

The main goal of our algorithm is to automatically analyze and classify large data sets coming from SMFS experiments like the ones described in the previous section. These membranes contain many different proteins. Some of these proteins might have been already crystallized and their three-dimensional structure might be known. Others might be completely unknown, leaving us with a F-x fingerprint asking for further identification. In general, a good computational procedure should be automatic, efficient and should require the least possible manual intervention.

The algorithm we developed can be divided in the blocks depicted in Figure 4.2. Initially, each F-x curve goes through a series of operations, summarized in the "Cutting & filtering block". In this block, all parts of the original trace that are physically irrelevant are removed, clearing the space to meaningful unfolding events only. Traces that are very short (below a user-defined length threshold) or completely negative, etc. are discarded (see subsection 4.2.1).

Next, two scores of equal importance are computed: the alignment score and the quality score. The alignment score estimates the similarity between two traces based on dynamic programming alignment [20] (see subsection 4.2.3). The quality score determines the consistency of the experimental data in each trace with the worm-like chain (WLC) model, proved to provide a proper quantitative description of the unfolding events (see subsection 4.2.2).

The distance used for classification combines the alignment score and the quality score. The
Figure 4.2: Block diagram of the algorithm developed by us.
power of this measure is that it is able to distinguish well-aligned good traces from well-aligned bad traces, where "good" or "bad" is quantified by the quality score. Then, a modified version of the density-peak clustering \cite{29} method is used to group the different meaningful F-x patterns into clusters (see subsection 4.2.5). The obtained clusters are subject to further interpretation in order to be assigned to the unfolding of a specific protein.

With our procedure a large amount of raw experimental data is processed in a reasonable time. It takes $\sim 1$ hour and 30 minutes to run the program on a set containing $10^5$ traces on a workstation with 16 CPUs.

4.2.1 Cutting and filtering the traces.

The first step in our procedure is data preprocessing. In subsection 2.0.6, we have explained why this step is necessary and so important. In summary, not all F-x curves contain successful unfolding events; most of the force spectra contain mainly noise or the force pattern, if any, can not be interpreted.

When a trace is read by our program, it is stored in two vectors: one containing the extension values and a second one containing the force values. In this vectors only points corresponding to successively increasing extension values are kept. All F-x curves with abnormally large extension and force values are discarded. The threshold value is 5,000 nm for the extension and 5,000 pN for the force. The presence of traces with out-of-range force and extension values is attributed to technical defects in the software used to process the raw experimental AFM data. Related to technicalities is also the presence of traces containing exclusively negative force values which are also discarded.

In general, each F-x curve starts with a region of highly negative forces coming from the upwards bending of the cantilever in the very beginning of the retraction cycle (Figure 2.9). The actual unfolding events appear in the positive force range. In order to exclude this initial negative part, for each trace, we find the first point at extension larger than 0 nm, followed by 20 consecutive points with positive force values. We call this point the starting point; the point in which the positive contact part begins. If we are not able to find at least 20 consecutive positive force values, the trace is discarded.

Next, we remove the non-contact part, the so-called tail of each F-x curve. This step requires the estimation of the standard deviation of the noise, $\sigma_{\text{NOISE}}$, in advance. Since its value depends on the spring constant of the cantilever, $\sigma_{\text{NOISE}}$ varies in different experiments. To estimate its value, we selected manually the eligible tails of 10 traces and computed the standard deviations of the force on these traces. This determines the value for $\sigma_{\text{NOISE}}$. Then we perform a linear fit to the last part of each trace. The core of the procedure is depicted in Figure 4.3. We start from 8 nm extending the fitting range towards the trace origin by approximately 2 nm$^1$ in a stepwise manner and we compute the standard deviation from the fit. When it exceeds $3\sigma_{\text{NOISE}}$ we stop. We assume

\footnotetext{1}{The exact spacing along the x axis depends on the sampling rate (the frequency in Hz) and the pulling speed.}
that this is an indication that the last force peak has been reached and the non-contact part has ended. The position of the detachment peak provides the trace length.

Figure 4.3: A schematical representation of the removal procedure of the non-contact part of a F-x curve. 

a. A drawing of a F-x curve. The last 8 nm in the non-contact part of the tail are highlighted in yellow. The following extension of 2 nm is highlighted in pink. b. The green line represents the linear fit to the last 8 nm (highlighted in yellow) of the tail plus three extensions of 2 nm each (highlighted in pink). The standard deviation of the noise is denoted as $\sigma$ and the threshold used to cut the traces is $3\sigma$. The highlighted blue area shows the upcoming 2 nm window in which the standard deviation of the point $i$, $\sigma_i$ is compared to the $3\sigma$ threshold. $\sigma_i$ is smaller than $3\sigma$ and the procedure continues. c. The standard deviation of the point $j$, $\sigma_j$ is larger than $3\sigma$ and the procedure stops. The point $j$ is the last point included in the trace. All points coming after $j$ are removed.
The non-contact parts of a large fraction of traces show unusual slopes and curvatures, which makes the traces ambiguous. To remove these traces, we apply an additional filter. By detecting the position of the last force peak, we obtain the total length of the non-contact part. This allows us to compute the standard deviation of the tail, $\sigma_{tail}$, from a horizontal zero-force line. If $\sigma_{tail} > 2\sigma_{NOISE}$ the trace gets discarded.

Another filter acts on short traces. If after the non-contact part is removed the trace is shorter than 50 nm the trace gets discarded.

### 4.2.2 Computing the consistency score.

The second step in our approach is the computation of the worm-like chain (WLC) consistency score. The consistency score quantifies how well the experimental data satisfy the WLC model predictions. The better the fit to the WLC model, the higher the total score for that trace. We compute the score by the following steps:

1. **Compute $L_c$ values.** We first perform a variable transformation from extension $x$ to contour length $L_c$ with the worm-like chain (WLC) model [44]. For this purpose, the WLC equation (Eq. 2.1) was converted into a third order polynomial following Bosshart et al. [21]. For each point in the F-x curve we compute $L_c$ value by solving the third order polynomial:

$$4\lambda^3 + \omega\lambda^2 - 1 = 0$$

where $\lambda = 1 - \frac{x}{L_c}$ and $\omega = \frac{4F}{\alpha}$ with $\alpha = \frac{k_BT}{l_p}$. The polynomial has three solutions: one real and two complex. Only one of them is physically relevant, in particular:

$$\lambda = -\frac{\omega}{12} \left(1 + \frac{\omega}{\beta\gamma}\right) - \frac{\beta\gamma}{12}$$

with

$$\beta = -\frac{1}{2} + i\frac{3^{1/2}}{2}$$

and

$$\gamma = (216 - \omega^3 + 12(324 - 3\omega^3)^{1/2})^{1/3}$$

By using $\lambda = 1 - \frac{x}{L_c}$ Eq. 4.2 gives:

$$L_c = x \times Re \left[ \left( \frac{\omega}{12} \left(1 + \frac{\omega}{\beta\gamma}\right) + \frac{\beta\gamma}{12} + 1 \right)^{-1} \right]$$

where $Re$ means taking the real part of the expression in the brackets. The persistence length we used was fixed to 0.4 nm. Each point of a trace is characterized by a value of $F$ and a value of $x$. The value of $L_c$ is computed by solving Eq. 4.5. The $L_c$ transformation is applied in the
force range from 30 to 500 pN due to the limitations of the WLC model (see subsection 2.0.4).

2. **Compute the histogram.**

We then estimate the histogram of the calculated $L_c$ values. The $L_c$ histogram of meaningful traces is characterized by the presence of a few maxima, separated by relatively deep minima. This structure comes from the unfolding of individual protein domains like the $\alpha$-helices present in many membrane proteins. A critical parameter for our algorithm is the bin width used for computing the histogram. If the bin width is too small, the histogram is noisy and the peaks corresponding to the unfolding of each separate domain are split. On the opposite, if the bin width is too large, meaningful peaks get merged. We have chosen a bin size of 8 nm, a value corresponding to approximately 20 a.a., which is close to the typical length of a single transmembrane helix in membrane proteins [34]. The choice of this bin size is benchmarked in subsection 4.5.2.

3. **Find minima and maxima.** In order to define the score we then find all the maxima and minima in the $L_c$ histogram. We denote by $n_{min}$ the total number of minima and by $n_{max}$, the total number of maxima. A trace is discarded if the last point in its histogram is a maximum, if it has only one maximum or if the number of maxima is larger than 10.

4. **Compute the score.** Now that all maxima have been detected, we compute the consistency score $W$ of each maximum. The consistency score is a peak quality factor. Ideally, we assume that a high quality peak has its two surrounding minima falling under $\frac{1}{2}$ of the peak height and in this case $W = 1.0$. If only one of the two minima satisfies this condition $W = 0.5$, otherwise $W = 0.0$. The closer to 1.0 the value of $W$, the deeper the maximum and the better the WLC fit. We implemented these requirements in the following functional form:

$$W = e^{-2.0 \left( \frac{p_{\text{min,left}}}{p_{\text{max}}} + \frac{p_{\text{min,right}}}{p_{\text{max}}} \right)^2}$$ (4.6)

with $P_{\text{min,left}}$ and $P_{\text{min,right}}$ the probability densities of the left and the right minima surrounding the maximum, whose probability density is $P_{\text{max}}$. If, for example, $P_{\text{min,left}} = 1$, $P_{\text{min,right}} = 2$ and $P_{\text{max}} = 16$, $W = 0.98$. If we change $P_{\text{min,left}} = 4$ and $P_{\text{min,right}} = 12$, $W = 0.60$ and for $P_{\text{min,left}} = 13$ and $P_{\text{min,right}} = 14$, $W = 0.24$.

In Figure 4.4 we provide a few examples of F-x curves, their $L_c$ histograms and the $W$-score of some of the peaks. The trace in panel a is a high quality trace and this is reflected in its histogram (panel b), which contains well-defined peaks with high W-scores. The trace in panel c is a good trace but it has also a peak with $W$-score 0.5 (panel d), which is a medium quality peak according to our definition. The third trace in panel e contains a low quality peak with $W$-score 0.27 and a high score peak with $W$-score 0.92 (panel f).
5. **Score assignment.** Once we have computed the WLC consistency score $W$ for each peak we assign a $w_i$ score to every point in each trace. A score is assigned to a point in two steps:

(a) We assign the peak’s score to all points in the $L_c$ histogram with $L_c$ values between the peak’s surrounding minima $L_c$ values. In this step we are excluding the points which are not accurately modeled by the WLC model namely, the points with forces below 30 pN.

(b) If a point has a force smaller than 30 pN we assign to it the same score $w$ of the first successive point whose force is larger than 30 pN. We apply this criterion only for points that are within 75 nm from the last point assigned to the peak (Figure 4.5). We selected this value by visual inspection of the traces, estimating the maximum widths of force peaks.
Figure 4.5: A scheme representing the second step in the score assignation process. At this step the peak score, depicted in pink is assigned also to points belonging to the peak but with forces below 30 pN and at distance 75 nm from the last peak point (the yellow area). The 30 pN threshold is set by the WLC model limitations. The 75 nm range was chosen based on visual inspection of the traces, looking at the maximum extension of relevant force peaks.

The threshold distance of 75 nm is introduced in order to avoid assigning too many points to a peak. For example, if there is a high score peak preceded by a long flat region, the points belonging to that part of the trace will be wrongly assigned to the peak’s score. When we apply the threshold they get zero score. Zero score is assigned also to points that do not satisfy the criteria pointed in (a) and (b), to points belonging to a peak with probability less than 1 % or to the points belonging to a peak corresponding to $L_c = 0$.

6. **Compute the global score.** The global WLC consistency score is computed as the sum of the $w_i$ scores for each trace,

$$S_w = \sum_{i=1}^{N} w_i$$

(4.7)

where $N$ is the total number of points in the F-x curve. The global score $S_w$ is a measure of the quality of that trace.

7. **Discard the traces with low global score.**

We use the ratio between the global score and the trace length to select high quality traces for subsequent analysis. If this ratio is smaller than 0.5, the trace is discarded. This is the same as saying: if more than half of the trace is inconsistent with the WLC model, it is a bad trace.
and we are not interested in analyzing it. On the contrary, if more than half of the trace is consistent with the WLC model, it is possibly a meaningful trace.

In Figure 4.6 we demonstrate this idea with an example. The data used in the figure is from a data set which contains a portion of manually selected traces describing the unfolding of the membrane protein, CNGA1 channel, and a larger portion of unknown traces which might be good or bad. A detailed description of the data set is given in section 4.4.

In Figure 4.6a we show the relation between the trace length and the global WLC consistency score. The traces with score-length ratio higher than 0.5 are shown as gray points and the selected CNGA1 traces are among them, depicted in black. The traces with score-length ratio lower than 0.5 are shown as red points. One can notice that for most of the CNG traces, the score is more or less equal to their length, which means that most of them satisfy well the WLC model and these traces are good according to our model (black points).

Figure 4.6a also shows that the dataset contains traces both much shorter and much longer than the CNG traces with high global scores with respect to their lengths (depicted with blue symbols). In Figure 4.6b, we plotted two of the shorter ones and two of the longer ones to illustrate that the combination of trace length and score can be used to detect meaningful traces.

The red points in Figure 4.6a have global scores much smaller than their length indicating poor agreement of the data with the WLC model.

4.2.3 The distance between two traces.

The final goal of our procedure is finding in an automatic and unsupervised manner meaningful F-x curves bearing a specific unfolding pattern and to group them into clusters based on their similarity to each other. To reach this goal a key ingredient is the distance between traces. The distance used in this work is based on a combination of the trace distance obtained by dynamic programming as in ref. [20] and of the quality score defined in the previous paragraph.

Dynamic programming finds the best match between the traces by allowing insertions and deletions. Given two traces, $a$ and $b$, denote by $S_D(i, j)$ the score between trace $a$ up to position $i$ and trace $b$ up to position $j$. In dynamic programming $S_D(i, j)$ is defined recursively starting from the beginning of the traces,

$$S_D(i, j) = \max(S_D(i - 1, j) - \mu, S_D(i, j - 1) - \mu, S_D(i - 1, j - 1) + M(i, j)) 
\quad (4.8)$$

where $\mu$ is the gap penalty and $M(i, j)$ is the match/mismatch score.
Figure 4.6: a. Trace length vs. global score in the CNGA1+noise data set. The gray points represent all traces in the data set after the score-length-based filtering, the black points represent the CNGA1 traces and the red points represent the traces which were excluded due to low score-length ratio. The blue symbols match the traces plotted in panel b. In panel b we show four high-scoring traces in data set I. Two of them are shorter than the CNGA1 channel and two of them are longer than the CNGA1 channel.
CHAPTER 4. AUTOMATIC CLASSIFICATION OF SMFS DATA FROM MEMBRANES.

The match/mismatch score is defined as follows:

\[
M(i, j) = \begin{cases} 
1 - \frac{|F_a(i) - F_b(j)|}{\Delta F_{\text{max}}} & \text{if } |F_a(i) - F_b(j)| < 2\sigma_{\text{NOISE}} \\
\frac{|F_a(i) - F_b(j)|}{\Delta F_{\text{max}}} & \text{otherwise}
\end{cases}
\] (4.9)

where \( F_a(i) \) and \( F_b(j) \) are the forces in points \( i \) and \( j \) in traces \( a \) and \( b \), \( \Delta F_{\text{max}} = \frac{F_{\text{max},a} + F_{\text{max},b}}{2} \) with \( F_{\text{max},a} \) the maximum force value in trace \( a \) and \( F_{\text{max},b} \) the maximum force value in trace \( b \) [20]. The gap penalty \( \mu \) is set to a value of 0.002 for force values in the first 10 nm of the trace and to 0.8 in the rest of the trace in agreement with ref. [20]. The higher the alignment score \( S_D(i, j) \) the bigger the similarity between the two force curves. All possible alignment scores \( S_D(i, j) \) make up the dynamic programming matrix. The optimal alignment is then found through traceback [99]. In the final cell of the matrix, \( S_D(N_a, N_b) \), the score of the best global alignment of \( a \) and \( b \) is stored. \( N_a \) is the length of the preprocessed trace \( a \) and \( \text{max}(N_a, N_b) \) is the length of the preprocessed trace \( b \). The optimal alignment is build in reverse starting from the final cell following the path through which the maximum alignment score is obtained. The optimal alignment length is denoted as \( N_D \) and the optimal score is denoted as \( S_D(i_k, j_k) \) for \( k = 1, \ldots, N_D \).

Now that the alignment score has been defined we can introduce the distance used in this work, \( d_{ab} \). The distance between two traces \( a \) and \( b \) is defined by combining the alignment score, \( S_D \), with the WLC consistency score, \( w \):

\[
d_{ab} = 1 - \frac{\sum_{k=1}^{N_D} S_D(i_k, j_k) \min(w_{i_k}^a, w_{j_k}^b)}{N_{\text{max}}}
\] (4.10)

where \( S_D(i_k, j_k) \) is the local optimal alignment score, \( N_D \) is the optimal alignment length, \( N_{\text{max}} = \text{max}(N_a, N_b) \); \( w_{i_k}^a \) and \( w_{j_k}^b \) are the WLC consistency scores in the aligned point \( k \) in traces \( a \) and \( b \). Negative alignment scores are considered zero in the summation.

This definition of the distance is a generalization of the one in ref. [20] and relies on two main contributions: the alignment score and the WLC consistency score. \( S_D \) quantifies how similar two traces are, while \( \min(w_{i_k}^a, w_{j_k}^b) \) accounts for their quality. In this way we are able to distinguish between well aligned good and bad traces. If two traces are both well aligned and satisfy well the WLC model, the distance will be small. If two traces are well aligned but their \( w \) score is low, their distance will be larger.

Our distance is approximately a metric: indeed it is non-negative, symmetric and it nearly satisfies the triangular inequality. The fraction of violations is \( \approx 0.0008 \) on a subset including 662 traces from data set I. In Figure 4.7 we show the probability distribution of \( q \), where \( q \) is equal to \( d_{13} + d_{23} - d_{12} \). The plot is showing that the probability of observing negative \( q \) values, which corresponds to violations of the triangular inequality is almost 0. We note however that by definition, the distance between one object and itself is not necessarily zero, violating the identity of the indiscernible axiom. This is due to the insertion of the WLC score in the distance. If we get rid
CHAPTER 4. AUTOMATIC CLASSIFICATION OF SMFS DATA FROM MEMBRANES.

Figure 4.7: Probability distribution of the quantity \( q = d_{13} + d_{23} - d_{12} \) in semilogarithmic scale, where \( d \) is our distance. Negative values of \( q \) indicate violations of the triangular inequality.

of this score the distance will be consistent with the one used by Marsico et al. [20] and the identity of indiscernibles will hold. The problem is that the similarity between two traces will be entirely governed by their shapes. The additional contribution of the WLC consistency score to the distance makes it more convenient for clustering. We are not only aligning two geometrical objects but we also weight them by the quantity of physically relevant information they contain. In this way, if two F-x curves have similar shapes, but they do not satisfy well the WLC model, the distance between them will be large. The other way around, if two curves satisfy well the WLC model, but they have very different shapes, the distance between them will be also large.

4.2.4 Proxy distance for speeding up the calculation.

The distances computation is the most expensive and time-consuming part in this computational approach since it scales quadratically with the number of traces. In order to make the whole procedure more efficient we define the proxy distance. The proxy distance is very cheap to compute and is used to select a relatively small sample of good quality traces similar to a trace. The proxy distance between traces \( a \) and \( b \), \( \tilde{d}_{ab} \), is defined as follows:

\[
\tilde{d}_{ab} = 1 - \sum_{i=1}^{N_{min}} e^{-\frac{(F_a(i) - F_b(i))^2}{2\sigma^2}} \frac{\min(w^a_i, w^b_i)}{N_{max}}
\]  

(4.11)
where $N_{min}$ is the minimum length between the length of the preprocessed trace $a$, $N_a$, and the length of the preprocessed trace $b$, $N_b$, or $N_{min} = min(N_a, N_b)$; $F_a(i)$ and $F_b(i)$ are the force values in point $i$ in traces $a$ and $b$; $\min(w^a_i, w^b_i)$ is the minimum WLC consistency score in point $i$ between traces $a$ and $b$; $\sigma = 25$; and $N_{max}$ is simply the maximum length, $N_{max} = max(N_a, N_b)$.

The similarity of a pair of traces is accounted for by the similarities in the forces and the consistency of the data with the WLC model. The similarity of the forces is taken into account by the exponential function while the consistency with the model is expressed by $\min(w^a_i, w^b_i)$. If two traces are very similar in terms of their force values and they satisfy well the WLC model, their proxy distance will be small indicating their similarity. If the traces differ a lot in the forces and they do not satisfy well the WLC model, their proxy distance will be large. Last but not least, if two traces have similar force values but they do not satisfy well the WLC model, their proxy distance will be shifted towards larger values.

For each trace we sort the computed proxy distances, $\tilde{d}_{ab}$, in ascending order and then we compute the true similarity distance measure, $d_{ab}$, for the first $n_{true}$ traces only. $n_{true}$ is a user-defined parameter with default value 5,000. If the total number of traces remaining after filtering is smaller than 5,000, the $n_{true}$ value is adjusted to the total number of traces. This is equivalent to computing the similarity distances for all pairs.

The true similarity distance and the proxy distance are well correlated with Pearson correlation coefficient $\sim 0.84$.

### 4.2.5 Clustering.

Once we have computed all distances, we use a modified version of the density peak clustering algorithm [29] with the $k$-nearest neighbor ($k$-NN) density estimator. An advantage of the method is that it doesn’t require the preselection of the number of clusters. The main idea behind density peak clustering is that the highest density points are located at relatively large distances from each other. The algorithm is the following:

1. The first step is computing the densities. In the $k$-NN method the density around a point is estimated as the ratio between $k$ nearest neighbors and the occupied volume; therefore,

$$\rho_i = \frac{k}{\omega_d r_{k,i}^d} \quad (4.12)$$

where $d$ is the intrinsic dimensionality (ID) of the dataset [100], $\omega_d$ is the volume of the $d$-sphere with unitary radius and $r_{k,i}$ is the distance of point $i$ to its $k$-th nearest neighbor. We then compute the logarithm of the density, $\log \rho_i = -F_i$, identified as the free energy at point $i$ [101]. Notice that using a $k$-NN estimator, the rank of $F_i$ is not influenced by the value of $d$, which therefore we do not compute. Eq. 4.12 reduces to:
\[ \log \rho_i = c_1 + c_2 \log \frac{1}{r_{k,i}} \]  

(4.13)

where \(c_1\) and \(c_2\) are constants, whose exact values do not influence the results.

2. We find the cluster centers, according to two conditions: (1) they should have highest density among their \(k\) nearest neighbors, and (2) they should not be among the \(k\) nearest neighbors of other points with higher density [101].

3. Subsequently, all points that are not centers are assigned to the same cluster as the nearest point with higher density.

### 4.3 Relation with previous works.

The first step in our procedure is quite similar to the one used by Kuhn et al. and Marsico et al. [19, 20]. We detect the detachment peak in each F-x curve and then remove the non-contact part. We remove also the initial negative contact part of each trace. Afterwards this part is not a subject of any physically relevant interpretation. In other approaches [20, 21, 22] the filtering of traces is done based on the expected contour length of the fully-stretched protein under investigation, which requires knowledge of the sample composition and the protein structure. This is a key assumption which reduces the number of analyzed traces tremendously. In fact, using such strong filtering condition can be useful in analyzing data coming from SMFS experiments with the scope of analyzing the structure of a specific protein. On the other hand, the analysis of SMFS data from experiments in native biological membranes, hosting a variety of membrane proteins, some of them unknown or with unknown structure, calls for a procedure which is able to automatically and simultaneously detect the characteristic unfolding fingerprints of several different proteins. In simple terms, the proper filtering procedure should be able to distinguish between good and bad traces based on their quality not on their length. For this purpose we developed the global WLC consistency score, which is a measure for the quality of each trace with respect to the WLC model. The WLC model describes the expected behavior of biopolymer molecules in AFM-SMFS experiments. If the peaks in a trace fit well with the model they should correspond to meaningful unfolding events, which is exactly what we are interested in, independently of the number of peaks, their positions and the trace length.

Instead of filtering traces based on their length, we filter traces based on their quality. We compute the ratio between the global WLC consistency score of the trace and its length, and if it is larger than 0.5, the trace is included in the subsequent analysis. This allows the analysis of large heterogeneous data sets without previous knowledge on the protein composition.

The alignment procedure we are using is the one reported by Marsico et al. [20]. The dynamic programming alignment gives as output the alignment score, which measures how similar two traces are. The larger the alignment score, the higher the similarity between the traces. The distance
metric used by Marsico et al. [20] is simply one minus the final alignment score. This is pretty much consistent with the metric used by Kuhn et al. [19] which is defined in the same way but with an alignment score not obtained using dynamic programming. With this metric two traces similar to each other have small distance, while two traces very different from each other have large distance. A problem arises if two traces are similar to each other but they are both of low quality. As long as they match each other they will have small distances. In order for the procedure to be successful only good traces should be included.

We address this issue by introducing the WLC consistency score in the distance metric. In this way, two traces can have small similarity distance only if they match each other and they satisfy the WLC model. If two traces have high alignment score but they do not fit well the model, which indicates that their quality is low, their distance will be large, and they will not form a cluster.

Finally, we are using a different clustering procedure with respect to other approaches. The major advantages of using density-peak clustering [29] is its simplicity and the fact that it doesn’t require knowing the number of clusters in advance.

The methods described in the literature [19, 20, 21, 22] include procedures for distinguishing different unfolding pathways of the same protein. We haven’t tackled this problem. A possible manner of tackling it would be using the path plot algorithm implemented in Fodis [22] within a cluster obtained with our program and see if different unfolding pathways are included. If yes, they can be divided in groups and further investigated.

### 4.4 Benchmark AFM-SMFS traces.

We tested our procedure on three data sets.

- **Data set I.**

  The first data set contains 101 manually selected traces ascribed to the unfolding of the CNGA1 channel and 4,027 traces from the same experiments containing traces of other proteins or noise. CNGA1 channels were expressed in *Xenopus laevis* oocytes with sample preparation and experimental procedure described in [13]. SMFS experiments were performed in the oocytes membrane with the AFM (NanoWizard 3, JPK). The cantilever was calibrated before the start of each experiment; its spring constant was $\sim 0.08 \text{ N/m}$. The AFM tip was pushed into the surface and a force of 1 nN was applied for 0.5 s to enhance the proteins adsorption. The tip was retracted from the surface at pulling speed 500 nm/s.

  The manual selection of the CNG traces was based on two criteria: the contour length of the curves and their force pattern. According to the interpretation of the experimental data made in ref. [13], the last peak in the CNG traces has a $L_c$ value larger than 220 nm and all CNG traces share a common unfolding fingerprint. The unfolding fingerprint consists of
a peak at $L_c$ around 100 nm corresponding to the unfolding of the cyclic nucleotide-binding (CNB) domain attached to the C-terminus; 3 or 4 force peaks between $L_c$ 120 nm and 250 nm corresponding to the unfolding of the six transmembrane helices and the detachment peak. The 101 CNG traces include traces that satisfy these criteria and some other traces that miss a peak in the middle or the last peak assuming different unfolding pathways as suggested in ref. [13]. Overall, the selected CNG traces can be divided in two groups based on the number of force peaks: traces with 5 to 6 major force peaks and traces with 4 major force peaks.

• **Data set II.**

The second data set contains a mixture of four manually selected groups of F-x curves corresponding to the unfolding of different proteins.

1. Group number 1 contains 35 F-x curves representing one of the possible unfolding pathways of rhodopsin in the plasma membrane of the rod outer segment (ROS) of *Xenopus laevis* as described in ref. [12]. The traces contain 4 to 5 major force peaks. The detachment peak has a $L_c$ value around 100 nm. The experiments were performed with the AFM (NanoWizard 3, JPK). The cantilever spring constant was $\sim 0.08$ N/m. The cantilever was calibrated before each experiment. A contact force of 1 nN was applied by the tip into the surface for 0.5 s. Subsequently, the tip was retracted at pulling speed 500 nm/s.

2. Group number 2 contains 61 F-x curves with a characteristic unfolding pattern that hasn’t been assigned to any protein yet. They come from SMFS experiments in the plasma membrane of primary hippocampal neurons using the cell unroofing technique described in section 4.1. The pulling experiments were performed on the membrane patch adsorbed on the glass arrow at pulling speed 500 nm/s. The cantilever spring constant was $\sim 0.08$ N/m. The cantilever was calibrated before each experiment. The contact force was 1 nN, applied for 1 s before retraction.

The 61 F-x curves belonging to this group contain 5 to 6 major peaks. The detachment peak is with $L_c$ value between 220 and 320 nm.

3. Group number 3 contains 46 F-x curves representing the unfolding of another unknown protein coming from the same experiments like group 2. The traces contain 3 to 4 main peaks. The last peak has a $L_c$ value between 150 and 220 nm.

4. Group number 4 is built from the manually selected 101 CNGA1 traces included in data set I.

• **Data set III.**

The third data set comes from unfolding experiments in the plasma membrane of the rod outer segment (ROS) of *Xenopus laevis* with experimental protocol described in subsection 3.1.2.
The plasma membrane hosts a variety of membrane proteins among which the CNG channels and rhodopsin are the most common [12].

The entire data set contains 386,756 traces.

4.5 Results.

4.5.1 Results in data set I.

Data set I contains 4,128 traces, 101 of which were manually selected by visual inspection and attributed to the unfolding of the membrane protein CNGA1 (for details see section 4.4). After filtering the traces, their number was reduced to 662. 91% of the manually selected CNG traces passed the filters.

By applying our procedure, we obtained six clusters. In Figure 4.8 the clusters are represented by the superimposition of their nine highest density members aligned to the cluster center depicted in orange. The alignment used for this graphical representation was accomplished in an automatic manner using the software Fodis [22]. All manually selected CNG traces were found in cluster number 1 except for two that were assigned to cluster number 6. Therefore, cluster 1 is the CNG cluster. The CNG unfolding pattern consists of a double peak around 100 nm, three major peaks between 120 nm and 200 nm and the detachment peak corresponding to \( L_c \) value \( \sim 290 \) nm. The peak at 100 nm is associated with the unfolding of the CNB domain, while the force peaks in the middle are associated with the pairwise unfolding of the six transmembrane helices in the CNGA1 channel.

The remaining five clusters contain traces much shorter than the CNG traces. The traces in cluster 2 have \( L_c \) values of the last peak between 70 and 80 nm and 2 major peaks. The cluster center has \( L_c \) value of the last peak \( \sim 70 \) nm and 2 major peaks at relatively low forces. The traces in cluster 3 have \( L_c \) between 90 and 120 nm and 1 or 2 major peaks. The last force peak is the only peak in the cluster center of cluster 3 and it has an \( L_c \) value of \( \sim 100 \) nm. Also the cluster center of cluster 4 has one peak but with a smaller \( L_c \) value of \( \sim 80 \) nm. The members of cluster 4 are characterized by \( L_c \) values between 80 and 90 nm and only 1 major peak. In cluster 5 the \( L_c \) is between 120 and 150 nm and 1 or 2 major peaks are present. The cluster center of cluster 5 has two force peaks, the first one is in the very beginning up to extension 20 nm. The \( L_c \) value of the last peak is \( \sim 130 \) nm. In cluster 6 the \( L_c \) values are longer, from 140 to 180 nm and 2 or 3 major peaks are present. The cluster center has two clean force peaks with \( L_c \) values \( \sim 95 \) and 145 nm. With the data that are available, we cannot aim at relating these clusters to proteins or further investigate their molecular origin.

Approximately 54% of the population of cluster 1 is made by the CNG traces. The overall content of the cluster is visualized in Figure 4.9 where we plot the cluster members ranked by their density in a descending order. The highest density traces are the CNG traces with 5 to 6 peaks (the
Figure 4.8: Superimposition of the nine highest density members in each cluster in data set I automatically aligned to their cluster center depicted in orange. The manually selected CNG traces are found in cluster 1.
blue area) followed by the CNG traces with 4 peaks (the green area). If we look at Figure 4.9 more closely, we notice a very thin gray line representing high density traces that haven’t been included in the CNG selection. We looked at these traces and found out that they are very similar to the cluster center of the CNG cluster (Figure 4.10). Therefore, these traces can be considered CNG traces which were not noticed in the manual selection. Remarkably, our procedure was able not only to detect them but also to group them together in the right cluster.

Figure 4.9: Graphical representation of the CNG cluster content. The cluster members are ranked by density in descending order. The blue area shows the manually selected CNG traces with 5 or 6 force peaks; the green area - manually selected CNG traces with 4 peaks; the red area - traces with contour length greater than 350 nm; the gray area - all traces.

Figure 4.10: a-d. Four high-density traces (in pink) included in the CNG cluster by our program which escaped the manual selection. The CNG cluster center is depicted in black.
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The long traces (length > 350 nm) in the cluster have significantly smaller densities. The shape of these traces varies a lot and a common unfolding pattern can not be easily found. Moreover, these traces are not similar enough to each other to form a separate cluster. For this reason we consider them as outliers in the periphery of the cluster. Consistently with this interpretation, their density is very low.

4.5.2 Validation of the main parameters.

Bin width of the \( L_c \) histogram.

An important parameter in our procedure is the size of the bin width of the \( L_c \) histogram. We chose it to be 8 nm, which corresponds to the typical \( L_c \) of a transmembrane \( \alpha \)-helix. In order to validate this choice, we applied our method on the CNG data set using different bin sizes, 1, 3, 7, 9 and 12 nm respectively. With bin width 1 nm, we obtained 2 clusters and with bin width 3 nm - 4 clusters. With bin widths 7, 9 and 12 nm we obtained 5 clusters, opposed to 6 clusters at bin width 8 nm. The CNG cluster was not found with bin widths 1 and 3 nm. In fact, only 2 of the manually selected CNG traces passed the filtering criteria defined in section 4.2 at bin width 1 nm, and 50 at bin width 3 nm. These results exclude the possibility of using such small bin sizes in our procedure. The 50 CNG traces remaining in the 3 nm case were assigned to cluster 4 but they do not form the core of the cluster. The first CNG trace has a density rank 119. With bin widths 7, 9 and 12 nm the CNG cluster remains well-defined. We will compare the results for these bin widths to the results for bin width 8 nm used as a reference.

With bin width 7, 9 and 12 nm, two of the reference clusters were merged but these two clusters were different at each bin width. Cluster 2 at bin width 7 nm is made from the merging of reference clusters 3 and 4, and cluster 5 at bin widths 9 and 12 nm is made from the merging of reference clusters 5 and 6. The change in the bin width leads also to a tiny difference in the number of traces remaining after applying the score-length ratio filter. In the reference setup, the number of remaining traces is 662. With bin width 7 nm, it is 657; with bin width 9 nm - 675 and with bin width 12 nm - 664.

More quantitatively, we use three descriptors to compare the results with the reference setup. The first descriptor measures how many among the ten highest density members in the reference cluster are found in the current cluster. It can take values between 0 and 10. The second descriptor counts how many of the ten highest density members in the reference cluster are among the ten highest density members in the current cluster. It can also take values between 0 and 10. The third descriptor gives the lowest density rank among the ten highest density members in the reference cluster. It can have values between 1 and \( M \), where \( M \) is the cluster size. For example, if we compare a reference cluster with itself, the corresponding sequence of descriptors will be 10-10-10.

We summarized the results in Table 4.1. If a given field of the table contains two sequences of descriptors this means that the corresponding cluster is made from the merging of two reference
clusters. The CNG cluster is present both at the three bin widths with very similar descriptor values. For bin width equal to 7 nm, its descriptors are 10-6-55. This means that all ten highest density members in the reference are assigned to the analogous CNG cluster, 6 of them are among the ten highest density members in the cluster and the reference member farthest away from the cluster center has density rank 55. In the example with bin width 9 nm this sequence almost didn’t change, it is 10-7-57. We do not compare the remaining 4 clusters at different bin widths with each other because they correspond to different reference clusters. A proper comparison is the one with the corresponding clusters at bin width 8 nm. For example, cluster 3 at bin width 7 nm corresponds to cluster 5 at bin width 8 nm, cluster 3 at bin width 9 nm corresponds to cluster 3 at bin width 8 nm and cluster 3 at bin width 12 nm corresponds to cluster 2 at bin width 8 nm.

We performed an additional check to see if the ten highest density members in the four non-CNG clusters obtained with bin sizes 7, 9 and 12 nm were found in the corresponding reference clusters and we confirm it for bin sizes 7 and 9 nm. With bin width 12 nm, this is valid for the first two clusters only, including the CNG cluster. The ten highest density members of the remaining three clusters at bin width 12 nm are split among the reference clusters. This suggests that the clustering results are robust with respect to small changes in the bin width.

There might be changes in the density rank between the different setups but the clusters content is not seriously affected, especially if the data set contains a significant number of high-quality traces corresponding to the same protein. Indeed, the CNG cluster is extremely robust with respect to changes of the bin width in the range 7-12 nm. Instead, reducing the bin width to 5 nm or less is for sure detrimental.

Different \( k \) values in the \( k \)-NN density estimator.

We performed tests with different values of the parameter \( k \) used in the \( k \)-nearest neighbors density estimator used in the clustering (Eq.4.13). The default value of \( k \) is 3 and we use these results as reference. We applied our procedure to data set I using \( k \)-values 2, 4, 5 and 10. We obtained different number of clusters for the different \( k \)-values. We present the results in Table 4.2. As \( k \)-increases, the total number of clusters decreases. The number of clusters we obtain changes significantly for small values of \( k \), for \( k \)=2 to \( k \)=3, from 12 to 6 clusters. If we increase \( k \) from 5 to 10, the number of clusters changes from 4 to 2, which is a small change with respect to the difference between the two \( k \) values. From Table 4.2, we also learn that in all setups the CNG cluster is found independently of the exact \( k \)-value. This shows that the CNG cluster detection is robust with respect to the value of \( k \).

We used the same three descriptors like in the bin width comparison to quantify the clustering results with different \( k \) values. The descriptors values are given in Table 4.2. The CNG cluster is found in all setups with relatively similar values of the descriptors. We tried to match the non-CNG clusters to the reference clusters and we were able to do it only for the remaining three clusters at
$k = 4$. We matched them successfully to the reference clusters at $k = 3$ and they remain quite robust. The only difference is that reference clusters 5 and 6 got merged into cluster 3. The large number of clusters obtained with $k = 2$ makes results hard to interpret. It was difficult to match the results for the remaining three clusters also at $k = 5$. The reason is that among the ten highest density members of these clusters, members from different reference clusters were found. For example, the highest density members of cluster 3 belong to reference clusters 2, 3, 4 and 5. Anyway, none of them contains members of the CNG cluster.

We relate the fact that the CNG cluster is identified at all $k$ values to the high density of high-quality CNG traces very similar to each other in this cluster. When the core of the cluster is made from a smaller portion of similar traces, the results become sensitive to the precise value of $k$. We observe this in the given examples with the non-CNG clusters. Different clusters get merged in the different setups. Moreover, the total number of clusters changes significantly for the different $k$ values. This can be explained as follows. When $k$ increases, the distances might become too large to detect separate clusters. The density estimation becomes inaccurate and clusters get wrongly merged. As we will see, most of the data sets we are working on are highly heterogeneous and the amount of high-quality traces similar to each other is very small (see subsection 4.5.4). This makes the $k$ value important for the final outcome of this procedure. We use as default a $k$ value of 3, which according to us is neither too small, neither too large for the adequate description of the data sets under consideration.
Table 4.1: Clustering results obtained with different bin widths of the $L_c$ histogram. Each sequence of numbers corresponds to the values of three descriptors measuring the difference of the results from the reference calculation, obtained with bin width 8 nm. The first descriptor indicates how many among the 10 highest density members in the reference setup are found in the cluster. It takes values between 0 and 10. The second descriptor measures how many of the 10 highest density members in the reference cluster are among the 10 highest density members in the current cluster. It takes values between 0 and 10. The third descriptor is the lowest density rank of a trace from the 10 highest density cluster member in the reference setup. It takes values between 1 and $M$, where $M$ is the cluster size.

<table>
<thead>
<tr>
<th>Bin Width, nm</th>
<th>CNG Cluster</th>
<th>Cluster 2</th>
<th>Cluster 3</th>
<th>Cluster 4</th>
<th>Cluster 5</th>
</tr>
</thead>
<tbody>
<tr>
<td>7</td>
<td>10 — 6 — 55</td>
<td>10 — 4 — 27</td>
<td>7 — 7 — 7</td>
<td>7 — 6 — 64</td>
<td>8 — 8 — 9</td>
</tr>
<tr>
<td></td>
<td></td>
<td>10 — 5 — 31</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>9</td>
<td>10 — 7 — 57</td>
<td>9 — 8 — 13</td>
<td>10 — 5 — 35</td>
<td>6 — 6 — 9</td>
<td>7 — 5 — 13</td>
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<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>10 — 4 — 24</td>
</tr>
<tr>
<td>12</td>
<td>10 — 6 — 41</td>
<td>8 — 5 — 17</td>
<td>4 — 3 — 13</td>
<td>5 — 1 — 37</td>
<td>8 — 5 — 20</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>10 — 3 — 34</td>
</tr>
</tbody>
</table>

Table 4.2: Clustering results obtained with different values of $k$ in the $k$-NN density estimator used in the clustering. In the first column, the total number of clusters obtained is listed. In the remaining columns, each sequence of numbers corresponds to the values of the same three descriptors of Table 4.1, with respect to the clustering results with the default $k$ value of 3, used as a reference. The question marks indicate undetermined descriptor values due to difficulties in matching the cluster to a single reference cluster. The dashes indicate non-existing clusters. The 12 clusters obtained with $k = 2$ are not listed in the table because they can not be easily related to the 6 reference clusters.

<table>
<thead>
<tr>
<th>$k$</th>
<th>Number of clusters</th>
<th>CNG Cluster</th>
<th>Cluster 2</th>
<th>Cluster 3</th>
<th>Cluster 4</th>
</tr>
</thead>
<tbody>
<tr>
<td>4</td>
<td>4</td>
<td>10 — 6 — 21</td>
<td>7 — 7 — 8</td>
<td>10 — 9 — 12</td>
<td>9 — 8 — 13</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>10 — 1 — 57</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>10 — 0 — 69</td>
<td></td>
</tr>
</tbody>
</table>
4.5.3 Results in data set II: Rhodopsin + Unknown 1 + Unknown 2 + CNGA1.

Data set II contains four different groups of manually selected traces and as a final result we would like to obtain four clusters. The first group of traces in data set II consists of 35 traces representing the unfolding of rhodopsin in the plasma membrane of the rod OS. The second group contains 61 traces associated with the unfolding of an unknown protein in the hippocampus, which we label as unknown protein 1. The third selected group contains 46 traces representing the unfolding of a different unknown protein in the hippocampus, labeled as unknown protein 2. The last group contains the 101 traces coming from the unfolding of the CNGA1 channel. The total number of traces in data set II is 243. After filtering, 194 traces, $\sim 80\%$, remained and were clustered together.

We obtained the three clusters depicted in Figure 4.11. The traces representing the unfolding of unknown protein 1 and the CNGA1 channel were properly identified and grouped into two separate clusters: clusters 2 and 3 (Figure 4.11b and c). The traces representing the unfolding of rhodopsin and unknown protein 2 were grouped together in a common cluster, cluster 1 (Figure 4.11a). The five highest density members in cluster 1 are rhodopsin traces, followed by five traces from the unfolding of unknown protein 2.

In order to better understand the reasons for this error we performed additional tests changing some of the parameters in our procedure. Given that data set II contains only good traces which were manually selected, we decided to change the program setup by using a slightly milder criterion for cutting the non-contact parts, considering as meaningful the last peak exceeding a $3\sigma_{\text{NOISE}}$ threshold instead of the default $2\sigma_{\text{NOISE}}$ threshold (see subsection 4.2.1). In this way, a larger portion of traces was kept after filtering - 216 traces, $\sim 89\%$ of the initial number of traces. As a result, we obtained four clusters corresponding to the four selected groups. The clusters superimpositions are shown in Figure 4.12. The members of the manual group selection for rhodopsin and CNGA1, were exclusively included in clusters 1 and 4 respectively. The majority of curves related to the unfolding of the two unknown proteins were found in clusters 2 and 3 respectively, but some of their manually selected members were spread between the other clusters.

These results indicate that a cluster is identified robustly and reliably if $\sim 100$ high quality traces corresponding to the same protein are present in the data set. If this number is smaller, or the traces are not so similar (like in the case of rhodopsin), the clustering partition becomes less robust, and can change (for better or worse) if an important parameter is modified.

4.5.4 Results in data set III.

We now describe the results for the data set which motivates our work and presents a true challenge for a traditional method for AFM-SMFS data analysis. The data is coming from experiments performed in the plasma membrane of the rod OS under native conditions without overexpression
Figure 4.11: Graphical representations of the three clusters obtained in data set II with the default parameters values. In each panel the superimposition of the nine highest density members automatically aligned to the cluster center (in orange) is plotted. Each panel represents a different cluster. The cluster number is indicated in the top right corner.
Figure 4.12: Graphical representations of the four clusters obtained in data set II with \(2\sigma_{\text{NOISE}}\) threshold for the removal of the non-contact part in each trace (see subsection 1.2.1). In each panel the superimposition of the nine highest density members automatically aligned to the cluster center (in orange) is plotted. Each panel represents a different cluster. The cluster number is indicated in the top right corner.

of certain types of membrane proteins. The data are therefore highly heterogeneous.

The number of traces in the raw data set is 386,756. After the removal of the non-contact part and the discarding of traces shorter than 50 nm, abnormal traces, etc. (for details go to subsection 4.2.1), 41,048 traces remained, \(\sim 11\%\) of the initial amount. The removal of such large portion of spectra is something we were expecting given that the minority of pulling experiments lead to the successful unfolding of a protein and the majority of the recorded spectra do not contain meaningful unfolding events or any unfolding events.

In Figure 4.13 we present the two-dimensional probability distribution function of the length and the score/length ratio. The subset of traces we used to generate the plot are the 41,048 remaining after the initial preprocessing and filtering. From this plot, we learn that most of the traces are very short (less than 100 nm) and with low scores (less than 0.1). The dashed area corresponds to traces falling under the 0.5 ratio threshold we use for filtering. Long traces (length above 250 nm) with high scores have the lowest probabilities. According to Figure 4.13, the high score traces with high probability present in the data set are relatively short, with maximum trace lengths around 100-120 nm. When we apply the 0.5 ratio threshold, we remain with 20,170 traces which is \(\sim 5\%\) of the initial data amount.

By applying our procedure to data set III, we find 29 clusters. In density peak clustering, all traces in the data set are assigned to a cluster but this is not always appropriate. If we take a cluster center and look at the distances of all cluster members to the cluster center, some of these distances
CHAPTER 4. AUTOMATIC CLASSIFICATION OF SMFS DATA FROM MEMBRANES.

Figure 4.13: Two-dimensional probability of the length and the score/length ratio in data set III. The data used to generate the plot contains 41,048 traces. The red areas correspond to regions with high probability, while the purple areas correspond to regions with low probability.

will be large and accordingly the similarity between the cluster center and the certain traces will be low. In order to interpret the results it is necessary to determine the size of a cluster, under the condition that all cluster members should be similar to the cluster center. In order to decide what "similar" means, we sorted the cluster members according to their distance to the cluster centers. An example is shown in Figure 4.14. The cluster center of cluster 24 is plotted in panel a. In panels b-f, different cluster members (in pink) are plotted together with the cluster center (in black). The distances for each pair are written in the top right corner of each panel. We can see how the similarity between the two traces decreases as the distance between them increases. At distances larger than \( \sim 0.3 \) we can no longer be confident that the cluster members and the corresponding cluster centers share the same features. We therefore fix to 0.3 the maximum distance at which we can be sure that the traces have been properly assigned to a cluster. The traces within this distance from the center will form what we will call the cluster core.

Next, we computed the average score and the average length of each cluster. We did this for the entire cluster and for the core of each cluster. We present the results in Table 4.3. We note that the majority of the obtained clusters contain short traces with lengths up to 100-120 nm. This is consistent with what we see in Figure 4.13, namely that the high score traces with high probabilities have short lengths. Not surprisingly, the average score computed for the core of each cluster is in general higher than the average score computed for the entire cluster with few exceptions. The size of
Figure 4.14: a. The cluster center of cluster 24 from data set III. b-f. Different cluster members (pink) manually aligned to the cluster center of cluster 24 (black). The distance from the cluster center is written in the top right corner of each panel.
each cluster changes drastically when we apply the distance threshold from the cluster center. This is consistent with having only few high quality traces similar to each other. In the case of cluster 10, the cluster population becomes zero. We show in Figure 4.15a the cluster center of cluster 10 and in panels b-d its alignment to the three nearest neighbors. The neighbors have lengths similar to that of the cluster center but their shapes show deviations from the cluster center shape and their similarity is low. Other examples are cluster 19 which remains with 2 cluster members, and clusters 13 and 27 with 4 cluster members.

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Table 4.3: Qualitative description of the 29 clusters obtained from data set III. The average score, the average length and the size of the core of each cluster and of the entire cluster are given. The core of each clusters contains only traces at distance from the cluster center smaller than 0.3.

In Figure 4.16 we show the representative traces of 6 clusters. We plot the superimpositions of
the nine highest density members of these clusters with the cluster centers highlighted in orange. The members of cluster 2 have 2 major force peaks and contour length values of the last peak between 70 and 80 nm on average (Figure 4.16a). The traces in cluster 4 have 2 or 3 major peaks and contour length values around 170 nm (Figure 4.16b). We already commented cluster 10, here we only show how large are the deviations of the cluster members from the cluster center, appearing already on the level of the nine highest density cluster members (Figure 4.16c). The cluster center of cluster 10 has three major force peaks and contour length 246 nm and is obviously a good trace but in this data set there are no other traces looking very similar to it. In cluster 11, the traces have 2 major peaks and contour length values ~ 80-90 nm (Figure 4.16d). Cluster 15 contains traces with 2 major peaks and contour length values around 90 nm (Figure 4.16e). Cluster 24 includes traces with 2 or 3 force peaks and contour length values around 80 nm (Figure 4.16f).

Given that the experiments were performed in the plasma membrane of the rod OS where rhodopsin and the CNG channel are the dominant proteins, one might expect to find a rhodopsin cluster and a CNG cluster. The contour length of a fully stretched rhodopsin with an intact disulfide bond is ideally 95.2 nm [12]. We obtain two decent clusters with contour lengths around 80-90 nm which might correspond to the unfolding of rhodopsin, clusters 11 and 24 (Figure 4.16d and f).

The contour length of the fully-stretched CNG channel is around 290 nm and we do not obtain clusters containing such long traces. We assume that even if there are such traces in the data set they are so few that they do not form a separate cluster.

We decided to test this hypothesis by adding to data set III the portion of manually selected CNG traces included in data set I. This test allows checking if we are still able to find the CNG traces in the bulk of hundreds of thousands of spectra. As a result, we obtain now 30 clusters, clusters corresponding to the previous 29 plus the CNG cluster. This result supports the hypothesis we made: if data set III contained a reasonable amount of CNG traces with both high quality and similarity, our method would have been able to find them. In addition, the obtained results lead to the conclusion that the amount of high-quality traces similar to each other in data set III, which can be uniquely assigned to the unfolding of a particular membrane protein, is relatively small.
Figure 4.15: a. The cluster center of cluster 10 from data set III. b-d. The three nearest neighbors (pink) of the cluster center (black) are plotted together with it. The corresponding distances are written in the top right corner of each panel.
Figure 4.16: a-f. Graphical representations of six manually selected clusters from data set III. The cluster number is given in the top right corner of each panel. In each panel the superimposition of the nine highest density members automatically aligned to the cluster center (in orange) is plotted.

4.6 Conclusions

In this chapter we described a fully-automatic procedure for the analysis of F-x curves coming from experiments performed in native cellular membranes. The method is designed to face a specific challenge: analyzing the huge amount of data obtained by AFM-SMFS experiments on native membrane patches. The algorithm is based on dynamic programming alignment, quality score computation and clustering. When it comes to F-x curves, an alignment step is mandatory to evaluate the similarity between two curves. In order to accomplish this step, we use dynamic programming which provides an alignment score, which is a measure for the similarity between two traces. High alignment score refers to a pair of similar traces. This approach has been already described by
Marsico et al. and applied to a hand-curated data set containing 135 traces from AFM-SMFS experiment on bacteriorhodopsin mutant P50A [20]. In addition, we defined a quality score measuring the consistency of each trace with the WLC model. It is well known that when a protein is stretched in an AFM experiment it behaves like a polymer chain described by the WLC model [44]. The WLC consistency score provides a quantitative measure for the quality of a trace. If the score is high, the trace is good and if the score is low, the trace is bad. One of the key components in our procedure is the distance which combines the alignment score and the quality score in such a way that high-quality traces similar to each other are characterized by a small distance, while low-quality traces, even if they are similar to each other, are characterized by a larger distance. This property of the distance is of crucial importance for the efficient analysis of non-curated data sets and membrane proteins data sets in general, because of the low amount of high-quality traces generated during the AFM-SMFS experiments [21]. The clustering method we use is a modified version of the density-peak clustering [29] with $k$-NN density estimator. The major advantages of this clustering approach are its simplicity, the fact that it is unsupervised and doesn’t require knowing the number of clusters.

Compared to other procedures [19, 20, 68, 22], our method provides filtering of spurious traces and selection of high-quality traces in a non-demanding manner. Mainly, it doesn’t require any previous knowledge on the sample composition and the proteins contour length. We achieve this by combining the WLC quality score with the trace length to measure the amount of experimental data consistent with the WLC model. In this manner we are able to reduce the amount of bad traces and to increase the method’s computational efficiency at the same time by filtering the traces based on their score/length ratio. We remark that our procedure is not appropriate for distinguishing different unfolding pathways, which is something that other methods in the literature can do [19, 20, 68, 22].

We benchmarked our method on a data set containing a manually-selected sample of CNG traces distributed among a larger portion of traces of unidentified quality. Our method successfully detected the CNG traces and grouped them in a separate cluster. Remarkably, it was also able to find CNG traces which escaped the manual selection and to properly assign them to the CNG cluster. This result demonstrates the ability of our method to detect high-quality traces similar to each other and to group them in the same cluster.

We obtained also other clusters whose molecular origin we are currently unable to identify. As we commented in Chapter 3, this issue might be managed through a combination of new experimental data with molecular modeling tools like the one we developed and described in the same chapter.

With our procedure, we were able to analyze a data set consisting of $\sim 400,000$ traces of unidentified molecular origin sampled from the plasma membrane of the rod outer segment (OS). The main motivation for the work presented in this chapter comes from data sets like this one. Our program turned out to be quite efficient taking only $\sim 90$ minutes to analyze the entire data set. The general observation we made from the obtained clusters is that the data set contains overall very few high-quality traces that are very similar to each other. Given that the protein composition
of the rod OS plasma membrane is dominated by rhodopsin and the CNG channel, one can expect to find clusters depicting the unfolding of these two membrane proteins. Indeed, two of the clusters we obtained are decent rhodopsin candidates, but we were unable to find a cluster describing the unfolding of the CNG channel. We decided to perform a test by adding the manually selected CNG traces from the previous data set to the large plasma membrane data set. The algorithm proved to be able to detect the selected CNG traces in the bulk of data and to group them in a separate cluster. This result support the observation we made, namely that high-quality traces, coming from the unfolding of the CNG channel, that are similar to each other are very rare. This makes the clustering of these traces extremely difficult and hinders the identification of characteristic unfolding patterns related to specific membrane proteins. In order to further validate this hypothesis we will have to analyze more data from native cell membranes.
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