Biophysical Characterization of SNARE Complex Disassembly Catalyzed by NSF and αSNAP

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Chapter 1

Introduction

To fulfill their specific function most if not all proteins interact with other proteins in some way or another. The assembly of protein complexes is therefore nothing unusual and one might ask why the mechanism of SNARE assembly should be interesting enough to be explained in great detail. The assembly of SNARE complexes is said to mediate fusion of biological membranes. But why is that so fascinating? How does the cell succeed to fuse membranes? How the right ones, at a sufficient speed, at the right times? Which are the key players, what is their mode of action and how are they being regulated? The main purpose of intracellular membranes being the enclosure and thereby division of different subcellular environments, they are usually spatially separated from each other which keeps them from interacting spontaneously. Furthermore, even if two membranes are in close apposition, the repulsion of charges between them still prevents fusion of different compartments from occurring—as long as no additional input of energy is provided. Therefore a reaction bearing the potential to drive membrane fusion obviously needs to fulfill at least two critical criteria: First, it needs to be able to mechanically draw the membranes close to each other, second it needs to generate enough energy to overcome the energetic barrier. These preconditions given, an additional level of complexity is added when we start to think of regulatory functions providing specificity. Tight regulation concerning the amount, speed and type of vesicles prone to fuse is also required to add to the quality of the fusion reaction. Nevertheless these regulatory mechanisms do not necessarily have to be inherent to the basal fusion machinery but could theoretically also be provided by other factors. Keeping these prerequisites in mind the underlying mechanisms bear some fascination. If the two main criteria, pulling membranes together and overcoming the energetic barrier were to be met by proteins—how should these proteins be structured, how would they interact? Since, as stated in the beginning, the SNAREs actually represent a family of proteins capable to fulfill this difficult task, studying their structure and mechanisms of interaction has helped to gain insights into membrane fusion during the last decades
and probably will continue to do so in the future.

1.1 Function of the SNARE Proteins

All intracellular transport processes ranging from secretion in yeast to neurotransmitter release in the brain, depend on the ability of membranes to fuse with each other. Due to the high variability regarding time-scale and function of these various processes, the group of proteins mediating membrane fusion must on the one hand show common characteristics which enable them to execute the event of fusion. On the other hand however, there must be diversities allowing for tight spatial and temporal regulation. SNAREs can be found in all eukaryotic organisms and their involvement in many steps of intracellular transport has been proven using various experimental systems. It is now commonly believed that they are involved in all vesicular fusion events, with different sets of SNAREs being responsible for different trafficking steps [1]. The SNAREs are usually short and C-terminally attached to the membrane by either a transmembrane region or a membrane anchor. One set of SNAREs typically consists of four proteins, which when combined in appropriate solutions spontaneously form extremely stable complexes via a region common to all SNAREs called the 'SNARE motif' or 'core-domain'. The association takes place along a highly exothermic reaction pathway which already early in SNARE research made it tempting to speculate that the assembly of SNARE complexes provides the driving force for membrane fusion [2, 3, 4]. Indeed, later they were shown to be able to autonomously catalyze liposome fusion in vitro without any additional factors [5].

1.2 A Closer Look at SNARE Assembly

But how can the assembly of proteins lead to the merger of two membranes? Even though the complete answer to this question has not been conclusively found, some by now well understood basic mechanistic principles of SNARE assembly have been of help to establish different models. One of the concepts of how SNARE assembly actually proceeds, the so called 'zipper'-mechanism, has gained more and more support over the years.

1.2.1 The 'Zipper'-Model—How Does SNARE Assembly Proceed?

The zipper model proposes that SNARE proteins residing on opposing membranes first interact at their membrane distal termini to form a loose complex. Starting from this point of interaction they consequently wind up towards their C-terminal
membrane anchors in a zipper-like fashion to form the four-helix-bundle, bringing the membranes into close proximity (illustrated in figure 1.1). The amounts of energy generated during assembly eventually suffice to allow for the membrane merger. In other words, SNAREs directly function as fusion catalysts [6, 7, 8, 9].

![Diagram of SNARE assembly](image)

Figure 1.1: SNAREs switch from 'trans' to 'cis' during fusion. The figure exemplary shows the distribution of the neuronal SNAREs before and after fusion. Before fusion, the Q-SNAREs reside on the plasma membrane whereas Synaptobrevin (Syb) is found on the vesicle. After fusion of the two membranes into one, all three SNAREs are depicted in complex, indicating that the complex has assembled during fusion. As described in the text, assembly does not only temporally coincide with but rather is instrumental to drive the fusion reaction. The driving force results from conformational changes which are schematically shown: Before fusion the SNAREs are mostly unstructured whereas during fusion a tight helical bundle is formed.

1.2.2 The Driving Force—Spontaneous Formation of Extremely Stable Complexes

Evidence for this model comes from various kinds of experimental systems. The first mechanistic insights resulted from experiments with recombinant neuronal SNAREs in solution. Most of the early experiments were performed using only the SNARE core domains, which were found to be sufficient to mediate SNARE assembly displaying similar biophysical characteristics as the full-length SNARE complex. A
common characteristic of these motifs is a lack of secondary structure as long as they are in a monomeric state [3, 4, 6, 7, 10, 11]. Only upon mixing with other SNAREs do certain combinations lead to the spontaneous association into complexes accompanied by major conformational and free-energy changes. Even though the two neuronal SNAREs Syntaxin1 and SNAP-25 (synaptosome-associated protein of 25kDa) alone form complexes displaying high $\alpha$-helical content, combining all three neuronal SNAREs produces amongst other side-products one uniquely stable outcome, which consists of all three neuronal SNAREs in a 1:1:1 stoichiometry. Not only is it resistant to higher amounts of SDS than all other resulting complexes and extremely high temperature—it also shows a pronounced hysteresis meaning that there is basically no spontaneous dissociation under assembly conditions [12]. Since all other side products are in equilibrium with the reactants, the degree of this respective complex increases with reaction time. Furthermore this ternary SNARE complex can be isolated from brain extracts indicating its existence in vivo [13] making it the most probable end product of the assembly pathway.

1.2.3 The Logistics—SNAREs Switch from 'trans' to 'cis' During Membrane Fusion

The SNAREs are in a 'trans'-state as long as fusion is incomplete (residing on opposite membranes). As soon as the membrane merger has taken place and all helices are anchored in one membrane the complex is considered to be in a 'cis'-configuration (figure 1.1). Originally the SNAREs were classified into vesicle (v-) and target (t-)SNAREs, according to their location on the membrane before fusion [14]. The drawback of that nomenclature is however that the functional distribution of some SNARE sets has not been characterized sufficiently to assign vesicle- or target-function and especially when it comes to homotypic membrane fusion this classification meets its limits. By comparison of the neuronal ternary complex crystal-structure to those of two other, only distantly related (endosomal) SNARE complexes, it became evident that all three complexes resemble an intertwined $\alpha$-helical bundle of parallel coiled coil domains consisting of four helices per complex [15, 16, 17]. The parallel alignment of SNAREs in the ternary complex mentioned in the context of the crystal structure was originally observed with the help of FRET- as well as EM-experiments [8, 18]. These SNARE complexes show a remarkable degree of conservation which led to a reclassification of SNAREs based on a structural property: The centre of the bundle contains 16 stacked 'layers' of interacting side chains which are largely hydrophobic, except for the very central one which contains three highly conserved glutamine (Q) residues and one highly conserved arginine (R) residue. Hence the SNAREs with a glutamine in this so called '0'-layer were grouped Q- and the ones with arginines R-SNAREs. The Q-
SNAREs can further be divided into Qa-, Qb- and Qc-SNAREs, each of these classes contributing one helix to the four helix bundle [6]. The neuronal SNAREs belong to a unique subset of SNAREs in the sense that rather than being part of two independent proteins, the Qb and Qc-SNARE motifs are accommodated in one SNARE called SNAP-25 (synaptosome-associated protein of 25kDa). It is tempting to speculate that it represents a fusion product of two reactants of the assembly reaction which has evolved because it may have led to a kinetic advantage over a four-component reaction. Thinking along these lines one can imagine that neuroexocytic events profit from this advantage possibly contributing to the high speed of fast firing neurons. Syntaxin and Synaptobrevin represent the second neuronal Q-SNARE and the R-SNARE, respectively. The neuronal SNAREs are distributed such that the R-SNARE is primarily located on the vesicle whereas the Q-SNAREs are predominantly located on the cell membrane. If the structural arrangement is a general feature of all SNARE complexes, their localization to the different membranes might also be conserved, but since characterization of different SNARE topologies has not come to an end yet, this at the moment is mere speculation.

According to this hypothesis one membrane would typically contribute three SNARE motifs whereas the second would contribute the fourth helix. On the one hand experiments on neuronal SNAREs and in yeast have led to this hypothesis. On the contrary homotypic in vitro-fusion of liposomes mediated by early-endosomal SNAREs has been shown to proceed with different SNARE topologies regardless of whether all Q-SNAREs are contributed by the same membrane or not [17]. Whether a 3Q/1R-rule holds true for all SNARE sets in a physiological context therefore remains a matter of debate and yet needs to be elucidated. Taken together, these findings are all in accordance with the zipper model: First of all the assembly reaction is exogenous and hence a source of energy, which spontaneously proceeds without additional factors. Second, the SNAREs reside on opposing membranes before, whereas they are found on the same membrane after fusion. Third, the helices are aligned in parallel after assembly, thus the N-termini interact with the N- whereas the C-termini interact with the C-termini of the partner SNAREs.
1.3 Postfusion-Time for Recycling of SNARE Complexes

After assembly, the complexed R- and Q-SNAREs reside in the same membrane and are no longer free to act in further rounds of fusion. As much as the above described characteristics like pronounced hysteresis, exothermal driving force of the reaction and the high level of complex-stability push the reaction into the direction of assembly they of course prevent the complex from voluntary disassembly under physiological conditions. Hence basically no spontaneous dissociation occurs. If there were no counteracting mechanism, all fusion events would come to an end as soon as all free SNAREs were used once. Of course the cell could constantly dispose of the fully assembled SNARE complexes and synthesize new free SNAREs in a single-use fashion, but keeping in mind that during assembly the SNAREs only undergo structural but no degradational changes, one can envision a faster and probably energetically more favorable solution to this problem: A factor which tears the complexes apart despite their apparent stability, e.g., by lowering the activation energy of this thermodynamically unfavorable reaction. The cell has exploited a family of ATPases which due to their functional variability have been assigned ‘ATPases associated with various cellular activities (AAA ATPases)’, to lower the activation energies of various reactions, mostly connected to folding or unfolding of proteins. One of these ATPases called NSF (N-ethylmaleimide sensitive fusion protein), with the help of its co-factors, the so called SNAPs (soluble NSF-attachment proteins), mediates the disassembly of the stable SNARE complexes and thereby recycles the fusion machinery [19]. To do so NSF manages to couple ATP hydrolysis to the highly endothermic dissociation of the 4-helix bundle.

1.3.1 NSF and the SNAPs – The Disassembly Machinery

NSF is ubiquitously expressed and therefore most likely capable of disassembling all existing SNARE complexes of a certain species [19]. Its universal mechanism is also underlined by the fact that to a certain extent functionality even between different species seems to be conserved. As an example the NSF yeast homologue Sec18p has been shown to stimulate exocytosis in permeabilized adrenal chromaffin cells [20].

NSF consists of three domains, two of which, termed the D1- and D2-domain, can bind ATP [21]. At one of these binding sites ATP is actively being hydrolyzed to provide energy during disassembly, whereas ATP-binding to the second site functions in oligomerization of the protein [22, 21], which is largely hexameric under equilibrium conditions [23].

SNARE complexes do not display any direct binding sites for NSF. In order for it to disassemble the complex, SNAPs are required which provide high affinity
binding sites for both, the enzyme as well as its substrate [24]. SNAPs therefore
serve as connectors between the SNARE complex and NSF which can disassemble
the SNARE complex in the presence of Mg$^{2+}$-ATP as soon as one or more, most
likely three, SNAPs have bound (figure 1.2).

\[ ... \]

\[ ... \]

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{figure12.png}
\caption{Disassembly of SNARE complexes. (A) The NSF-molecule consists of
three subdomains which are schematically depicted. The N-domain includes the binding
region for $\alpha$SNAP, the D1 domain actively hydrolyses ATP during disassembly and the D2
domain plays a role in oligomerization of the molecule, which is mostly hexameric under
equilibrium conditions.
(B) Most likely three $\alpha$SNAPs bind to one SNARE complex to form a recognition site for
one hexamer of NSF. Since no high resolution structure of NSF in its hexameric state has
been solved it is shown schematically. In the presence of Mg$^{2+}$ NSF then disassembles the
SNARE complex, a process which is fuelled by hydrolysis of ATP into ADP+P$_i$.}
\end{figure}

1.3.2 How Does This Transition of Energy Take Place?

Compared to the assembly reaction the molecular details of disassembly are much
less understood. Electron microscopic images of NSF gave some insight into the
mechanism in the sense that clearly distinguishable structures were visible, depend-
ing on the respective nucleotide bound [8]. These pictures suggest that hydrolyzing
ATP to ADP+P$_i$ leads to conformational changes in the whole NSF-hexamer which
might exert mechanical force onto the complex it is bound to.
Hexamers have a donut-like shape, which gives rise to the speculation that the hole in the middle might fulfill some kind of function, especially since other likewise hexameric AAA ATPases are known to use their holes to pull interaction partners through [25, 26]. On the one hand, it is well imaginable that NSF drags one of the SNAREs through its center pore during disassembly, thereby separating it from the other SNAREs. On the other hand the size of the pore in an NSF-hexamer would theoretically be large enough to accommodate an unfolded protein, like e.g. monomeric synaptobrevin, but according to EM-data meet its limitations as soon as folded domains were to fit through [8]. In some sets of SNAREs each of the four proteins includes one domain with inherent tertiary structure which makes this model improbable unless drastic conformational changes facilitating pore enlargement take place during ATP-hydrolysis.

Alternatively, the NSF-hexamer might, while sitting on the scaffold provided by SNAPs, create a rotational force through conformational changes which moves the SNAPs in a circular manner. The SNAPs being connected to the SNARE complex would insert the same rotational force onto the complex. Since this again is rigidly attached to the membrane it is not difficult to picture that as a consequence the complex would be unwound into its individual SNAREs, like operating a mandrel in the opposite direction would unwind a thread into its individual fibers.

Originally it was widely believed that disassembly proceeds in a ‘symmetric’ fashion, all six NSF-molecules synchronously hydrolyzing ATP and only then leading to conformational changes in the ring-like structure which consequently enable dismantling of the SNARE complex. More recently an alternative possibility has been raised, which is based on studies undertaken on another AAA-protein called ClpX [27]. Here the authors could show that the single subunits of one ClpX-hexamer were capable to exert their catalytic function independently from each other, whereas a concerted action of several subunits apparently increased the enzymatic efficiency in a proportional dependence. If a comparable mechanism were to be utilized by NSF, it would also be conceivable that, depending on the number of NSF-subunits participating in a specific reaction, the amount of αSNAPs per disassembly event is similarly flexible. Nevertheless no attempts have been made to investigate whether NSF-activity also works in an asymmetric manner, or whether all six subunits need to bind and hydrolyze ATP in concerted fashion.

1.3.3 α-, β- and γ-SNAP – Specialized or Redundant Isoforms?

Besides αSNAP two more SNAP-isoforms called βSNAP and γSNAP are known [28]. Of all three mammalian SNAPs, the structure has only been solved for γSNAP using crystallographer [29]. Despite limited sequence homology, the γSNAP structure
strongly resembles that previously solved for the SNAP homolog in yeast, Sec17 \([30]\). Even though \(\alpha\)SNAP alone is sufficient to serve as a co-factor for disassembly in \textit{in vitro} experiments, it remains unclear whether the other known SNAP isoforms \(\beta\)- and \(\gamma\)-SNAP function as positive or negative regulators or simply provide redundancy despite relatively low sequence homology \([31, 32, 33, 34]\).

In fact so little is known about the mechanism that the number of \(\alpha\)SNAPs required is still debated. Three might be needed, but alternatively one or two might also suffice. In any case participation of more than three can be excluded. It still remains unclear how exactly one SNARE complex, one to three \(\alpha\)SNAPs, but six NSF molecules (meaning one hexamer) can reasonably be put together. No structure of the complete disassembly machinery, also termed the '20S'-complex \([24]\), has been solved leaving room for speculations.

1.3.4 Input and Output of the Machinery – Which Other Substrates or Products are Feasible?

Neither all substrates nor products of the disassembly-reaction are known. For instance SNAREs could theoretically be fully disassembled into monomers but alternatively might also only be partially disassembled up to an intermediate stage. SNAREs contributed by one membrane could for example be kept together in order to decrease the complexity of the subsequent fusion reaction.

Likewise even though there is no doubt that the ternary SNARE complex is a target of NSF, it cannot be excluded that other complexes e.g. assembly intermediates or unwanted dead-end-complexes also serve as substrates for NSF-mediated disassembly. If this were to be the case, NSF might in addition to its indirect effect on SNARE assembly, in terms of SNARE recycling, also directly regulate complex assembly by influencing the steady-state concentrations of complex intermediates.

1.3.5 Potential Regulatory Mechanisms of SNARE Disassembly

Even though rapid SNARE-complex disassembly is essential for the cell in order to provide sufficient amounts of free SNAREs to readily replenish SNARE pools after fusion, it is nevertheless of high importance that SNARE disassembly, like all vital physiological processes, is tightly regulated. It should for example be granted that uncontrolled disassembly does not deprive the cell of factors which at the same time are needed for other vital processes. \(\alpha\)SNAP as well as NSF have been reported to be involved in various other cellular functions independently of each other, making it feasible that there are mechanisms which inhibit disassembly as soon as one of them becomes limiting. Likewise, since ATP is a fuel needed for a large variety of other
vital cellular processes as well, NSF activity might be restricted under conditions in which ATP levels are low. Furthermore, disassembly of SNARE complexes prior to completion of assembly would, according to current knowledge, prevent final membrane merger and should hence also be regulated by the cell.

SNARE-complex disassembly being mediated by the combined efforts of two proteins brings about the question of whether regulatory mechanisms act on the level of the enzyme, the adaptor or either of them. Finally the SNARE target might also be the targets of regulation. Up to date, several groups found potentially regulating factors, covering all three of these possibilities.

The enzyme – Regulation of NSF activity

For NSF, nitrosylation and phosphorylation, both having an apparent inhibitory effect on its function, have been reported. Phosphorylation at residue Ser-237 by a Serine/Threonine-kinase supposedly results in a hexameric form which does not bind SNAP/SNARE complexes [35]. Likewise, phosphorylation at the tyrosine-residue 83 has been suggested to lead to reduced binding of αSNAP [36]. In vitro phosphorylation has also been witnessed for αSNAP where it apparently led to a ten-fold reduction in SNARE-complex affinity [37].

The adaptor – Regulation on the level of the SNAP

Additionally, proteins able to compete with αSNAP for SNARE complex binding have been described. Amongst these are the so called Complexins (also known as Synaphins) which were originally identified by their interaction with the neuronal SNARE core complex [38, 39].

Complexins The Complexins represent a family of small proteins without folding similarities to any other protein family and are present in all multicellular eukaryotes. Complexins have originally been reported to displace αSNAP from SNARE complex when added in excessive amounts, leading to speculations that they might have inhibitory properties with respect to SNARE disassembly [39]. Notwithstanding these findings, in vitro assays directly monitoring SNARE disassembly do not show any negative influence of Complexin 1 or 2 on SNARE-complex disassembly [40]. However, due to a time resolution in the range of several minutes, these experiments are preferably suited for comparing 'all-or-nothing' disassembly defects rather than minor kinetic differences which could hence not be ruled out. Meanwhile the mode of binding between Complexin and the SNARE core complex has been investigated. An α-helical central stretch consisting of 58 amino acids was found to bind the SNARE complex in an anti-parallel fashion [41], and the structure of complexin bound to the neuronal SNARE complex has been solved [42]. Seemingly contradictory outcomes
of knock-out experiments and over-expression studies in various animals and cell-types have complicated the determination of complexin function, leaving a range of proposed models in the field up to now. Most of these studies report a reduction of exocytosis in presence of excess Complexin which would be in line with a negative impact on fusion. However, at the same time the absence of complexin appears to result in impaired exocytosis as well. Recently, three different studies resulted in findings which led to a similar interpretation, according to which Complexin might act as a so called fusion clamp binding the SNARE complex in a partially zippered state prior to complete fusion and thereby 'clamping' it until the signal for the final fusion step is received [43, 44, 45]. Even though this scenario primarily envisions Complexin to function as an inhibitor of the last step of SNARE assembly, it does not exclude the possibility that the 'Complexin clamp' serves as a protection against unwanted disassembly at the same time. Altogether the apparently contradictory results could in principle also be explained by a fine-tuning capability which for instance might depend on a physiological Complexin concentration. If this were to be the case, too little as well as too much Complexin would harm the balanced fusion process and might lead to similar phenotypes. Complexins’ probably complex role and possible implications on SNARE disassembly hence require further elucidation.

The membrane – Another check-point of SNARE regulation?

Even though SNARE complexes readily assemble and with the help of NSF and αSNAP disassemble in solution, they are naturally located on membranes, which might also influence their mechanism as well as speed of assembly or disassembly. SNAREs have been inserted into membranes but only little has been done to dissect the process on membranes in detail. What has been done up to now has led to contradictory results. EPR-measurements have attributed a possible role of the membrane to control the capability of SNARE motifs to enter SNARE complexes. They indicated that a short membrane proximal region of Synaptobrevin-2 is dipped into the membrane and thereby inhibits the formation of SNARE complexes [46], a regulatory mechanism that was called 'Synaptobrevin restriction' which could in this study be overcome by mutation of two membrane proximal tryptophan residues. This scenario would to some extent contradict the results mentioned above, which suggested that SNARE zippering proceeds in an N- to C-terminal direction. Notwithstanding these findings, other groups have witnessed fusion of native Synaptobrevin membranes and Synaptobrevin has recently been shown to be constitutively active, regardless of whether incorporated into membranes or not [47]. Here Synaptobrevin could be driven into SNARE complexes both in isolated synaptic vesicles and in proteoliposomes. Even though the membrane thus does not seem to regulate SNARE assembly as such, it might influence one of the other steps of the SNARE
Recent, Munc-13 was found to interact with SNARE complexes when these are incorporated into liposomes, whereas they hardly interacted in solution [48].

Likewise, SNARE disassembly might be affected by the membrane. αSNAP and its yeast homologue Sec17 have for instance been suggested to directly interact with membranes independently of the well established interaction with membrane bound SNARE complexes, which of course also confines them to the membrane [49, 50]. This is in line with the observation that αSNAP is an amphipathic proteins which binds to plastic surfaces [51]. Steel et al. report that the stimulation of the NSF ATPase-activity by αSNAP is more pronounced in the presence of lipids [50]. Lastly, NSF is released inefficiently from many cellular membranes under conditions which allow for 20S-complex disassembly. This might be an evidence for NSF-binding to lipids as well. Nevertheless, no experiments directly assessing the impact of membranes on SNARE disassembly have been performed so far, leaving room for further investigations.

1.3.6 The Yeast Homologous Protein Family – How Conserved is the Disassembly Mechanism?

Like all other SNARE-complex structures which have been solved so far, the crystal structure of the yeast plasma membrane complex is very similar to that of the known neuronal and endosomal SNARE complexes [52]. The yeast SNAREs constituting the complex are Sso1 (Syntaxin-homologue), Sec-9 (SNAP25-homologue) and Snc1 (Synaptobrevin homologue). Interestingly, the structures of the neuronal and the S. cerevisiae complex, both of which are involved in plasma membrane fusion, are even more similar to each other than they are to the endosomal SNARE complexes. As opposed to the neuronal SNARE complex, the yeast SNARE complex seems to be less stable. It neither is SDS resistant nor, with a melting temperature of 55 C, is it as heat resistant as the neuronal SNARE complex, which only melts at ∼90 C.

The homologues of αSNAP and NSF in yeast are Sec17 and Sec18, respectively. The Sec17 structure as well as that of the Sec18 N-terminal have been solved crystallographically [30]. In analogy to the neuronal SNARE complex, in vitro binding studies of Sec17 yeast complex coupled to GST-beads revealed that the stoichiometry of Sec17 to SNARE complex is 3:1 [53]. The crystal structures of the NSF and Sec18 N-terminal domains reveal a remarkable degree of similarity [54, 55, 56], suggesting that the mode of interaction with their respective SNAP adaptors might also be conserved. As a matter of fact this structural similarity extends to function in the sense that Sec18 can efficiently replace NSF in biochemical and membrane trafficking assays [57, 58, 20].
1.3.7 Summary

To sum it up, the molecular details of SNARE disassembly remain poorly understood even twenty years after NSF discovery. This is rather astounding, since even though the original hypothesis that αSNAP/NSF might be the factors actually mediating the last step of membrane merger has meanwhile been proven wrong by various convincing findings, their nowadays widely acknowledged function as the mediators of SNARE recycling, does not weaken their fundamental importance for intracellular membrane fusion. On the contrary, keeping in mind that the disassembly reaction actually fuels membrane fusion by means of energy transfer into SNAREs one might still consider it as the motor of membrane fusion. It is also conceivable that NSF acts at several steps of the SNARE cycle, on the one hand disassembling SNARE complexes post fusion, on the other hand disassembling complexed SNAREs either not wanted or not able to form fusion competent complexes. Nonetheless it is clear that this motor does not act at the fusion step itself but is temporally uncoupled from actual membrane merger strongly suggesting that there are mechanisms to prevent its action at either the wrong time or on the wrong substrate. Furthermore, since the reaction uses parts of the cellular energy storage, there most likely exist regulatory factors ensuring balanced energy levels at all times. Such modulatory proteins or conditions might either completely abolish or, in other cases, only decrease or enhance the disassembling activity of NSF and/or the SNAPs. Insights into how exactly this might happen may be reached in studies directly investigating the requirements for optimal SNARE disassembly. Comparison of disassembly kinetics under various conditions and in presence or absence of putative regulatory factors might help to shed light on the mechanistic details of the disassembly reaction.

1.4 Aim of This Study

The aim of this study was to investigate the molecular basis of SNARE disassembly including SNAP homologues, putative regulatory factors and the degree of functional conservation. Obviously, fine-tuning activities would best be observed using dynamic assays with a high time resolution. Secondly, experiments should be easy to manipulate in a defined manner. Additionally, NSF and αSNAP have been reported to promiscuously interact with several other cellular factors as well, making it difficult to distinguish abnormalities in disassembly from other phenotypic effects possibly resulting from other NSF- and SNAP interactions. Therefore the experiments should be placed in a defined background. Many of the findings collected with regard to αSNAP and NSF so far are based on findings in living cells.

Others have used biochemical assays, which either determine the binding properties of the involved proteins or NSF activity by ATP consumption. Both of these
biochemical approaches are indirect in the sense that they do not allow for conclusions concerning the actual dissociation of the SNARE complex.

Others have monitored SNARE disassembly in vitro using SDS page, which is a direct read-out, albeit with only limited time resolution [6]. A dynamic in vitro system allowing for direct monitoring of SNARE interaction would therefore be ideally suited for quantitative determination of disassembly kinetics and putative influencing factors. Since fluorescence spectroscopy of SNARE proteins is well established in our lab, it was the most obvious option to also employ fluorescence spectroscopy for the investigation of their disassembly.

Moreover, the experimental set-up should then be exploited to gain further insights regarding the disassembly mechanism, in particular on which parts or interactions are the bottlenecks of the reaction and thus might represent regulatory targets.

In order to be able to also assess regulatory mechanisms and factors which do not act in an 'all-or-nothing' but rather a modulatory fashion, the conditions during the experiment should be as close to nature as possible. For this it is e.g. desirable, that none of the proteins involved is required in unreasonably high amounts for optimal function. As a prerequisite to addressing regulatory and mechanistic questions on the level of all proteins involved, this in vitro system should thus be optimized in such a way that ATP, αSNAP and NSF can be applied in quantities one would expect according to stoichiometric (or enzymatic) considerations or, if known, endogenous concentrations.

Subsequently, the impact of some regulatory factors should be addressed. Finally, the degree of functional conservation of the disassembly machinery was to be investigated with the help of yeast SNARE proteins and their disassembly machinery.
Chapter 2

Materials and Methods

2.1 Chemicals

Acrylamide/Bisacrylamide solution, Roth (Karlsruhe, Germany)
Agarose (low EEO), Applichem (Darmstadt, Germany)
Ammoniumpersulfate (APS), Sigma (Deisenhofen, Germany)
ATP
Bradford-Reagent, Biorad (Richmond, USA)
Calcium chloride dihydrate, Fluka (Switzerland)
Coomassie Brilliant Blue R-250, Serva (Heidelberg, Germany)
Dipotassium hydrogen phosphate, Merck (Darmstadt, Germany)
Disodium hydrogen phosphate, Merck (Darmstadt, Germany)
Dithiothreitol (DTT), Roth (Karlsruhe, Germany)
EDTA Titriplex III, Merck (Darmstadt, Germany)
Glycine, Sigma (Deisenhofen, Germany)
HEPES (N-(2-hydroxyethyl) piperazine - N-2-ethanesulphonicacid)
Imidazole, Sigma (Deisenhofen, Germany)
Isopropyl-D-thiogalactoside (IPTG), Roth (Karlsruhe, Germany)
Magnesium chloride, Sigma-Aldrich (St.Louis, USA)
MPD (2-methyl-2, 4 pentanediol), Fluka (Switzerland)
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PMSF (paramethyl sulphonyl fluoride), Roth (Karlsruhe, Germany)
Potassium dihydrogen phosphate, Merck (Darmstadt, Germany)
Sodium hydrogen carbonate, Merck (Darmstadt, Germany)
Sodium chloride, Merck (Darmstadt, Germany)
Sodium cholate, Sigma-Aldrich (St. Louis, USA)
Sodium dodecyl sulfate (SDS), Biorad (Richmond, USA)
TEMED (N, N, N, N- Tetramethylethylenediamine), Biorad (Richmond, USA)
Tricine, Applichem (Darmstadt, Germany)
Tris (hydroxymethyl)-aminomethane (Tris), Merck (Darmstadt, Germany)
Triton X-100, Merck (Darmstadt, Germany)
Tween - 20, DAKO (Carpinteria, USA)
Urea, Merck (Darmstadt, Germany)

2.2 Chemical solutions

Acetic acid, Merck (Darmstadt, Germany)
Hydrochloric acid, Merck (Darmstadt, Germany)
Isopropanol, Merck (Darmstadt, Germany)
Methanol, Merck (Darmstadt, Germany)

2.3 Culture media

Tryptone, Roth (Karlsruhe, Germany)
Yeast extract, Roth (Karlsruhe, Germany)
Agar-agar, Roth (Karlsruhe, Germany)
LB (Luria Bertani) media: 10g tryptone, 5g yeast extract and 5g NaCl, add 1 liter with distilled water added 12g agar for solid media.
TB (Terrific broth) media: 13.3g tryptone, 26.7g yeast extract, 4.4ml glycerol.
TB salt: 0.17M KH2PO4, 0.72M K2HPO4.

Before inoculation, the media were autoclaved and brought to room temperature. In the case of TB media, after autoclaving, TB salt was added.

### 2.4 Bacterial strains

The following strains were obtained from (Stratagene, La Jolla, CA, USA):
Escherichia coli BL21 (DE3) B F- dcm ompT hsdS (rB- mB-) gal(DE3)
Escherichia coli XL-1-Blue recA1, endA1, gyrA96, thi-, hsdR17, (rk-, mk+), supE44, relA1, lac-, [F’, traD36, proAB, lacIqZ_M15, Tn10 (tetR)]
Escherichia coli M15[pREP4] Escherichia coli TBK1 B F dcm ompT hsdS(rB mB) gal (DE3) [pTK Tetr]

### 2.5 Enzymes

Restriction endonucleases, New England Biolabs (Ipswich, USA) or MBI Fermentas (Ontario, Canada)
Lysozyme, Applichem (Darmstadt, Germany)
DnaseI, Applichem (Darmstadt, Germany)
Thrombin, Merck (Darmstadt, Germany)
2.6 Fluorescent labels

Oregon Green(r) 488 iodoacetamide, Invitrogen (USA)
Texas Red(r) C5 bromoacetamide, Invitrogen (USA)
Alexa Fluor(r) 594 C5 maleimide, Invitrogen (USA)

2.7 Instruments

AEKTA explorer, Amersham Pharmacia Biotech (Uppsala, Sweden)
Fluoromax-2, Jobin Yvon-Spex (Edison, NJ, USA)
Flurolog, Jobin Yvon-Spex (Edison, NJ, USA)
SMART chromatography sysyem, GE Healthcare VP-ITC, Microcal
UV-spectrophotometer, UV-2401 PC Shimadzu (Japan)
Luminescent image analyzer, LAS-1000 Fujifilm (Japan)

2.8 Constructs

The neuronal SNARE proteins used in the study were Syntaxin1a, SNAP-25A and Synaptobrevin2 and various deletion mutants as indicated in the results section. Norway rat (Rattus norvegicus) cDNAs, made from the transcript encoding these proteins, were used to make the expression constructs. The yeast SNARE proteins were soluble portion of Sec9, Sso1p, and Snc2 as well as transmembrane variants of Snc2 and Sso1. The SNAPs used in this study were αSNAP and mutants, βSNAP and Sec17. The AAA ATPases used in this study were chinese hamster NSF, Sec18, NSFY83E.

The following expression constructs were used:
• Syntaxin variants
  - H3 domain of Syntaxin 1a (residues 180-262),
  - full-length soluble Syntaxin (1-262).
  - H3 with the transmembrane region (residues 183-288),
  - Syntaxin with transmembrane region (1-288),
  - single-cysteine mutant (183-262 cys 225).

• SNAP25A variants
  - cysteine-less variant of SNAP25A (residues 1-206),
  - partial SNAP25A (residues 1-197),
  - partial SNAP25A (residues 1-188),
  - point-mutated SNAP25A (residues 1-206, M71A/I192A),
  - single-cysteine mutant (residues 1-202 cys 130).

• Synaptobrevin variants
  - partial Synaptobrevin (residues 1-96),
  - partial Synaptobrevin (residues 1-70),
  - partial Synaptobrevin (residues 1-116)
  - single-cysteine mutants (residues 1-96 cys 28 and residues 1-96 cys 61).

• Sec9 variants
  - partial Sec9 (residues 403-651),
  - single-cysteine mutant Sec9 (residues 403-651, cys 587)

• Snc2 variants
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- partial Snc2 (residues 1-93) with and without TMD,
- single-cysteine mutant Snc2 (residues 1-93 cys 24).

• Sso1
  - partial Sso1 (residues 179-264) with and without TMD.

• αSNAP variants
  - partial αSNAP (residues 1-295),
  - αSNAP L295A (residues 1-295),
  - αdelSNAP (residues 33-295) in pET28a vector (His tag).

• βSNAP (residues 1-298) in the pGex-2T vector (GST tag).

• Sec17p (residues 1-292) in a pQE vector (His tag).

• NSF variants
  - Chinese hamster ovary NSF (residues 1-744) in pET28a (His tag),
  - Chinese hamster ovary NSF (residues 1-744, Y83E) in a pET vector (His tag).

• Sec18p (residues 1-758) in a pQE vector (His tag).

All the SNARE proteins were in his-tag pET-expression vectors (Novagen).
Chapter 3

Methods

3.1 E.coli competent cells

The electrocompetent cells were made employing the following procedure; 10ml of overnight grown E.coli culture were added to 1L LB medium. The culture was grown at 37C in a shaker until an O.D. of about 0.5 - 0.7 was reached. The cells were later pelleted and resuspended for two consecutive times in 1L and 500 ml ice-cold HEPES buffer (1mM HEPES, pH-7.0), respectively. The cells were again pelleted and were resuspended in 20ml HEPES buffer with 10% (v/v) glycerol followed by pelleting and resuspension in 3 ml 10% (v/v) glycerol. They were frozen rapidly in liquid nitrogen. The cells were later stored at -80C. The heat-competent cells were made as described in [95].

3.2 Transformation

3.2.1 Electroporation

The electrocompetent cells were thawed on ice. Then 20ng of plasmid DNA were added and the reaction kept on ice for 10 min. The cells were then transferred to a 0.2cm electroporation cuvette (Bio-Rad). The electroporation was carried out with a gene pulser (Bio-Rad) with an 25F, 1.8kV, and 200 ohms pulse. After electroporation 800µl LB medium were added to the cuvette. The cells were transferred to a microfuge tube and were grown at 37C for 40 min in a shaker. Then 50 microliter of the culture were plated on an LB agar plate that contains the appropriate antibiotic for selecting the transformed cells. The plates were kept overnight at 37C to get transformed colonies.
3.2.2 Heat-shock

An aliquot of the heat-competent cells was thawed on ice and 30ng plasmid DNA were subsequently added. It was incubated for 20 min on ice. The cells were then subjected to heat shock by keeping them for 90 s in a water bath that was kept at 42°C. The cells were then kept on ice for 1 min until 500 microliter of LB medium were added. The cells were grown at 37°C for 40 min in a shaker. 50 microliter of the cells were plated on LB agar plates with the appropriate antibiotic and the plates were kept overnight at 37°C to get the transformed colonies.

3.3 Cloning of the αdelSNAP mutant

The αSNAP mutant αdelSNAP (residues 33-295 of αSNAP) was first amplified from αSNAP wild-type DNA using appropriate primers. Afterwards, the PCR product was cleaned up, cleaved with NdeI and XhoI and finally ligated into the NdeI XhoI cleaved empty pET28a vector (all according to the protocols available online at the Fermentas webpage). The sequence was confirmed by sequencing.

3.4 Protein expression and purification

All proteins were expressed as follows: LB medium was inoculated with the appropriate transformed colony of the E. Coli-respective strain. The cells were grown overnight in a shaker at 37°C. For expression, 20 ml of overnight grown culture were used to inoculate 1L of appropriate antibiotic containing TB or LB medium, the cells were grown until they reached an OD600 of 0.9. They were then induced to express the protein by the addition of IPTG (60 mg (w/v) per liter). The cells were grown for 3 hours at 37°C to allow expression of the recombinant protein.

The cells were pelleted and resuspended in 20 ml of resuspension buffer (20 mM Tris pH-7.4, 1 mM EDTA, 500 mM NaCl for the SNARES and the SNAPs, or HEPES pH-7.4, 500 mM NaCl, 0.5 mM ATP, 2 mM MgCl₂, for NSF, NSF²⁸³E, phosphorylated NSF and Sec18). The cells were lysed with the combination of lysozyme treatment and sonication. To reduce the viscosity of the solution DNase was added; PMSF (200 mM) was added to inhibit the activity of proteases. The lysate was centrifuged at 13,000g to pellet the cell debris. The supernatant was used for the affinity purification. All the proteins were soluble and contained a cleavable His- or GST tag for affinity purification.

Affinity Purification  Purification of His-tagged proteins (αSNAP, sec17, all SNAREs, all triple A ATPases): In the case of His-tagged proteins Ni²⁺-nitrilotriacetic acid beads (Qiagen) (1 ml/1L of expressed culture) were added for affinity purification. The beads were washed with excess of wash buffer 200 ml (20 mM Tris pH -7.4, 500 mM NaCl, and 8 mM imidazole (+ additionally 0.5 mM ATP, 2 mM MgCl₂ in case of the ATPases). The protein was eluted using two bead volumes of elution buffer.
(20mM Tris pH-7.4, 500mM NaCl, 400mM imidazole (plus additionally 0.5mMATP and 2mM MgCl₂ in case of the ATPases)). The His-tag was always cleaved during the subsequent dialysis by the addition of thrombin (100U per 12 liter of culture medium during expression).

Purification of GST-tagged proteins (βSNAP): For βSNAP, GST-beads (Qiagen, 2.5ml/6l of expressed culture) were added for affinity purification. The beads were washed with excess of wash buffer 200ml (20mM Tris pH -7.4, 500mM NaCl, 1mM DTT and 0.5mM EDTA). Subsequently, the beads were transferred to a falcon tube using approx. 5ml of washing buffer and the protein was directly cleaved on the beads by addition of Thrombin (100u) and incubation o.n. Using two bead volumes of wash buffer the cleaved protein was seperated from the beads via a column the next morning.

The affinity purification was followed by a 2nd step of chromatography on an AEKTA system (Amersham Biosciences).

Second step of Chromatography This second step of Chromatography was ion-exchange chromatography for all the SNAREs and all the SNAPs (isoforms, homologs and mutants) and gel-filtration for the ATPases (NSF, pNSF, and NSF^{YS3E}). Prior to ion exchange chromatography the SNAPs and SNAREs were dialyzed against a buffer containing 20mM Tris pH-7.4, 1mM EDTA, 1mM DTT and 50mM NaCl. The dialysis of the ATPases prior to gelfiltration contained 50mM HEPES, 0.5mM ATP, 2mM EDTA, 200mM NaCl, 2mM DTT and 10% glycerol. The same buffer was used for gelfiltration via a superdex200 column. Sec18 activity could not be reproducibly recovered after gelfiltration and Sec18 preparations were hence used directly after the Ni/NTA affinity procedure for experiments.

All the proteins obtained after these purification procedures were more than 95% pure as seen from the SDS-PAGE except from Sec17 and Sec18 (see results). All proteins were quantified either by their absorption at 280nm or using the Bradford assay (ATP interferes with the absorption at 280nm).

The yeast SNARE proteins were kindly provided by Xiong Chen and the transmembrane SNARE proteins by Ursel Ries and Alexander Stein.

in vitro phosphorylation of NSF For in vitro phosphorylation of NSF a special E.Coli strain was used. This strain, E.Coli Tkbl1, harbors a second plasmid which codes for a tyrosine kinase. NSF was first expressed like usual, except that tetracycline (12.5μg/μl) was included in all bacterial media to select for the plasmid harboring the kinase. After three hours of expression, cells were pelleted and resuspended in TK-induction medium as per the manufacturer’s instructions (Stratagene) Cells were then grown to an optical density of 0.5 (Abs600) and harvested as described above.

Assembly and purification of SNARE complexes. For the formation of a ternary complex the proteins Syntaxin (or its H3 domain): SNAP25 (or its mutants)
and Synaptobrevin (or its mutants) were assembled in a 1:1:1.5 molar ratio. They were incubated at room temperature for 2 hours before purification on a Mono-Q column to separate the unassembled monomers from the complex. The proteins were quantified using absorption at 280nm.

3.5 Gel Electrophoresis

For the SDS gel electrophoresis the protocol, as suggested by Laemmli.U.K [59] was used. After electrophoresis the stacking gel was discarded and the separation gel was fixed and stained for \( \sim 20 \) min in staining solution (50\% (v/v) methanol, 10\% (v/v) acetic acid and 0.2\% (w/v) Coomassie Brilliant Blue R-250) under agitation. The gel was destained for \( \sim 20 \) min and then in Destain 2 (5% (v/v) methanol and 12.5\% (v/v) acetic acid) until no background staining was visible. After scanning, the gel was dried in a gel dryer and preserved for further analysis.

3.6 Gel based disassembly

3.6.1 Disassembly of the minimal core complex

Purified minimal core complex (2.4M) was disassembled by addition of equal concentrations of NSF and a 14-fold excess of -SNAP, 2mM MgCl2, and 2.5mM ATP in 50mM HEPES, pH 7.4, 120mMKGlut/20mM KAc, for 1, 2, 5 and 10 minutes at 37\(^\circ\)C. The reaction was stopped by adding SDS sample buffer. As a control, the ATPase activity of NSF was abolished by replacing MgCl2 with 10 mM EDTA. All samples were analyzed by SDS-PAGE and Coomassie Blue staining. Unless heated, the minimal core complex (MC) runs as a single band of 34 kDa and thus can be separated from the monomeric SNARE components generated during disassembly.

3.7 Protein labeling

The single cysteine mutants were labeled using the sulphahydryl-reactive fluorophores Oregon Green 488 iodoacetamide, Texas Red C5 bromoacetamide, or Alexa Fluor 594 C5 maleimide (Molecular probes- Invitrogen). The proteins were labeled as per manufacturer’s instructions. After labeling the labelled protein was separated from the dye by size exclusion chromatography on a Sephadex G-25 column (Amersham biosciences). The labelling efficiency was later calculated according to the manufacturer’s instructions.
3.8 Fluorescence measurements

The fluorescence measurements were carried out in the Fluorolog-3 or Fluoromax-2 spectrofluorometer (Jobin Yvon). A fluorescence cuvette with magnetic stirrer, with a path length of 10 x 4 mm or with a pathlength 10 x 10mm (Hellma) was used in the experiments. If not stated otherwise in the figure legend, all the measurements were taken in buffer containing HEPES(50mM)/KGlue(120mM)/KAc(20mM), and an additional 0.5% of chaps in case of the FRET experiments. Whenever disassembly was monitored, the buffers additionally included 2mM ATP and 5mM MgCl₂ if not stated otherwise.

3.9 Fluorescence anisotropy

The fluorescence anisotropy of proteins labelled with Texas red or Alexa594 was measured with excitation and emission wavelengths, 590nm and 610nm respectively. For an Oregon green labeled protein, excitation and emission wavelengths of 488nm and 520 nm were used. The G factor was calculated using G = IHV/IHH. The anisotropy (r) was calculated using r = (IVV - G x IVH) / (IVV + 2 x G x IVH). Here, r refers to the fluorescence intensity, the first subscript is the direction of the exciting light and the second the direction of the emitted light.

3.10 FRET

Disassembly: Foerster resonance energy transfer (FRET) was monitored between proteins labelled with Oregon green and proteins labelled with Texas red. The labelled proteins were used at the concentrations indicated in the respective figure legends. The donor, Oregon green was excited at 488nm, the donor fluorescence and acceptor fluorescence (that of Texas red) were measured simultaneously at 520nm and 610nm respectively.

3.11 Preparation of proteoliposomes

Lipids (Avanti, Alabaster, AL) were mixed in chloroform to yield molar ratios as follows:

- Standard liposomes: phosphatidylcholine(5), phosphatidylethanolamine(2), phosphatidylserine(1) and cholesterol (1).
- PE/PC-liposomes: PE(1) to PC(4)
- PC liposomes: PC only

After drying, the lipids were resuspended in 20 mM HEPES/KOH pH 7.4 100 mM KCl, 5 mM DTT, 5% (w/v) sodium cholate at a total lipid concentration of 13.5 mM. SNARE-proteins in 1% chaps were added (lipid to protein ratio of 200:1 n/n); followed by size-exclusion chromatography on a PC 3.2/10 fast desalting column (GE Healthcare) with a sample/column volume ratio of 1:15, pre-equilibrated in 20 mM HEPES/KOH pH 7.4, 150 mM KCl, 1 mM DTT.
For the preparation of NBD/Rhodamine labeled liposomes 1.5% (n/n) 1,2-Dioleyl-sn-Glycero-3-Phosphoethanolamine-N-Lissamine Rhodamine B Sulfonyl, 1.5% (n/n) 1,2-Dioleyl-sn-Glycero-3-Phosphoethanolamine-N-(7-nitro-2,1,3-benzoxadiazol-4-yl) and 17% (n/n) phosphatidylethanolamine were used instead of 20% (n/n) phosphatidylethanolamine.

3.12 Fluorescence spectroscopy on liposomes

Disassembly 10 to 20 microliter of labeled liposomes were diluted in a total volume of 1.0 ml disassembly buffer (HEPES(50mM)/KGl Glu(120mM)/KAc(20mM), 2mM ATP and 5mM MgCl\(_2\) if not stated otherwise) resulting in final protein concentrations of approx. 35nM to 100 nM (indicated in the respective figures). FRET or fluorescence anisotropy were measured and normalized as described above and at the respective figures.

Detergent titration into liposomes 20µl of Nbd/Rho liposomes were diluted in a total volume of 1.0 ml buffer (HEPES(50mM)/KGl Glu(120mM)/KAc(20mM)/2mM ATP and 5mM MgCl\(_2\)). Dodecylmaltoside was then titrated to the reaction at steps of 62.5nM per addition and the emission spectra recorded. To excite the Nbd, an excitation wavelength of 460nm was used and the emissions of Nbd and Rhodamin were recorded at 538nm and 585nm, respectively.

3.13 Monoclonal antibodies

A monoclonal antibody was generated against the NSF N-terminal domain (residues 1-205). The rabbits were immunized with 300 g of protein emulsified in Freud’s adjuvant. It was followed by booster injections that were given every three weeks for 3 to 4 months. Eventually, the antiserum was obtained from the ear veins. Selection of clones was done in cell culture. Finally, the three single clones used for the experiments (141.1, 141.2 and 141.3) were amplified and concentrated using a bioreactor.

3.14 Western Blotting

For semi-dry western blotting the proteins were transferred to a nitrocellulose membrane after gel electrophoresis. The gel and the membrane were sandwiched between blotting papers (GB003 from Schleicher & Schuell). Before blotting, the blotting papers and the membrane were soaked in transfer buffer (25mM Tris/HCl pH-8.3, 193mM glycine, 20% (v/v) methanol, 0.1% (w/v) SDS). The transfer was carried out at 50mA constant current for 1 hour. After transfer, the membrane was either stored at 4C or it was directly proceeded with blocking. To the membrane, blocking solution (5 %( w/v) dry milk powder in PBS) was added and kept in a shaker for 30
min. The membrane was then washed three times, 10 min each, using the washing solution (0.1% Tween-20, 200mM Tris-HCl pH 7.7, 0.15M NaCl). After washing, the primary antibody, which was diluted 1000 times in blocking solution, was added to the membrane. It was either left overnight in a shaker at 4°C or kept in a shaker for 1 hour at room temperature. To remove unspecifically bound antibodies the membrane was washed three times, 10 min each, with washing solution. After washing, the secondary antibody that was diluted 2000 times in blocking solution was added. The membrane was kept in a shaker for 30 min at room temperature. The membrane was washed three times, 10 min each, with washing solution to remove the unbound secondary antibody. The secondary antibody was coupled to horseradish peroxidase which emits luminescence after the addition of chemiluminescence reagent (Perkin Elmer). The luminescence was detected using the luminescent image analyzer.

3.15 Cell Culture

Rat pheochromocytoma (PC12, clone 251) cells were grown in 75 cm² tissue culture flasks which contained 25 ml of growth medium - DMEM with high (4.5 g/l) glucose (Cambrex, New Jersey, USA). DMEM was supplemented with 10 % horse serum (Biochrom, Berlin), 5 % foetal calf serum Gold, 4 mM L-glutamine, 60 U/ml penicillin, 60 g/ml streptomycin (both from Cambrex) which all have been sterile filtered. For maintenance of a Mycoplasma-free cell culture, 6.25 g/ml of Plasmocin (InvivoGen, San Diego, USA) was present in cell culture throughout entire period of its growth. Cells were grown at 37°C in 10 % CO2 at 90% relative humidity. For passaging, one flask of confluent cells (approximately 240000 cells/cm²) was used. Growth medium was decanted followed by wash with 3ml of trypsin/EDTA solution (Cambrex). Cells were detached from their substrate with 3ml trypsin/EDTA. Trypsin activity on cells was blocked by addition of 27 ml of growth medium. Suspension of cells was centrifuged at 235 x g av for 5 min at 20°C (Varifuge 3.0R, Heraeus-Kendro Sepatech, Langenselbold, Germany). Medium was removed and the resulting cell pellet was titrated in 10 ml of fresh growth medium. Cells were diluted at 1:2 1:4, transferred into 75 cm2 uncoated tissue culture flasks (Sarstedt, Nuembrecht, Germany) and passaged every 48-96 hrs. To avoid any undesired age-dependent differentiation of culture, cells were used for only 20 passages after defrosting. For seeding on glass coverslips, cells were detached from their substrate as described above, titrated and diluted to give a final concentration of approximately 600,000 cells/ml. 500 l of this suspension (3 x 104 cells/cm2) were evenly plated onto each poly-L-lysine-coated coverslip. After plating, cells were transferred to incubator and allowed to settle onto the coverslips for at least 30 min. 3 ml of growth medium were then added. Cells were used for experiments 48-72 hrs after plating.
3.16 Disassembly on membrane sheets

Adherent cells are cultured on cover slips and subsequently subjected to a gentle ultrasound pulse of 100ms, which 'unroofs' the cells. Fluorescent cis-complexes were generated by reacting membrane sheets with Alexa594-labelled synaptobrevin 2 (lacking its TMR) that forms complexes with membrane sheet associated syntaxin 1 and SNAP-25. Membranes were then directly fixed or fixed after incubation for the indicated times in presence of variable concentrations of NSF and excess of $\alpha$–SNAP (2 µM), or variable concentrations of $\alpha$–SNAP and excess of NSF (40nM) as indicated in the respective figures. Remaining fluorescence after disassembly was related to the directly fixed value. For each time point 3-9 independent experiments were performed. Values are given as mean ± SEM.

3.17 Isothermal titration calorimetry

ITC was performed on a VP-ITC instrument (Microcal) at 25C. Samples were dialysed against degassed HEPES buffer (20 mM HEPES, pH 7.4, 150 mM NaCl, 1 mM DTT). Titrations were carried out by 10 ml injections. The stoichiometry was determined using a single-site model and measured heat released on binding was integrated and analysed with Microcal Origin 7.0 using a sequential-site binding model, yielding the equilibrium association constant $K_a$, the enthalpy of binding $\Delta H$, and the stoichiometry $n$. 
Chapter 4

Results

4.1 Basic In Vitro Characterization of the Disassembly Reaction

4.1.1 Purification of Active, Hexameric NSF

In order to investigate the disassembly reaction on a molecular level, at first all proteins involved had to be purified. As opposed to the neuronal SNARE proteins and αSNAP, whose purification was already well established in the lab, NSF turned out to be a challenging protein to purify in its active state. The original idea was to purify the histidin-tagged NSF via Ni/NTA- followed by anion exchange chromatography and finally separate the different homo-oligomers from monomers and one another.

Unfortunately, even though it expresses well, is soluble, eagerly binds to anion exchange materials and nicely elutes at a defined NaCl-concentration, it soon became evident that a major amount of activity is irreversibly lost during anion exchange chromatography rendering reproducible data collection impossible. Therefore, to optimize purification conditions, in addition to size- and purity-control via SDS-gel electrophoresis, NSF activity was tested after each step of purification using the FRET assay explained in more detail in section 4.1.4.

This led to the insight that Ni/NTA purification yielded best amounts of non-degraded NSF if pursued in presence of Mg\(^{2+}\)/ATP, a condition that allows for constitutive disassembly. A fast procedure and frequent buffer-exchange are mandatory to minimize degradation and retain activity, probably because constitutive ATPase activity can lead to limiting amounts of ATP, which have been described to cause NSF to fall apart into its monomers [8]. Monomers are presumably more accessible to bacterial proteinases than hexamers, and subsequent to longer purification procedures, bands at the size of approximately 25kDa and a 60kDa can be witnessed after SDS-PAGE. These bands coincide with those of the D1D2- and the N-terminal domain respectively.

Furthermore, anion exchange chromatography turned out to be inappropriate
for purification of NSF. The anion exchange step would have been advantageous because as a positive side-effect of purifying it is also suited to concentrate the sample. Instead, pure and hexameric NSF had to be purified via gel filtration chromatography (Superdex200) after dialysis following the elution from the nickel beads. Best results were obtained when the presence of Mg\(^{2+}\) and hence constitutive ATPase activity was kept as short as possible and ongoing degradation could be prevented by immediate dialysis against a buffer containing ATP and EDTA after elution from the Ni-column. Since NSF precipitates at low salt levels, NaCl concentration during purification should be kept above 175mM.

### 4.1.2 Monitoring Disassembly Via SDS-PAGE

An easy way to monitor the disassembly reaction exploits the fact that the ternary SNARE core complex is stable in SDS sample buffer, unless heated, and therefore appears as one single band after SDS gel electrophoresis [12]. Incubation of purified SNARE complex with NSF and αSNAP under disassembly conditions prior to gel electrophoresis will lead to SNARE-complex disassembly which can then be seen as a loss of intensity of the complex band and the appearance of additional bands representing the single SNARE proteins. This is illustrated in figure 4.1, where the kinetics of NSF disassembly are compared to that of a phosphomimetic mutant (NSFY83E) which has been reported to show a defect in αSNAP binding.

Even though this readout allows to differentiate between samples which show a dramatically different degree of disassembly, it is only semi-quantitative in the sense that the efficiency of disassembly can only be estimated by the intensity of the Coomassie stain on the gel. Secondly, the time resolution is too low to detect subtle differences. Furthermore, the reaction can not be monitored online, making the observation of reversible changes improbable. For example, differences in speeds of re-assembly between different SNARE targets, which can lead to apparently different amounts of disassembly, would presumably be misinterpreted to represent differences in disassembly. Hence, the next step was to establish a quantitative method with high time resolution which also allows to monitor the reaction online.

### 4.1.3 Online Monitoring of Disassembly Via Fluorescence Spectroscopy

Since fluorescence spectroscopy using fluorophore-labelled SNARE proteins is well established in the lab, the idea was to also monitor the disassembly reaction this way. Two types of experiments, exploiting different properties of fluorescence were set up. The first, FRET spectroscopy, makes use of the fact that energy transfer can take place between certain pairs of fluorophores, provided that the emission spectra of one (donor) overlaps with the excitation spectra of the other (acceptor). The amount of this so called fluorescence resonance energy transfer (FRET) is proportional to the inverse sixth power of the distance between the fluorophores, a property which
allows for monitoring of protein interactions. If the fluorophores are attached to
different, putatively interacting proteins under study, then upon interaction donor
emission will decrease while acceptor emission will simultaneously increase.

The second, fluorescence anisotropy, depends on the influence of rotational motion
on the fluorescence scattering of fluorophores in solution. It is proportional
to the ratio of parallel to perpendicular scattered light with respect to the plane of the
polarized excitation beam. Rotation of fluorophores shifts the plane of scattered
emission and hence changes the anisotropy. Since small particles rotate faster than
larger ones, the anisotropy can often be correlated to the size of the protein carrying
the fluorophore, and conclusively serve as an indicator of protein binding or complex
formation which slows down particle rotation.

To generate fluorescent substrates, maleimide-conjugated fluorophores can specifically
be attached to the sulphur atom of cysteines via a covalent bondage. Various single-cysteine mutants of the neuronal SNARE proteins were already available (e.g. [60]), providing good candidates to function as suitable substrates during fluorescence based disassembly assays.

### 4.1.4 The FRET Assay

At first, to find a FRET pair suited for disassembly, FRET pairs known to lead
to good signal changes during SNARE assembly were tested. An example is given
in figure 4.2. Since here we are looking at complex dissociation, i.e. the distance between donor and acceptor increases, donor fluorescence will increase upon disassembly of the SNARE complex. Effectively, we therefore observe ‘de-FRETting’ rather than ‘FRETting’ during disassembly. For simplicity, only the donor spectra are shown in the following.

The SNARE complex used here was generated by incubation of stoichiometric amounts of SNAP25, the soluble part of Synaptobrevin and the H3 domain of Syntaxin1A. The donor dye (Oregon green) was attached to the 28th position of Synaptobrevin (Sb\textsubscript{28OG}) and the acceptor (Texas red) to the 130th position of SNAP25 (SNAP25\textsuperscript{130TR}).

Figure 4.2: Monitoring disassembly of a core SNARE complex using FRET spectroscopy. Approximately 80nM of each FRET complex component (H3/SNAP25/Sb\textsuperscript{28OG}) were mixed with 1.8μM of αSNAP, 2mM ATP and NSF as indicated, and the reaction was started by addition of Mg\textsuperscript{2+}. (A) The half-time of the reaction correlates with the NSF concentration. (B) Addition of ∼400nM of unlabelled SNARE complex to the reaction results in a 5-fold decrease of apparent disassembly. This indicates that the labelled, ‘visible’ complex is disassembled at comparable speeds as the unlabelled (invisible) SNARE complex and hence disassembly is not impaired.

To ensure that the labelling of the SNARE complex does not influence disassembly kinetics of the wildtype SNARE complex, a five-fold excess of unlabelled SNARE complex was added to the reaction. Even though this unlabelled complex can not be observed in the spectrum, the apparent half-time of the labelled complex visible in the emission spectrum should increase by a factor of six if both complexes are disassembled equally well. The half-time indeed increased from 40 to 280s, indicating that the labelled complex is disassembled at comparable speeds as the unlabelled and hence labelling does not influence disassembly. Knowing this, various buffering conditions were tested for optimal rates of disassembly, eventually leading to the conclusion that the reaction dramatically slows down in presence of NaCl or KCl concentrations higher than 120mM, whereas a combination of KGlu/Kac at concent-
trations of 120mM and 20mM respectively still allows for efficient disassembly. The chelating agent EDTA, which depletes the reaction of Mg\(^{2+}\), can be used to stop disassembly. If the concentration of free SNAREs is high enough at this point, reassembly takes place. Furthermore the reaction is inhibited by high amounts of ADP and can be irreversibly stopped by addition of AlF\(_3\), which supposedly freezes NSF in a transition state. Likewise, NEM (N-ethyl-maleimide) blocks the disassembly reaction in this assay.

4.1.5 Fluorescence Anisotropy as a Readout for Disassembly

As for the FRET assay, different labelling positions were tested for their suitability to serve as the target in fluorescence anisotropy measurements. Since here the readout depends on the flexibility of the dye, which may or may not change upon interaction with other proteins, it was less predictable, which positions would be useful and which would not. Furthermore, as opposed to the FRET assay, interaction of the dye with components of the disassembly machinery might also lead to anisotropy changes undifferentiable from those caused by SNARE interactions. Setting up an anisotropy measurement, therefore requires a close look at stepwise anisotropy changes after addition of each respective component to understand which signal changes are actually caused by which interaction. As an example, the spectra of a SNARE complex labelled with Oregon green at the 28th position of Syb2 (Sb\(_{28}OG\)) during addition of \(\alpha\)SNAP and NSF as well as disassembly are shown in figure 4.3.

Note that adding \(\alpha\)SNAP causes a significant increase of fluorescence anisotropy before the reaction is even started, which can also be seen when an \(\alpha\)SNAP mutant known to be able to bind but not disassemble the SNARE complex, is added. The addition of NSF does not lead to a signal change, which is reasonable if one considers that enzymatic amounts of NSF are used. Upon triggering disassembly, the reaction including wt \(\alpha\)SNAP, but not the one including mutant \(\alpha\)SNAP, efficiently proceeds as can be told from a gradual decrease of fluorescence anisotropy.

4.1.6 Optimizing Protein Amounts for Fast Disassembly Kinetics

Having optimized purification- and buffer-conditions which lead to best NSF performance in the *in vitro* fluorescence assays, I subsequently determined how much of the components of the disassembly machinery are needed for optimal disassembly. From the stoichiometry of 20S complexes [61] one would assume that a molar ratio of NSF monomer to \(\alpha\)SNAP to SNARE complex of 6:3:1 should be used for the disassembly reactions. Anyhow, considering that NSF is an enzyme, substoichiometric amounts for NSF are probably more reasonable in a functional assay. For NSF it should hence be possible to find a range in which the amount of enzyme correlates with functional efficiency in a linear fashion. Similarly, \(\alpha\)SNAP could be sufficient.
Figure 4.3: Monitoring αSNAP binding and disassembly of a purified core complex using fluorescence anisotropy. 150nM of purified SNARE complex labelled at Sb position 28 (Sh28TR/H3/SNAP25) are disassembled in presence of 2mM ATP and 3,6µM of αSNAP. The addition of 1,2µM αSNAP or αSNAP\textsuperscript{L294A}, respectively, at the time point indicated leads to an increase of fluorescence anisotropy, whereas NSF addition at t=500s does not significantly influence the spectrum. Finally, adding MgCl\textsubscript{2} at t=540s leads to SNARE disassembly in the αSNAP wt-reaction, which can be observed as a gradual decrease of fluorescence anisotropy.

In enzymatic amounts, but might as well be required in a 3-fold excess with respect to the substrate, to saturate all complexes and thereby always keep the possible NSF acceptor-sites at a maximum. The reactions shown in figures 4.1.4 and 4.1.5 were conducted using excess amounts of ATP and αSNAP to ensure that NSF performance would not be underestimated due to limitations brought about by other components of the disassembly machinery. To open up possibilities to also employ the fluorescence assays for experiments investigating the reaction on the fuel- as well as the adaptor-level, it nevertheless is important to know as well, how much of these factors is required. Hence, NSF-, ATP- and αSNAP-dependence of SNARE disassembly were investigated under the as yet optimized buffer conditions.

The enzyme – NSF dependence of core complex disassembly in solution

In order to determine the concentration range of NSF at which it is able to efficiently disassemble SNARE complexes under the conditions used in the fluorescence assays, disassembly kinetics of approximately 130nM labelled SNARE complex using increasing amounts of NSF in presence of excess αSNAP (3,6µM) and ATP (2mM) were recorded. As shown in figure 4.4, as little as 1nM of NSF is sufficient
Figure 4.4: NSF dependence of core complex disassembly in solution. Disassembly kinetics of 130nM SNARE complex (H3/Sb28TR/SNAP25) using increasing amounts of NSF in presence of excess αSNAP (3,6µM) and ATP (2mM) were recorded. The arrow indicates the start of the reaction by addition of MgCl₂.

for optimal NSF performance. Up to 16nM enzyme, NSF-dependence is in a linear range, with speeds of disassembly of ∼15 pmol of SNARE complex per µg NSF per minute as calculated by the respective half-times. Due to ongoing re-assembly at later time points of the reaction caused by the increasing concentrations of free SNAREs, these rates are probably underestimates.

The fuel – How much ATP?

Subsequently, ATP-requirements for efficient SNARE disassembly under the conditions used in the fluorescence assay were investigated. To do so, disassembly kinetics of 75nM SNARE complex using increasing amounts of ATP in presence of excess αSNAP (1,25µM) and 10nM NSF were recorded. As shown in figure 4.5, the ATP concentration sufficient for optimal NSF performance is between 75 and 125µM.

The adaptor – αSNAP dependence of core complex disassembly in solution.

In order to determine the αSNAP dependence under the conditions used in the fluorescence assays, disassembly experiments with ∼90nM SNARE complex using increasing amounts of αSNAP in presence of excess ATP (2mM) and 10nM NSF were performed (figure 4.6). Notably, in line with published in vitro assays [62, 41], stoichiometric amounts of αSNAP did not suffice to disassemble the SNARE
Figure 4.5: ATP dependence of the reaction. Disassembly kinetics of ∼75nM SNARE complex comparing increasing amounts of ATP in presence of excess αSNAP (1,25µM) and 10nM NSF were recorded. The arrow marks the start of the reaction by addition of 5mM MgCl₂.

complex. On the contrary, between 420nM and 1,25µM of αSNAP are required for optimal speeds.

As will be seen in some later reactions, this may even be an underestimate due to the fact that at higher αSNAP concentrations the disassembly speeds reach the time resolution limit of the assay (1,25µM and 3,75µM have half-times of 15s).

As mentioned in the introduction, these are not likely to represent physiological amounts, evoking the need to clarify the reasons for the apparent lack of αSNAP-efficiency before any quantitative comparisons on the adaptor-level can be conducted.

4.2 How robust is the reaction?

It as yet remains unclear, which parts of the SNARE complex are important for disassembly and, with respect to the assumed three sites of αSNAP/SNARE interaction and even six sites of NSF/αSNAP interaction to which degree the machinery is robust in the sense that the specific interruption of single interaction sites only decreases reaction rates or leads to a complete abolishment of disassembly. If one found a way to abolish NSF/SNAP interaction by an alteration of either NSF or αSNAP, this would likely affect all SNAP/NSF interactions equally and consequently lead to a complete abolishment of SNARE disassembly. The interaction between SNAP and the SNARE-complex target however, is mediated by at least two and most likely
three structurally different sites, whose 'loss-of-interaction' effects can be addressed independently. The minimal SNARE regions required for SNARE disassembly [63] as well as mutants which may or may not inhibit SNARE disassembly have been described [64, 62, 65]. In the following section, some of these findings are being re-investigated and additional SNARE-complex alterations assessed, exploiting the increased time resolution of the fluorescence assays as compared to the approaches used in the original publications.

4.2.1 Inhibition of the Enzyme

The N-terminal domain of NSF has been reported to be responsible for αSNAP binding [66]. An antibody against this domain might therefore abolish SNAP-binding and hence disassembly, if its epitope overlaps with the critical area.

Generation of a functionally inhibitory antibody against the NSF N-terminal domain

To generate an antibody which blocks NSF-function, the purified NSF N-terminal domain (residues 1-205) was used for immunization. After expression in E. Coli the protein was purified via Ni/NTA- followed by cation-exchange chromatography. The pure N-domain was then given to Michaela Hellwig who used it for immunization of rabbits and subsequently raised monoclonal antibodies from these. After selection, the supernatants of different clones were tested in the FRET assay, one of which
turned out to block disassembly. This clone was then amplified and concentrated using a bioreactor and afterwards purified via ProteinA-sepharose by Michaela Hellwig. As shown in figure 4.7, 2.5 µl of antibody after elution from the ProteinA column prevent 1.5 nM of NSF from disassembling a soluble SNARE complex, which usually promote fast disassembly as shown in figure 4.4. Increasing the NSF concentration then allows to overcome the inhibition. The antibody could be shown to specifically recognize only NSF in rat brain cytosol as shown in figure 4.7 and none of the other proteins present in the disassembly assay. It can hence be concluded that binding of the anti-N antibody to NSF interferes with its function.
Figure 4.7: A monoclonal antibody directed against the NSF N-terminal domain inhibits SNARE-complex disassembly. (A) 2.5 µl of anti-N domain antibody (clone 141-2) were pre-incubated with 3nM NSF and ~40nM of FRET-SNARE complex (labels Syb^{28TR} and SNAP^{130OG}) in reaction buffer. Subsequently, 1.5µM of αSNAP were added and the reaction started by addition of MgCl₂ at t=0 seconds. Since no disassembly was observed, an additional 3nM of NSF were added after 175s, leading to the fast disassembly expected for 3nM NSF. (B) The indicated samples were blotted against clone 141-2 of the antibody to ensure specificity.
4.2.2 Influencing the Reaction on the Target Level

Full length soluble Syntaxin versus the Syntaxin H3 domain

All the experiments described so far were performed with the full soluble domains of SNAP25 and Synaptobrevin-2, whereas of Syntaxin1A only the SNARE motif (H3 domain) was used. The large N-terminal domain of Syntaxin1A known to be involved in the regulation of SNARE-complex assembly, was deleted assuming that it does not influence disassembly kinetics. To prove this notion right, disassembly kinetics of a ternary complexes with either the full-length soluble portion or the H3 Domain of Syntaxin1A were compared. Complexes were assembled using equimolar amounts of SNAP25\textsuperscript{130\textit{TR}}, Synaptobrevin\textsuperscript{28\textit{OG}} and Syntaxin or SyntaxinH3, respectively. After completion, disassembly was carried out as described above. Figure 4.8 illustrates that no major change in the disassembly rate was observed.

![Figure 4.8: The Habc domain of Syntaxin1A is not required for SNARE complex disassembly. SNARE complexes either containing the full-length cytoplasmic part of Syntaxin1A or only the Syntaxin SNARE domain (H3) were subjected to disassembly by NSF and αSNAP as before. The reaction was started at t=90s, the time point at which the beginning of a the gradual increase can be observed in both graphs.](image)

Various single or dual mutations in close vicinity impair SNARE disassembly only slightly, if at all

To elucidate whether certain regions of the SNARE complex are essential for disassembly, various SNARE complex mutants were to be investigated next.
Mutations in the SNARE-complex '0'-layer do not disturb SNARE disassembly

Figure 4.9: SNARE complexes with mutations in the '0'-layer are as prone to disassembly as wildtype complexes. (A) Pre-assembled FRET SNARE complexes (FRET pair Sb$^{28OG}$ and SNAP25$^{130TR}$ (80nM of each component) were mixed with a five-fold excess of unlabelled purified complex containing either Sb$^{Pro56}$ or wildtype Sb. Subsequently 1,9µM αSNAP and 4nM NSF were added and finally MgCl$_2$ at t=180s. (B) FRET-SNARE complexes assembled with either H3 or H3$^{Q226R}$ (FRET pair Sb$^{28OG}$ and SNAP25$^{130TR}$, 80nM of each component) are disassembled by 1,9µM αSNAP and 4nM NSF. The sample represented by the red graph additionally contained 2µM unlabelled Sb to render re-assembly invisible. MgCl$_2$ was used to trigger the reaction at t=220s. After 1100s EDTA was added to the wildtype reaction to show ongoing re-assembly.
First, SNARE mutations in the '0'-layer were investigated. For Synaptobrevin this meant a mutation of the arginine residue 56 to proline (Sb$^{R56P}$), the '0'-layer glutamine of Syntaxin was mutated to arginine. In section 4.1.4, mixing unlabelled SNARE complex with Sb$^{28OG}/$SNAP25$^{130TR}$/H3 FRET complex was used to demonstrate that the labelling does not impair disassembly. Now, the reaction from section 4.1.4 is shown in comparison to another one, which instead of wildtype complex included the same amount of Sb$^{Pro56}$-complex. Visible SNARE disassembly resulting from 'defretting' is as fast as for the wildtype reaction, indicating that both complexes are equally prone to disassembly and the Sb$^{Pro56}$ mutant conclusively does not impair disassembly.

To also re-investigate the Syntaxin '0'-layer mutant reported to cause a disassembly-defect in Scales et al. [64], H3$^{Q226R}$ was used for the second disassembly experiment illustrated in figure 4.9, where the disassembly kinetics of the mutation are compared to wildtype disassembly. Again, the reaction does not seem to be impaired, but what can be noted is some re-assembly very soon after disassembly is completed. This can not be seen in the wildtype reaction under these conditions, where EDTA needs to be applied to allow for visible re-assembly. In a third reaction, an excess of unlabelled Sb was added to the reaction to render re-assembly invisible. Here, the speed of disassembly seems slightly slower, which may be due to an increased invisible assembly-rate caused by the excess Sb which would increase target concentration and hence slow down the rate of visible disassembly towards the end of the reaction. Together, the slight kinetic differences observed do not support the notion that disassembly is changed significantly but might as well be explained by changes of one of the other interaction parameters, possibly an increased rate of re-assembly of the Sx$^{Q226R}$-mutation.
C-terminal and layer-5 mutations of SNAP25 do not inhibit SNARE disassembly

C-terminal deletions of SNAP25 and point mutations exchanging the two amino acids which SNAP25 contributes to the '5'-layer of the neuronal SNARE complex do not inhibit SNARE-complex disassembly. This was observed in experiments depicted in figure 4.10, which were carried out after the respective SNAP25 mutants were pre-assembled into complexes with a fluorescent dye attached to Synaptobrevin. In presence of αSNAP, NSF and ATP disassembly proceeds to the same extent at comparable speeds. To once more ensure that this is also true when compared to wildtype reactions labelled at a different position (which does not interfere with disassembly), the SNAP deletion mutants are shown in comparison to wildtype complex labelled with Texas red at Synaptobrevincys44.
Figure 4.10: Neither the disassembly of complexes containing C-terminally deleted SNAP25 nor that of SNAP25 with two point mutations in the 5th hydrophobic layer is impaired. (A) Approximately 200nM of pre-assembled SNARE complex labelled at Syb\textsuperscript{cys28} containing either SNAP25\textsuperscript{M71AI92A} or wildtype SNAP25 were disassembled by 1.5µM αSNAP, 2nM of NSF and 2mM ATP. MgCl\textsubscript{2} was used to trigger disassembly at t=850 s. The reactions were performed in presence of soluble unlabelled Synaptobrevin (2µM) to prevent reformation of labelled complexes. (B) Disassembly of different pre-assembled SNAP25-mutant SNARE complexes (Syb\textsuperscript{28TR}/H3/SNAP25 mutant as indicated, 200nM of each component) compared to disassembly of wildtype complex (Syb\textsuperscript{44TR}, H3 and SNAP25 wt). The reactions were performed in the presence of 1.6µM αSNAP, 4nM NSF and 2mM ATP and started by addition of MgCl\textsubscript{2} at t=265s. All traces represent the changes in fluorescence anisotropy during the reaction.
C-terminal mutations of Sb, Sx or both do not or partially inhibit SNARE disassembly

Next, C-terminal deletion mutants of SxH3 (183-240) and Sb(1-70) were scanned for disassembly defects. Sb deletion did not impair disassembly at all, whereas Sx-deletion led to a slight inhibition (see figure 4.11). The Sx mutant however did not reduce total amounts of disassembly, but only slightly impaired kinetics at the later stages of the reaction. Even when both, Sb and H3, were concertedly deleted, the inhibitory effect did not increase.

In summary, none of the various mutations examined in the last sections completely inhibited SNARE disassembly. Most deletions did not disturb disassembly at all. Partial impairment could only be seen when the C-terminal region of H3 was deleted, but even though the mutation slowed down disassembly kinetics towards the end of the reaction, all substrate present was sufficiently disassembled. Even the combination of two C-terminal SNARE deletion mutants (Sb and H3) did not increase the slight inhibition witnessed for H3 alone. One reason for this might be, that in all these cases only one SNAP/SNARE interaction site was disrupted. Hoping to finally abolish SNARE-complex disassembly completely, two non proximal regions were to be targeted simultaneously, as will be shown in the next section.
Figure 4.11: Disassembly of SNARE-complex deletion mutants. Different neuronal ternary FRET complexes, containing either an N-terminally shortened Synaptobrevin (1-70) or H3 (183-240) or both, were assembled (A) and subsequently disassembled in presence of 1.2µM αSNAP, 1.5nM NSF and 2mM ATP by addition of MgCl₂ at t=145s (B). The 'parts of disassembly' are defined as the fraction of donor signal increase at a given point divided by the change of donor signal observed during full assembly of the complex \((F_{ass\:complete})/(F_{dis\:complete}-F_{ass\:complete})\). \(F_{dis\:complete}\) and \(F_{ass\:complete}\) representing the donor fluorescence at complete disassembly or assembly, respectively.
4.2.3 Evidence for a Robust Target Recognition of the Dis-assembly Machinery

Inhibiting SNARE disassembly with the use of ternary complex-antibodies

A different approach to investigate whether SNARE disassembly can be blocked on the target level, is not to delete parts of the complex but rather attach a factor, which either hides one or more interaction sites or sterically hinders conformational changes required for disassembly. Good candidates for proteins, whose attachment might influence function, are antibodies.

![Graph showing FRET values over time with different antibodies](image)

Figure 4.12: Two antibodies against the ternary SNARE complex act as synergistic inhibitors of SNARE disassembly. Approximately 70nM FRET-SNARE complex (Sb$^{280G}$/SNAP25$^{130TR}$/H3) were incubated with the respective antibodies in disassembly buffer. Subsequently, NSF (1.5nM) and αSNAP (1.2µM) were added and the reaction started with MgCl$_2$ at t=-40s.

Three different antibodies raised against the purified SNARE 'mini'-complex (Dirk Fasshauer & Michaela Hellwig) had already been mapped to recognize different regions of the SNARE complex by Tabrez Siddiqui. Each of them alone as well as combinations of the antibodies were tested for their ability to inhibit SNARE disassembly using the FRET assay. As is shown in figure 4.12, all three antibodies partially impair disassembly, clones 131.1 and 131.2 having a slightly more pronounced effect than clone 131.5. Interestingly, if clones 131.1 and 131.2 are combined, they completely block disassembly. Considering that only half of the amounts used in the previous single-antibody experiments was applied in order to

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1Tabrez Jamal Siddiqui, *SNARE assembly and regulation on membranes*, PhD thesis
exclude a dosage dependent cause of the block, this finding indicates that the antibodies act synergistically, probably by interfering with the disassembly mechanism at two different sites of the complex.

**The combination of two non-inhibitory mutations can lead to inhibition of disassembly**

The last finding suggests that the NSF machinery might indeed be robust enough to function even though one of the interaction sites between substrate and machinery is weakened or blocked. Considering that the antibodies have been mapped to different regions of the complex, it is not likely that the two synergistic clones 131.1 and 131.2 interfere with the same site of interaction. This might indicate that even if one site is blocked, other sites might compensate for the defect and suffice to mediate disassembly. Similar to the observation made for the anti-complex antibodies, other structural alterations of the neuronal SNARE complex might also lead to inhibitions of function and maybe even show synergistic effects if they interfere with different sites of interaction. As will be shown in the next section, this synergy indeed turned out to be the case for two alterations, both of which did not exhibit any defect in disassembly on their own.

**Attachment of fluorescent dyes to Sb\textsuperscript{61} and Sx\textsuperscript{225} alone does not impair disassembly**

As pointed out in section 4.1.4, the FRET pair used throughout all experiments up to now (Sb\textsuperscript{28OG}/SNAP25\textsuperscript{130TR}) is equally prone to disassembly as wildtype ternary complex. This does not necessarily need to be true for all FRET pairs, since the attachment of labels at different positions might have a negative effect caused by steric hindrance or ablation of an important interface in any way. Hence, for every new FRET pair used, this possibility needs to be excluded first. Figure 4.13 therefore compares the speed of SNARE disassembly of a FRET pair which will be used in some of the following experiments (Sb\textsuperscript{61OG}/H3\textsuperscript{225TR}), to that of the one used hitherto. As can be seen, there is no significant difference between the disassembly kinetics of the two different FRET pairs.

**The FRET pair Sb\textsuperscript{61}/Syntaxin\textsuperscript{225} combined with C-terminal SNAP25 deletions severely impairs disassembly**

Notwithstanding the fact that each of the two alterations alone discussed in sections 4.2.3 and 4.2.2 did not influence disassembly kinetics, the combination of both severely impairs disassembly as can be witnessed in figure 4.14. Here, the kinetics of SNAP25 C-terminally deleted SNARE complexes carrying the FRET pair Sb\textsuperscript{61OG}/H3\textsuperscript{225TR} are significantly slower than the wildtype reaction. Furthermore, of the SNAP25\textsuperscript{layer5−}, SNAP25\textsuperscript{1−197−} and SNAP25\textsuperscript{1−188−} complexes no more than 10%, 20% or 30%, respectively, are successfully disassembled, compared to 75% in the wildtype reaction.
Figure 4.13: The two different FRET complexes Sb^{28OG}/SNAP25^{130TR}/H3 and Sb^{61OG}/H3^{225TR}/SNAP25) were disassembled in presence of 1,2µM αSNAP, 1,5nM NSF and 2mM ATP, triggered by addition of MgCl2 at t=160s. The parts of full disassembly are defined as the fraction of signal increase at a given point divided by the change of signal observed during full assembly of the complex (Fass_{complete}/(Fdis_{complete}-Fass_{complete}). Fdis_{complete} and Fass_{complete} representing the fluorescence at complete disassembly or assembly, respectively.

To exclude the possibility that the reactions with the SNAP25-mutations show a pronounced re-assembly, possibly due to slight inaccuracies with respect to the protein determination, the impact of an enhanced assembly and subsequent re-assembly was tested in further experiments, where the concentration of the respective SNAP25 was raised by a factor of four to promote faster assembly. As shown in figure 4.15, even though the acceleration has a subtle influence on disassembly, the differences between the mutant and the wildtype reaction remain much more prominent than those between the wildtype and the defective mutants.
Figure 4.14: Two structural alterations of the SNARE complex show a pronounced synergy. Different SNAP25 mutants (BotA mutant: residues 1-197, layer-5' mutant: M71A/I192A) or wildtype SNAP were assembled with their cognate SNAREs to form complexes carrying the FRET pair (Sb6^{OG}/H3^{225TR} introduced in section 4.2.3 (A) and subsequently disassembled in the presence of 1,2μM αSNAP, 1,5nM NSF and 2mM ATP by addition of MgCl2 at t=0s (B). The ‘parts of disassembly’ are defined as the fraction of signal increase at a given point divided by the change of signal observed during full assembly of the complex (F-F_{ass \_complete})/(F_{dis \_complete}-F_{ass \_complete}), F_{dis \_complete} and F_{ass \_complete} representing the fluorescence at complete disassembly or assembly, respectively.
4.2.4 Inhibition on the SNAP Level?

Having observed that certain structural changes on the substrate level can result not only in additive but even synergetic defects of disassembly, one may wonder how this can be explained on a mechanistic basis. One tempting speculation would be that these effects relate to the number of αSNAP bound to the complex during disassembly. Unfortunately, it is not possible to use stoichiometric – not to mention substoichiometric – amounts of αSNAP at this stage, which would be needed to test this hypothesis, because as pointed out in section 4.1.6, optimal disassembly requires enormous amounts of αSNAP in the fluorescence assays. It was therefore substantial to figure out first, how the efficiency of αSNAP could somehow be improved in our assays.
4.3 Why So Much αSNAP?

In order to finally be able to address questions related to αSNAP action, the following sections will try to shed light on why the αSNAP-performance is so unexpectedly inefficient in the fluorescence assays and how this nuisance can be resolved. In principle the low αSNAP efficiency could be inherent to the recombinant αSNAP, possibly due to invisible degradation or a lack of posttranslational modifications in vivo, which do not occur when expressed in E. coli. Alternatively, it might be caused by non-optimal conditions in the fluorescence assays. To test, whether the recombinant αSNAP shows any intrinsic defects, different methods were employed.

Figure 4.16: αSNAP binding to core complexes in solution. (A) Signal changes upon binding of increasing amounts of αSNAP to ~75nM SNARE complex were recorded and are shown as multiples of the anisotropy before αSNAP addition. Note that even at 1.25µM of αSNAP, saturation is not reached. (B) Mean values of the graphs on the left are plotted against the αSNAP concentration.

As outlined above, fluorescence anisotropy also allows to monitor αSNAP-binding. Figure 4.16 illustrates that for αSNAP binding the amounts needed to saturate the system are extremely high. Even at 1,25µM αSNAP the maximum signal change caused by αSNAP binding is not reached. The binding capacity saturates somewhere between 1,25µM and 3,75µM of added αSNAP (here relating to a ratio of approximately seventeen αSNAPs to one complex and fifty αSNAPs to one complex, respectively).

4.3.1 αSNAP Binding to the Complex Monitored Calorimetrically

The fluorescence experiments in section 4.1.6, investigating αSNAP-binding, indicated that more than 17 αSNAPs per complex were needed to achieve saturation. This finding dramatically differs from the 3:1 ratio one would expect and might be explained either by a reduced affinity or efficiency or alternatively by degradation, which renders a large portion αSNAP incompetent with respect to SNARE-complex binding. To check, whether the recombinant αSNAP binds the SNARE complex in
Why So Much αSNAP?

A stoichiometric ratio of 3:1 when used at higher concentrations, isothermal titration calorimetry was performed. The calorimeter detects temperature changes with such high sensitivity that it can be used to quantify the small amounts of heat released or consumed during protein/protein interactions. One protein is titrated into the other in distinct steps in an adiabatic jacket, and enthalpy changes as well as the stoichiometric ratio can be directly measured.

![Graph showing isothermal titration calorimetry of αSNAP binding to the core complex.](image)

Figure 4.17: Isothermal titration calorimetry of αSNAP binding to the core complex. αSNAP (70µM) in the syringe was titrated in ten steps into purified ternary SNARE complex (H3/SNAP25/Sb, 2.5µM) in the cell. Shown are the enthalpy changes during the experiment on top, and the integrated areas normalized to the amount of αSNAP (kcal/mol) versus the molar ratio of αSNAP to the SNARE complex below. The solid line represents the best fit to the data for a three site sequential binding site model using a nonlinear least squares fit.

As can be seen in figure 4.17, the enthalpy changes are very low obscuring the possibilities to extract much thermodynamical information. The stoichiometry 'n'
of the SNARE complex could nevertheless be determined to be 3 αSNAPs per complex in these experiments. Notably, this is exactly what is expected and hence allows us to conclude, that the majority of the recombinant αSNAP is able to bind SNARE complex, excluding the possibility that degradation is the reason for the low potency. Notwithstanding the low enthalpy values of the reaction, I still tried to fit the reaction in order to roughly estimate the αSNAP affinity. The thermodynamic data could not be properly fitted using a single site binding model, which assumes that αSNAP has the same affinity to all SNARE binding sites. Using a three-site sequential fit however, the curve could nicely be fitted (see figure 4.17). This suggests, that αSNAP binding to SNARE complexes is not equal to all three SNAP/SNARE interaction sites but the sites rather display different affinities. The affinities resulting from this fit were 10nM for the first, 50nM for the second and 380nM for the third binding site. Nevertheless, keeping in mind the low enthalpies and also considering that only a low number of data points per binding site were recorded, the affinities of αSNAP to the SNARE complex determined might not be very reliable.

4.3.2 αSNAP Dependence of SNARE Disassembly on 'Membrane Sheets’

Having established that αSNAP binds the SNARE complex at the expected ratio of 3:1, two possibilities remain: Either does the recombinant αSNAP have a reduced affinity as opposed to the endogenous protein, or a factor important for binding is missing in our fluorescence assay. Hence, the next step was to test the recombinant αSNAP’s efficacy under the more physiological conditions of a different system, the so called membrane sheets. These membrane sheet experiments were conducted together with Dana Bar-On.

Membrane sheets are generated by subjecting cultured, adherent PC12 cells to a gentle ultrapulse. The upper half of the cells is consequently ripped off, leaving behind an intact native membrane bilayer attached to a cover slip. The preparation provides the full set of membrane lipids and proteins in their native environment, including steric and conformational restriction, membrane fluidity, cytoskeletal components, lipid component diversity, and associated regulatory factors.
Figure 4.18: Disassembly of SNARE complexes on membrane sheets. After completion of complex formation, the membranes were incubated with increasing concentrations of NSF and αSNAP in the presence of Mg$^{2+}$/ATP, allowing for disassembly of the formed SNARE complexes. Upon disassembly the soluble fluorescent synaptobrevin is released into the buffer making the observed loss of fluorescence from membrane sheets a direct read-out of the disassembly reaction (100% represent total fluorescence before disassembly).

Cis-SNARE complexes in this experiment were generated by pre-incubation of membrane sheets with the fluorescent recombinant Synaptobrevin2 also used in the fluorescence assays. As in solution, the Synaptobrevin spontaneously assembles with its cognate t-SNAREs, here the endogenous Syntaxin and SNAP25 sitting on the PC12-cell membrane. After completion of complex formation, the membranes were incubated with increasing concentrations of NSF and αSNAP in the presence of Mg$^{2+}$/ATP(5mM), allowing for disassembly of the formed SNARE complexes. Upon disassembly, the soluble fluorescent synaptobrevin is released into the buffer making the observed loss of fluorescence from membrane sheets a direct read-out of the disassembly reaction. As illustrated in figure 4.18, fluorescence decreases with time in an NSF- and αSNAP-concentration dependent manner.

Surprisingly, under the conditions of the membrane sheet assay ~100nM of αSNAP are sufficient to mediate disassembly equally well as 2µM indicating saturation at 100nM. Of course the amount of SNARE target in these assays is not
known, not permitting a conclusion about the stoichiometry of αSNAP and SNARE complexes in this assay. Nevertheless, together with the ITC measurements which indicated a molar ratio of 3:1, these findings strongly suggest that a factor missing in the FRET and anisotropy experiments but present on membrane sheets is responsible for the higher αSNAP affinity.
4.3.3 Disassembly of SNARE Complexes Incorporated into Liposomes

Having excluded that the recombinant αSNAP differs from the endogenous in a way that causes the low SNAP efficiency, the reason for the low power in the fluorescence assays was to be searched elsewhere. Since the same recombinant NSF and also identical buffer conditions to the ones used in the fluorescent assays were used in the membrane sheet assay, these two factors could also be neglected. Presumably, an additional factor or domain potentiating the αSNAP function had to exist, which was present in the membrane-sheet assay but as yet not in the cuvette.

Possible candidates were the SNARE transmembrane domains as well as the plasma membrane itself. To check, whether the presence of either of these improves αSNAP efficiency, SNARE complexes were incorporated into liposomes prior to αSNAP/NSF mediated dissociation. And indeed, in the presence of liposomes 300nM of αSNAP suddenly sufficed to disassemble with the same kinetics as 1.5µM in solution in the FRET as well as the anisotropy assay (figure 4.19). Furthermore, it could be excluded that this was only due to low amounts of SNARE complex incorporated into the liposomes, because doubling the amount of complex as well as the amount of NSF still did not necessitate an increased αSNAP-concentration to finish disassembly in the same time-period as before. When disassembling liposomal complexes, αSNAP is hence no longer limiting at 300nM.
Figure 4.19: Liposomal SNARE complexes increase αSNAP efficacy. FRET spectroscopy (A) and fluorescence anisotropy (B) measurements of complexes incorporated into liposomes. Disassembly kinetics of SNARE complexes (Sb(1-116)$^{TR28}$/H3/SNAP25$^{130OG}$ and Sb(1-116)$^{TR28}$/H3/SNAP25, respectively) incorporated into liposomes were recorded. The reactions contained the αSNAP concentration as indicated, 6nM of NSF and 35nM complex. A third reaction shown in (A) also contained 300nM of αSNAP but 12nM NSF and 70nM complex. All reactions were started at $t=190s$ by addition of MgCl$_2$.
Figure 4.20: αSNAP dependence of SNARE-complex disassembly on liposomes using FRET spectroscopy. Disassembly kinetics of ~35nM FRET-SNARE complex (SbTMD_{28TR}/SNAP25_{130OG}/H3) with decreasing amounts of αSNAP in presence of 6nM NSF and excess of ATP (2mM) were recorded. Donor (A) and acceptor (B) fluorescence are shown. The reaction was started at t=180s by addition of MgCl₂. Note that αSNAP binding leads to an increase of acceptor signal in a concentration dependent manner prior to the start of the reaction. Note that between 60 and 120nM of αSNAP suffice for optimal disassembly.
Figure 4.21: αSNAP dependence of disassembly on liposomes using fluorescence anisotropy. Disassembly kinetics of ~35nM SNARE complex (SbTMD28TR/SNAP25/H3) using decreasing amounts of αSNAP in presence of 6nM NSF and excess of ATP (2mM) were recorded. The reaction was started at t=190s by addition of MgCl₂. Differences in fluorescence anisotropy prior to the start of the reaction result from different amounts of αSNAP bound to the complex (see figure 4.16). Note that again 120nM of αSNAP promote optimal disassembly.

To now determine the limiting amount for αSNAP under these improved conditions, decreasing amounts of αSNAP were used for disassembly until a concentration not capable of disassembling liposomal SNARE complexes was reached. This is illustrated in figure 4.20 using the FRET assay and in figure 4.21 using fluorescence anisotropy.

Together, these data show that αSNAP is capable of efficiently disassembling SNARE complexes at a stoichiometric ratio of three to one or even less, even at such low concentrations. The affinity was therefore significantly increased by the incorporation of SNARE-complexes into liposomes via the Synaptobrevin transmembrane domain.

Figure 4.22 shows the comparison of disassembly kinetics in solution and on liposomes using limiting amounts of αSNAP for the respective condition. Here one can see that 45nM of αSNAP on liposomes are comparably efficient as 1,1µM in solution, leading to the conclusion that αSNAP is at least 20-fold more efficient on liposomes than in solution.
Figure 4.22: αSNAP is roughly 20 times more effective on liposomes than in solution. Disassembly kinetics of ∼70nM SNARE complex (H3/SNAP25130OG/Sb28TR) either incorporated into liposomes via the Sb transmembrane domain or in solution were recorded. The reaction included 5nM NSF, αSNAP as indicated and excess of ATP (2mM). Disassembly was triggered after 120s by addition of MgCl₂.
Figure 4.23: NSF is equally potent on liposomes as in solution. Liposomal and soluble complexes (H3TMD, SNAP25\textsuperscript{130OG}/Sb) were disassembled in presence of 1,25\(\mu\)M \(\alpha\)SNAP, 6nM NSF and 2mM ATP by addition of MgCl\(_2\) at \(t=150\)s. (A) Traces represent the fluorescence anisotropy measured. (B) The solution signal has been shifted upwards to directly compare the kinetics of disassembly.

4.3.4 NSF Performance is Unchanged on Liposomes

To check whether NSF behaviour is changed on liposomes as well, disassembly at excess amounts of \(\alpha\)SNAP (1,5\(\mu\)M) was compared between liposomes and solution at identical NSF-concentrations. Figure 4.23 shows that NSF disassembles at comparable rates, regardless of whether the SNARE complex is incorporated into liposomes or not, as long as \(\alpha\)SNAP is not limiting.

Furthermore, the same amount of anti-N-domain antibody (141.2) is needed to block NSF function on liposomes as in solution. This is illustrated in figure 4.24, where the anisotropy of liposomal SNARE complexes is recorded in presence or absence of 2,5\(\mu\)l of the ProteinA-eluted antibody 141.2. When NSF and the antibody have time to interact prior to \(\alpha\)SNAP addition, disassembly is completely blocked. The block is nevertheless relieved upon addition of a further 3nM of NSF, identical to what was observed in solution in section 4.7.
Figure 4.24: As in solution, 2.5µl of the anti-N-domain antibody block 3nM of NSF. Black trace: Approximately 35nM of SNARE complex (H3/SNAP25/SbTMD<sup>28TR</sup>) were mixed with 3nM of NSF and 2.5µM of clone 141.2. After 50s, 3µM of αSNAP were added and anisotropy recorded. The disassembly reaction was subsequently triggered by addition of MgCl₂ at t=0. Red trace: Everything as above except that αSNAP and the antibody were added together, prior to NSF, to take away the antibodies advantage of being pre-incubated with NSF.

Monitoring NSF-binding using stoichiometric amounts

Figure 4.25: NSF-binding and speed of the reaction using stoichiometric amounts of NSF. (A) Red graph: 150nM αSNAP were added to ∼35nM of liposomal SNARE complex (SbTMD<sup>28TR</sup>/H3/SNAP25) at t=-600s. Subsequently NSF was added in a stepwise manner as indicated by black arrows. Finally, the reaction was started at 150s by adding MgCl₂. Black graph: Everything was performed as for the red graph, except that αSNAP was added at 20s and 15nM of NSF were added in a single step at 80s. (B) Close-up of the disassembly time interval between 120 and 200s shown on the left to illustrate the high speed of the reaction.
To see, whether the binding of NSF to the SNARE-complex can be noted in the spectrum as well if stoichiometric instead of enzymatic amounts of NSF are added, increasing amounts of NSF were titrated to a complex already saturated with αSNAP (150nM). This is illustrated in figure 4.25 (A), where each addition of NSF is indicated by a black arrow. An increase of anisotropy is observable using such high amounts of NSF, but it is not large enough to assess stoichiometry. As can be seen on panel (B), at higher NSF concentrations the rate of disassembly soon becomes too fast to be resolved with this experimental set-up. The apparent half-times of disassembly here are 5 and 8 seconds, which probably is an underestimate since this time period is in the range of the time required for efficient mixing of the reaction solution.

A time resolution high enough to obtain a spectrum of one disassembly cycle using stoichiometric amounts of the SNAREs, SNAPs and NSF might be achieved using a stopped-flow apparatus. Some preliminary experiments have already been performed using such a set-up, pointing to a speed of less than four seconds per cycle at room temperature (data not shown). Since NSF has been observed to disassemble significantly faster at 37°C, an even faster rate of disassembly can be expected at that temperature. Here, it should nevertheless be stated again that these *in vitro* experiments cannot provide absolute numbers with respect to physiological processes but only serve as a platform to pursue comparative investigations.
4.4 Membrane Anchorage of SNAREs Potentiates the Power of αSNAP

Looking at the experiments explained in the last section it became evident that much less αSNAP is needed for efficient disassembly using complexes incorporated into liposomes via the transmembrane domain (TMD) of Synaptobrevin. Nevertheless it is as yet still unclear, whether the liposomes or the TMD of Synaptobrevin are the factor causing the improved function.

4.4.1 The Transmembrane Domain of Synaptobrevin is not Essential for αSNAP Potentiation

To test whether the Synaptobrevin transmembrane domain increases αSNAP function, SNARE complexes lacking the Synaptobrevin transmembrane domain were incorporated into liposomes via the Syntaxin transmembrane domain. As judged from my previous findings, 300nM αSNAP are not capable of mediating sufficient disassembly in solution but are sufficient for efficient disassembly on liposomal complexes incorporated via the Sb-transmembrane domain.

To now determine, whether liposomal complexes incorporated via the Syntaxin- and hence missing the Sb-transmembrane domain mediate disassembly at 300nM αSNAP, populations of complexes with a TMD of either H3 or full length Syntaxin were disassembled by two different concentrations of αSNAP (300nM and 1.5µM, respectively).

Notably, as shown in figure 4.26, 300nM αSNAP were not limiting during complex disassembly. This can be concluded from the fact that the speed of disassembly at the two concentrations used was identical. Furthermore, the binding of both amounts of αSNAP results in a comparable increase of fluorescence anisotropy, once more indicating that the system is saturated at 300nM αSNAP. Moreover, the disassembly kinetics were still identical when the NSF concentration was raised from 1.5nM to 6nM, indicating that even at higher NSF concentrations 300nM of αSNAP are not limiting as opposed to what would be expected in solution.

Indeed, performing the same experiment in solution (figure 4.27) confirmed that the lower αSNAP concentration here only allows for slow disassembly of the soluble complexes. The slow disassembly may not even be improved by raising the NSF concentration. Disassembly is hence improved on liposomes at low αSNAP levels even in the absence of the Synaptobrevin TMD, showing that it is not the TMD of Synaptobrevin which leads to the observed increase of αSNAP affinity on liposomes.
Figure 4.26: 300nM αSNAP are not limiting even in the absence of the Synaptobrevin transmembrane domain. (A) Full-length Syntaxin complexes (SxTMD/SNAP25<sup>130OG</sup>/Sb) were pre-assembled and incorporated into liposomes via the Syntaxin transmembrane domain. (B) Complexes containing Syntaxin without Habc domain (H3TMD/SNAP25<sup>130OG</sup>/Sb) were pre-assembled and incorporated into liposomes via the Syntaxin transmembrane domain. The 35nM complex per reaction were dissociated using 1.5nM or 6nM of NSF and 300nM or 1.5µM of αSNAP and started at t=190s by addition of MgCl<sub>2</sub>. Notably, the binding of both amounts of αSNAP results in a comparable increase of fluorescence anisotropy (t=50s).
Figure 4.27: 300nM of αSNAP are limiting using the same amount of soluble complexes. The procedure and conditions were identical to figure 4.26, except that soluble complex (H3/Syb/SNAP25130QG) not incorporated into liposomes was disassembled. The 35nM complex per reaction were dissociated using 1,5nM or 6nM of NSF and 300nM or 1,5µM of αSNAP and started at t=190s by addition of MgCl₂. Note that the anisotropy shift caused by αSNAP binding (at t=30s) is far more pronounced for the binding of 1,5µM than for that of 300nM αSNAP.
4.4.2 The Potentiation of $\alpha$SNAP Efficacy is Independent of the Lipid Composition

Now knowing that liposomes increase $\alpha$SNAP efficacy, it remained to be elucidated whether this effect is caused by a specific lipid component. The liposomes up to this stage were constituted of phosphatidylcholine (PC), phosphatidylethanolamine (PE), cholesterol (Chol), phosphatidylserine (PS), phosphatidylinositol (PI) in order to closely resemble the lipid composition of the brain. To repeat the experiments with a simplified lipid composition, the liposomes used in the following experiment were constituted of PC and PE only. Even this simple lipid composition did not alter the efficiency of $\alpha$SNAP (figure 4.28). As shown here, 60nM of $\alpha$SNAP were sufficient to drive the reaction, which is comparable to the findings for the more complex liposomal preparations used in the previous experiments. It turned out that even pure phosphatidylcholine liposomes (data not shown) suffice to support the increased potency of $\alpha$SNAP. The effect is hence caused not by any special lipid but rather independent of the composition of lipids.

Figure 4.28: Liposomes consisting of PC and PE only are sufficient to bring about the observed higher efficacy. The experiment was performed as in figure 4.21 except that cholesterol, PI and PS were omitted from the lipid mixes used to generate the liposomes. The reactions were started at $t=190$s.
4.4.3 Disassembly on Partially Solubilized Liposome Populations (Mixed Micelles)

To ensure that it is not the mere presence of lipids in the environment (which e.g. might stabilize fragile regions of αSNAP and thereby promote a positive effect), but indeed the anchorage of the complex to the membrane, the following strategy was employed: First, it should be investigated how much of a respective detergent is needed to solubilize only a fraction of the liposomes, leaving the rest intact. Afterwards, disassembly was to be tested at different ratios of micelles to liposomes caused by different amounts of detergent. Under such conditions, the amount of disassembly in presence of low αSNAP should correlate with the fraction of complexes still incorporated into liposomes, whereas dissociation of the remaining complexes should only proceed when the αSNAP concentration would be increased to levels sufficient for disassembly in solution.

Disassembly in Presence of Dodecylmaltoside

To determine the detergent concentration required for partial solubilization, detergent was titrated into liposomes that contained 1.5% NBD (Nitrobenzoxadiazole)- and 1.5% Rhodamine-labelled lipids, respectively, two dyes which can function as a FRET pair. With increasing detergent concentrations, FRET efficiency between the lipids will at first be slightly perturbed by insertion of detergent molecules into liposomes and at some point dramatically decrease due to solubilization of the liposomes, which obviously leads to an increased distance between the lipids. Several detergents were tried, the biggest difficulty being the circumstance that most detergents, when used at concentrations that solubilize liposomes, harm the efficiency of the disassembly reaction as such. These are consequently not suited for experiments comparing disassembly at different detergent concentrations. Dodecylmaltoside (DOM) however, a detergent which due to its low critical micelle concentration (cmc) succeeds to solubilize liposomes already at low concentrations, turned out not to have a negative impact on disassembly activity in the respective concentration range.

In figure 4.29 the increase of donor fluorescence during DOM-titration into labelled liposomes is shown and the fraction of solubilized liposomes estimated. Below, disassembly is performed on mixed micelles, representing a mixture of liposome-attached and detergent-solubilized FRET-labelled SNARE complexes at the respective detergent concentrations. For the first step of disassembly, enough αSNAP (i.e. 120nM) to disassembly liposomal but not soluble complexes according to what we learned in section 4.3.3 was applied. After stabilization of the signal subsequent to this first part of disassembly, a high concentration of αSNAP suited to disassemble complexes in solution was added and the remainder of disassembly proceeded. Figure 4.30 lists the correlation between the fraction of liposomes expected to exist and the amount of disassembly proceeding at the low αSNAP concentration under the respective concentration of detergent.
Figure 4.29: The percentage of disassembly requiring low αSNAP correlates with the fraction of liposomal complexes in mixed micelle solutions. (A) The donor fluorescence during titration of DOM into Nbd/Rho-labelled liposomes was recorded, used to estimate the fraction of solubilized liposomes (\(\% \text{ solubilized liposomes} = \frac{F-F_{\text{start}}}{F_{\text{end}}-F_{\text{start}}}, F_{\text{start}}\)) and plotted against the concentration of DOM. (B) Disassembly is performed on mixed micelles at DOM concentrations of 125, 375 and 500nM, respectively. After 160s, 120nM αSNAP were added to allow for disassembly of liposomal SNAREs. At t=440s, a further 1.5µM of αSNAP were added to also allow for disassembly of the soluble SNARE complexes.
Figure 4.30: The percentage of disassembly requiring low αSNAP correlates with the fraction of liposomal complexes in mixed micelle solutions.
4.4.4 An αSNAP Mutant Lacking the Putative Membrane Interaction

The data collected so far made it tempting to speculate that a direct interaction between αSNAP and the membrane lipids is responsible for the increased efficacy on liposomes. If this were true, the lipid binding property might possible be mapped to a certain region of αSNAP. No crystal structure has been solved for αSNAP so far, but the structure of the αSNAP isoform γSNAP as well as the yeast αSNAP homolog Sec17 have been solved [30]. Based on the Sec17-structure and αSNAP/SNARE-complex interaction studies using various point-mutated αSNAPs, a model of αSNAP bound to the SNARE complex [67] has been proposed by Marz et al. Looking closely at the model one can observe that the very N-terminal 32 residues form an arm-like structure pointing away from the complex, the most far-out region of which includes a strand of mostly hydrophobic amino acids. If this region were the interaction site of αSNAP with the membrane, its deletion should abolish membrane binding and hence the higher αSNAP efficiency on membranes.

![Figure 4.31](image-url)

Figure 4.31: Purification of the αSNAP mutant ’adelSNAP’. A sample after elution from the Ni column, αSNAP wildtype for size comparison as well as the elution fractions after anion-exchange chromatography are shown. Fraction 11 was used for all of the functional experiments.

A deletion mutant comprising aa 33-295 of αSNAP was hence cloned into the pET28a vector and subsequently expressed and purified. It will be referred to as ’adelSNAP’ from now on. The purification of the His-tagged protein was carried out by Wolfgang Berning-Koch via Ni/NTA- followed by anion-exchange chromatography. It worked according to the standard protocol used for αSNAP 4.31.

First, I tested whether adelSNAP is able to disassemble SNARE complexes using the previously described FRET assay. Indeed, adelSNAP is able to disassemble –
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![Graph showing disassembly kinetics](image)

Figure 4.32: \( \alpha \)delSNAP efficiency is comparable on liposomes and in solution. Disassembly is shown \( \sim 70nM \) of FRET complexes (H3/SNAP25\(^{130OG}/Sb^{28TR} \)) either in solution or incorporated into liposomes via the TMD of Sb. At the start of the reaction the samples contained the amount of \( \alpha \)del indicated. The reactions were triggered after 140s. 1.5\( \mu \)M of \( \alpha \)delSNAP were added twice more to the reactions represented in black and green, at 400s and 680s.

albeit at lower efficiency than \( \alpha \)SNAP. At an \( \alpha \)delSNAP concentration of 1.5\( \mu \)M, disassembly was slow and only a third of the substrate was disassembled. In order to improve the rate of disassembly, more \( \alpha \)delSNAP had to be added. 4.4\( \mu \)M succeeded to efficiently disassemble most of the complex (figure 4.32).

Next, to address the question of whether the membrane boost observed for wt-\( \alpha \)SNAP in the previous sections is indeed abolished in the mutant, the kinetics of \( \alpha \)delSNAP mediated disassembly in solution and on liposomes were compared. As illustrated in figure 4.32, for all concentrations of \( \alpha \)delSNAP tested the kinetics in solution are identical to the kinetics on liposomes. This result indicates that the membrane mediated \( \alpha \)SNAP-potentiation was indeed abolished by deletion of its N-terminal region.

To strengthen the finding that \( \alpha \)delSNAP does not differentiate between liposomal and soluble complexes with respect to disassembly efficiency, the anisotropy read-out was employed next and \( \alpha \)delSNAP directly compared to wildtype \( \alpha \)SNAP. Since in the FRET-experiment it already became obvious that \( \alpha \)delSNAP is less effective than wildtype \( \alpha \)SNAP in general, the extent of this difference was determined at first. To do so, various amounts of \( \alpha \)delSNAP and \( \alpha \)SNAP were compared with respect to their ability to disassemble SNARE complexes in solution. As shown in figure 4.33, 3.6\( \mu \)M of \( \alpha \)delSNAP disassemble complexes with the same kinetics as 600nM \( \alpha \)SNAP. Likewise, 1.8\( \mu \)M of \( \alpha \)delSNAP are as efficient as 300nM \( \alpha \)SNAP. It can hence be concluded that \( \alpha \)delSNAP is six times less efficient than \( \alpha \)SNAP in solution (figure 4.33).
Results

Figure 4.33: αdelSNAP is less efficient than αSNAP in solution. The fluorescence anisotropy changes upon disassembly of purified SNARE complexes (H3/SNAP25/Sb28TR) with either αSNAP or αdelSNAP as the NSF/SNARE-adaptor protein are shown. The signal at a given time is divided by the signal prior to the adaptor-addition (A/A0). The reaction was triggered after 120s as indicated.

Subsequently, the efficiencies of αdelSNAP and αSNAP with respect to disassembly were compared on liposomes. First, I employed the concentrations of αdelSNAP and αSNAP which led to comparable kinetics in solution. On liposomes, 600nM of αSNAP disassembled the SNARE complexes significantly faster than 3.6µM of αdelSNAP (figure 4.34). At lower αSNAP concentrations, which are more appropriate for αSNAP disassembly on liposomes, the differences between αdelSNAP- and αSNAP-mediated kinetics at a ratio of 6:1 are even more pronounced: 280nM αdelSNAP do not permit any disassembly at all, whereas 45nM of αSNAP mediate fast and complete disassembly (figure 4.35). Even after increasing the concentration of αdelSNAP up to 2.3 µM the rate of αdelSNAP mediated disassembly was slower than that of the wildtype reaction (data not shown). These results are in line with the outcome of the FRET assay, that αdelSNAP does not preferentially disassemble on liposomes.
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Figure 4.34: The difference between αdelSNAP and αSNAP is far more pronounced on liposomes than in solution. Anisotropy changes during disassembly of ~70nM liposomal SNARE complexes (H3/SNAP25/Stb28TMD) at 3.6μM αdel or 600nM αSNAP in the presence of 5nM NSF. Disassembly was triggered at 130s by MgCl2. Anisotropy was normalized to parts of full disassembly to ease quantification.

Figure 4.35: The low αdelSNAP efficacy on liposomes is even more obvious at low SNAP concentrations. Anisotropy traces of SNARE-complex disassembly (H3/SNAP25/Stb28TMD) at 280nM αdelSNAP or 45nM αSNAP in the presence of 5nM NSF. Disassembly was triggered at 125s by MgCl2.

Theoretically, from the findings we have so far, one can predict the factor by which wt-αSNAP should be more efficient than αdelSNAP during SNARE-complex
Results

Figure 4.36: 45nM αSNAP disassemble liposomal SNARE complexes at a comparable speed as 3.6 μM of αdelSNAP. Anisotropy traces of liposomal SNARE-complex disassembly (H3/SNAP25/Sb28TRTMD) at 3.6μM αdel or 45nM αSNAP in the presence of 5nM NSF. Disassembly was triggered at 125s by MgCl2 as indicated.

Disassembly on liposomes. We have seen in this section that αdelSNAP is approximately six fold less efficient than αSNAP in solution, and in section 4.3.3 that the efficiency of wildtype αSNAP is increased by a factor of 10-20 in the presence of membranes. According to this, an efficacy factor in the range of 60-120 (roughly six times ten to twenty) would be expected on liposomes. Indeed, figure 4.36 illustrates that 45nM αSNAP disassemble liposomal SNARE complexes at similar speed as 3.6 μM αdel or 45nM αSNAP, indicating an 80-fold higher efficiency for αSNAP. These findings strongly suggest that the N-terminal arm-like structure observable in the crystal structure of αSNAP is essential for a protein/lipid-interaction, which mediates the potentiation of complex disassembly at low αSNAP concentrations.

4.4.5 Potentiation Through Membrane Anchorage – General Feature of SNAPs or αSNAP-Specific Phenomenon?

The adaptor – αSNAP versus its brain specific isoform βSNAP

As described in the introduction, αSNAP is not the only adaptor protein of NSF. Two more αSNAP isoforms are known, called β- and γSNAP, respectively. γSNAP is ubiquitously expressed like αSNAP, whereas βSNAP is a brain specific isoform. Up to date there have been contradictory results about βSNAP function and its ability to disassemble SNARE complexes in particular and it yet remains unclear whether βSNAP functions as a positive or negative regulator or simply provides redundancy despite relatively low sequence homology [32, 33, 31, 34].
Membrane Anchorage of SNAREs Potentiates the Power of αSNAP

With the following experiments I sought to answer, whether βSNAP is capable of disassembling SNARE-complexes and if so, whether its properties differ from those of αSNAP. For instance, it is as yet not known, whether αSNAP has the same affinity for all three binding sites and whether all three binding sites need to be occupied to allow for disassembly. If the SNAP binding sites are not exchangeable there might be differences of β- and αSNAP affinities to any of the binding sites. If so, it is for instance feasible that α- and βSNAP are more or less efficient in different concentration ranges, respectively and that mixing of both SNAP isoforms would lead to different kinetics than the sum of both isoforms alone. Furthermore, an interesting question would be, whether the membrane boost observed for αSNAP on liposomes is conserved between SNAP isoforms.

The properties of α- and βSNAP are similar in solution To first establish, whether βSNAP disassembles neuronal SNARE complexes in general, I purified recombinant βSNAP and used it for disassembly experiments exploiting the fluorescence anisotropy.

As can be seen in figure 4.37, βSNAP was able to disassemble the SNARE complex. Like previously observed for αSNAP, the binding of βSNAP to the SNARE-complex results in an increase of fluorescence anisotropy. All experiments were also performed with αSNAP as a direct comparison. It should be noted that βSNAP turned out to be fragile during purification, making it impossible to estimate the true power of βSNAP as compared to αSNAP. The following experiments are hence not intended to provide absolute numbers but only aim to investigate whether there are mechanistical differences between the SNAP isoforms.

As is shown in figure 4.37, 3.4μM βSNAP disassembled the complexes with the same kinetics as 600nM αSNAP. The βSNAP preparation used was hence about four times less effective than αSNAP at this concentration. To see, whether this difference is reproducible at much lower concentrations, a second experiment using only 600nM βSNAP and 150nM αSNAP was carried out. Again, the kinetics were comparable. Further experiments were pursued using different β- and αSNAP concentrations, all of which lead to comparable kinetics of α- and βSNAP-containing reactions, when for times more βSNAP preparation than αSNAP was used (data not shown). Furthermore, mixing of limiting amounts αSNAP and βSNAP resulted in a reaction rate which would be expected for the sum of the two isoforms (data not shown). None of these solution-experiments hence pointed towards a different function of βSNAP and αSNAP, rather suggesting that both isoforms are mechanistically exchangeable in solution. Whether the efficiencies of the isoforms differ, can not be judged from these experiments, as pointed out above.
Figure 4.37: βSNAP is approximately four times less effective than αSNAP during SNARE-complex disassembly in solution. Anisotropy changes during disassembly of ∼70nM FRET-complex (Sb^28TR/H3/SNAP25) at high adaptor-protein concentrations (as indicated in A) and low adaptor-protein concentrations (as indicated in B) with 6nM NSF are shown. The reactions were triggered by MgCl_2 at 780s (A) or 650s (B).

The membrane boost observed for αSNAP is conserved for βSNAP  To now address the question of whether the potentiation of αSNAP efficiency by incorporation of the complexes into liposomes is conserved for βSNAP, disassembly of the two SNAP isoform was compared on liposomes as well. In order to exclude that putative differences between liposomes and solution are caused by different qualities
of the preparations used, the same preparations of $\alpha$SNAP and $\beta$SNAP as in the last experiment were used and reactions carried out on the same day. As illustrated in figure 4.38, 45nM of $\alpha$SNAP disassembled almost as efficiently as 160nM $\beta$SNAP. The rates of disassembly in presence of $\alpha$- or $\beta$SNAPs are thus comparable when about four times as much $\beta$SNAP as $\alpha$SNAP was used like previously observed in solution. The fact that the efficiency-difference between the two SNAPs is the same on liposomes as in solution, indicates that the efficiency of $\beta$SNAP on liposomes increases to the same extent as has earlier been determined for $\alpha$SNAP. The positive effect of complex incorporation into liposomes hence is conserved amongst at least two of the SNAP isoforms, namely $\alpha$SNAP and $\beta$SNAP.

Figure 4.38: $\beta$SNAP versus $\alpha$SNAP on liposomes. 160nM $\beta$SNAP or 45nM $\alpha$SNAP were pre-incubated with $\sim$70nM FRET-complex (H3/SNAP25$^{130\text{OC}}$/Sb$^{28}$Alexa594TMD), NSF and ATP and the disassembly initiated at $t=180s$ by MgCl$_2$. 
4.5 How is the Reaction Being Regulated?

The newly discovered SNAP potentiation through membrane anchorage now opens up possibilities to investigate putative regulatory mechanisms of SNARE disassembly which act on the level of the SNAP adaptor protein.

4.5.1 Influence of Complexin1 on Core-Complex Disassembly

Besides SNAP isoforms, putative competitors of αSNAP with respect to SNARE-complex binding have been suggested, one of these proteins being Complexin1 (Cpx1). Complexin has originally been reported to replace αSNAP from SNARE complexes and meanwhile its crystal structure bound to the SNARE complex has been solved. Gel-based experiments investigating the effect of Complexin on SNARE disassembly have been performed [41] but did not confirm the proposed inhibitory role. Anyhow, as mentioned earlier, such experiments only have a limited time resolution and are thus mainly suited to trace 'all-or-nothing' defects. Therefore, I re-investigated the Complexin influence on SNARE disassembly using the fluorescence assays. First, as shown in figure 4.39, I tested whether Complexin can influence SNARE disassembly in the first place. Even though both, αSNAP and Complexin, were used at very high concentrations compared to the SNARE substrate (3.6μM and 2μM, respectively at 90nM complex), Complexin was able to slightly inhibit SNARE complex disassembly. The extend of inhibition increased, when the αSNAP concentration was reduced to levels which make αSNAP limiting when disassembling in solution (1.1μM). As a result, the half-time of the reaction increased by a factor of five from 20s to 100s. Complexin was added first, to allow for binding, in all cases.

To now estimate the ratio of Complexin- to αSNAP-concentration at which Complexin is able to modulate αSNAP-function in solution, increasing amounts of αSNAP were titrated into a FRET complex under disassembly conditions, either in presence or absence of 150nM Complexin1 (data not shown). Comparison of the two resulting donor spectra indicated that the sample containing Complexin was less efficiently disassembled at equal αSNAP concentrations up to an αSNAP concentration of 625nM. A more than four-fold excess of αSNAP was hence needed to fully compensate the inhibition. Figure 4.40 illustrates that 370nM Complexin are sufficient to strongly inhibit SNARE disassembly at an αSNAP concentration of 1.1μM.
Figure 4.39: Influence of Complexin on core complex disassembly. Fluorescence anisotropy was recorded during SNARE-complex disassembly (90nM of purified SNARE-core complex labelled with Texas red at $\text{SH}_{28}$) either with or without $2\mu M$ of Complexin1 (1-134). $3.6\mu M$ (A) versus $1.1\mu M$ (B) of $\alpha$SNAP are compared. Proteins were added as follows: (A) Complexin was added after 240s, and can be seen as a small increase in the green graph. Subsequently, $\alpha$SNAP was added to both reactions at 320s. Note that both reaction show a similar increase, independent of whether they contain Complexin or not. Finally NSF was added at 380s and $\text{MgCl}_2$ at 420s. (B) Complexin was added after 40s, $\alpha$SNAP after 520s, NSF after 580s in the red graph; $\alpha$SNAP at 500s and NSF at 550s in the red graph. The reactions were triggered by $\text{MgCl}_2$ at 620s.
Figure 4.40: Complexin is dominant over αSNAP in FRET experiments in solution. Disassembly at 1,1μM αSNAP was triggered after 140s in presence or absence of 370nM Complexin. The reactions included 4,5nM NSF and ∼75nM complex (H3/SNAP25130OG/Sb28TR).

Figure 4.41: Influence of Complexin on disassembly of liposomal complexes. FRET spectra of SNARE complex (H3/SNAP25130OG/Sb28TR-TMD) disassembly were recorded at 45nM αSNAP and various concentrations of Cpx1.

If αSNAP and Complexin really were to compete in the cell, performing competition experiments in presence of the membrane would naturally be more reasonable. These might lead to a completely different outcome than in solution, depending on whether the membrane influences Complexin’s efficiency during disassembly in a similar way as αSNAP or not. An additional advantage of investigating the Complexin/SNAP interplay on liposomes is the possibility to employ stoichiometric amounts of αSNAP. As pointed out earlier, this is not possible in solution experiments, because αSNAP needs to be employed in a μM range even at low substrate
How is the Reaction Being Regulated?

Figure 4.42: Full block of disassembly via an excess of Complexin at very low amounts of αSNAP (22.5nM). The donor fluorescence of FRET experiments looking at disassembly of 22.5nM αSNAP in presence of various amounts of Cpx1 is shown. All reactions were mediated by 5nM of NSF and triggered by MgCl₂ at 120s. Note that disassembly is blocked completely at 370nM of Complexin.

Figure 4.41: Competition experiments were also performed on liposomes as shown in figure 4.41. To this end, SNARE complexes were incorporated into liposomes and disassembled in the presence or absence of Complexin1. Here, 75nM of Complexin partially inhibited disassembly mediated by 45nM αSNAP, and 7.5µM of Complexin inhibit disassembly almost completely. Nevertheless, the inhibition by 75nM Cpx1 is overcome by raising the αSNAP concentration to 90nM. Even the more pronounced inhibition caused by 7.5µM of Cpx1 is partially recovered by 90nM αSNAP and almost completely relieved when αSNAP concentration is raised to 1.1µM.

Interestingly, Complexin completely blocked SNARE disassembly when very low amounts of αSNAP were used (22.5nM) as illustrated in figure 4.42. Yet, this block mediated by 370nM Cpx1 was partially relieved, as soon as the αSNAP concentration was increased to 45nM (A). Interestingly, the extent and the rate of this reaction are identical to the rate in presence of 75nM Cpx1 and 22.5nM αSNAP, maybe indicating that the extent of inhibition is confined to discrete levels. On the contrary to what has been observed in solution, 450nM αSNAP (in other words a 30% excess of αSNAP with respect to Complexin) then suffice to completely relieve the block. A significantly higher ratio of Complexin:αSNAP is hence needed on liposomes than in solution, to successfully impair disassembly.
Figure 4.43: αSNAP is dominant over Complexin on liposomes. FRET SNARE-complex and NSF concentration are identical to figure 4.42, the Mg$^{2+}$-trigger was added after 130s. (A) Complexin/αSNAP ratios of 3:1 (75nM Cpx:22.5nM αSNAP) and 6:1 (370nM Cpx:45nM αSNAP) inhibit disassembly to the same extent. (B) Addition of 450nM αSNAP at t=200s to the reaction containing 370nM Cpx1 completely relieves the inhibition.

Altogether, the data suggest that αSNAP function is not as fragile with respect to Cpx1 as in solution, where even a 3-fold excess of αSNAP could not recover normal disassembly speeds. This is probably due to the fact that much lower amounts of αSNAP are needed for disassembly on membranes compared to solution. The affinity of Complexin hence does not seem to vary between liposomal and soluble complexes. Complexin therefore has a selective advantage over αSNAP in solution.

### 4.5.2 Does Phosphorylation of NSF Have an Impact on Function?

The competition between αSNAP and Complexin discussed in the last section affects the interface between the SNAPs and the SNARE complex. An additional level of regulation is feasible at the interaction sites between the SNAPs and NSF. For example, phosphorylation of NSF at a tyrosine at position 83 has been reported to reduce NSF’s affinity for αSNAP [36]. This hypothesis is best to be tested under conditions which allow for the use of limiting amounts of αSNAP, making the fluorescence assays on liposomes ideally suited.

To this end, two NSF mutants were expressed and purified: First, a wildtype construct of NSF was expressed in an E. coli strain in which proteins can be in vitro-phosphorylated directly inside the bacteria during expression. The strain contains a second plasmid coding for a kinase whose expression is under control of a tryptophane operon and can hence be induced independently. Expression of the kinase is triggered by changing the expression medium to a medium containing indole-acrylic-acid (IAA) as an inducing agent after three hours of NSF production. The kinase then phosphorylates NSF which can afterwards be purified according to the protocols used for unphosphorylated NSF. Phosphorylation was subsequently confirmed...
by Western blotting (data not shown). Using this approach the degree of phosphorylation can however not be determined. Therefore, we also employed a second approach to have a control which should behave like NSF phosphorylated to saturation: A phosphomimetic point-mutant (NSF$^{Y83E}$) of NSF was expressed, which was purified according to the protocols for wildtype NSF as well. During purification of NSF$^{Y83E}$ it became evident that the monomeric peak was relatively higher, whereas the hexameric was relatively smaller compared to wildtype NSF preparations (data not shown). This may have a physiological background, but could as well simply be due to an increased fragility of the mutant under the conditions used during purification.

As shown in section 4.1.2, the phosphomimetic mutant indeed performed less efficient than wtNSF in the gel-based disassembly assay. However, as mentioned in section 4.1.2 as well, this approach does not allow to differentiate between incomplete disassembly and partial re-assembly. The interpretation of such a finding is thus more difficult than in the ‘real-time’ fluorescence approach.

Consequently, both of these mutants were tested in the FRET assay, where they behaved similarly. Therefore, disassembly on liposomes is exemplified only for NSF$^{Y83E}$ (figure 4.44). Liposomal SNARE complexes were incorporated either via the Sb- or the H3-transmembrane domain and afterwards disassembled by either wildtype NSF or NSF$^{Y83E}$. For both complexes the mutant performed slightly less efficient than the wildtype. In case of the Sb-TMD complex, disassembly was only slightly slowed down whereas for the H3-TMD complex, only half of the complex was disassembled by the mutant at all. Increasing the mutant’s concentration by 50% anyhow led to disassembly comparable to, or in case of the H3-TMD complex even better than, the wildtype indicating that the difference was not very pronounced.

On liposomes, the differences between wildtype and phosphorylated NSF can be diminished by increasing the mutant’s concentration by 50%, a difference which
might as well result from reduced stability of the mutant. At this stage it should again be pointed out that NSF is known to be a fragile enzyme and it is difficult to compare the preparations of the wildtype and the point mutation quantitatively, since it can never be excluded that differences between the enzymes stem from buffering conditions which, even though optimized for NSF as described in section 4.1.3, do not mimic the enzymes natural environment.

To next investigate, whether the reported phosphorylation-mediated defect of NSF function is dependent on the Habc domain of Syntaxin, the experiments were also performed with liposomal complexes containing full-length Syntaxin, again once incorporated via the Sb- and once via the Syntaxin-transmembrane domain. Again, using 1.5-fold the amount of mutant sufficed to disassemble at comparable speeds as the wildtype, indicating that the Sx Habc domain does not influence the reaction (see figure 4.45).

Figure 4.45: NSF^{Y83E} efficiently disassembles liposomal complexes containing full-length Syntaxin. Everything was carried out as in figure 4.44 except that complexes contained full-length Syntaxin (including the Habc domain) (Sx/SNAP25^{130OG}/Sb^{28TR}) (A) or the Sx-TMD (B), respectively.

To clarify, whether the disassembly differences observed in the gel-based assay only show up in solution, the FRET assay was carried out in solution next. As shown in figure 4.46, phosphorylated NSF once more disassembled slower and less efficiently than wildtype when used at identical concentrations. As opposed to the liposomal findings however, this deficiency could not be overcome by adding more phosphorylated NSF. Here, it is therefore less likely that the imperfect disassembly by pNSF is due to a lower concentration or stability. Due to the reported defect in αSNAP binding, I subsequently tested whether disassembly would improve in presence of more αSNAP. Indeed, the defect is almost completely abrogated by increasing the αSNAP concentration from 1,1µM to 2,2µM. Whether this is due to a reduced αSNAP affinity of NSF^{Y83E} in solution remains to be elucidated. Further experiments hence need to be performed in order to get further insights into whether the phosphorylation indeed leads to a mechanistical difference of the mutant or whether the mutant is simply less stable in my hands than the wildtype.
Figure 4.46: pNSF performance at 1,1µM αSNAP is less efficient in solution. Everything was carried out as in figure 4.44 except that soluble complexes and NSF and αSNAP as indicated were used. The reaction was triggered by MgCl₂ after 130s. The Black arrows indicate further additions of pNSF after equilibrium has been reached.
4.6 How Conserved is the Reaction Mechanism?

4.6.1 SNARE Disassembly Using Yeast Homologues

The fact that αSNAP and NSF are ubiquitously expressed in all tissues and mediates the recycling of most if not all intracellular SNARE proteins involved in intracellular fusion processes suggests that the mechanism of the different disassembly reactions throughout the cell is strongly conserved. Considering that the four-helix-bundle structures of the different SNARE complexes show a high degree of similarity, the overall sequence homology between the different complexes studied thus far is surprisingly low. It is hence possible that the SNAP/SNARE machinery does not care about conservation of specific residues but rather recognizes its target proteins on the basis of their quaternary structure. To elucidate, whether even between two organisms as distant as yeast and mammals the degree of conservation is high enough to provide functionality of one machinery in the other organism, components of the mammalian machinery were tested on yeast SNARE complexes and vice versa. Subsequently disassembly experiments were performed using various combinations of the yeast and the mammalian components, to determine which of the parts or interfaces are especially conserved and whether the mechanisms found for the neuronal SNARE complex (as e.g. the ’membrane boost’) are conserved in yeast.

4.6.2 Disassembly of Neuronal Complexes Using the Yeast Machinery

To this end, the yeast homologues of αSNAP and NSF (Sec17 and Sec18, respectively) were expressed and purified according to the protocols used for their mammalian counterparts. Unfortunately, Sec18 activity could not be recovered after gel filtration. Furthermore, the ratio between monomeric and hexameric Sec18 after gel filtration varied drastically between preparations and was shifted towards monomeric enzyme with respect to the average NSF preparation. This could indicate that Sec18 is either less stable than NSF under the purification conditions used or that the equilibrium is naturally shifted to the monomeric (i.e. inactive) fraction in case of Sec18. Anyhow, since activity was not recovered after gel filtration, Sec18 was directly used after Ni/NTA-chromatography for all of the following experiments. Sec17 did not express well and still contained impurities after purification. Furthermore, it soon became evident that both, Sec17 as well as Sec18, tend to loose activity very rapidly (in a day or two from purification). Sec17 was therefore also employed directly after elution from the Ni-beads, omitting the second chromatographic step. All experiments using the yeast machinery, therefore do not aim to determine absolute activities or rates of disassembly. Even though the true efficiency of the yeast machinery can not be determined that way, qualitative studies can be carried out. As will be shown in the next section, some questions regarding the degree of conservation of the disassembly mechanism can be resolved merely on the basis of comparative measurements.
'Component Swapping’ between the yeast and the mammalian reaction

First, all possible combinations (αSNAP/NSF, Sec17/Sec18, αSNAP/Sec18 and Sec17/NSF) were tested for disassembly neuronal SNARE complexes. αSNAP and NSF were most efficient, followed by the complete yeast machinery (Sec17 and Sec18) and the combination of Sec18 and αSNAP, which worked equally efficient. As mentioned above, this does not allow to draw the conclusion that the yeast machinery is less efficient than the mammalian in general, but what can be seen from this experiment is that the yeast machinery is capable of disassembling neuronal complexes. Furthermore, the efficient disassembly mediated by αSNAP and Sec18 indicates that the interface and basic mechanism on the level of the adaptor-enzyme interplay seems to be conserved (figure 4.47). To ensure that Sec17 is not limiting, a second reaction (data not shown) using a four-fold excess of Sec17 was performed, which led to the same kinetics. What is also striking in this experiment is the fact that the combinations mammalian complex/mammalian adaptor/yeast enzyme and mammalian complex/yeast adaptor/yeast enzyme, both containing one interface between non-cognate proteins, display similar kinetics. Interestingly, the combination of Sec17 and NSF did not promote disassembly of the neuronal complex at all, even though used at the same concentrations as in the other combinations which all led to efficient disassembly (see figure 4.47). Spontaneously, two scenarios are feasible: Either the interface between Sec17 and NSF has diverged so much during evolution that the two cannot productively interact and disassemble at all. Alternatively, the connection could only be weakened but still be able to function, as long as the cognate substrate for Sec17 were provided. This would result in a combination containing only one interface between non-cognate proteins and might successfully disassemble like the other combinations with one incorrect interface. When disassembling the non-cognate neuronal complex however, disassembly might already be hampered by an imperfect interaction between Sec17 and the neuronal complex and then completely be abolished by the second impeded interaction between the non-cognate adaptor and enzyme.

The membrane boost is conserved for the yeast adaptor protein Sec17

To now investigate, whether the membrane-caused potentiation of efficiency found for αSNAP is conserved for the yeast adaptor protein Sec17, further experiments using more Sec17 were carried out, in order to see whether or not the combination of Sec17 and NSF could not be brought to function at least partially. If this were to be the case, the same amounts of Sec17 to NSF could be applied to disassemble SNARE complexes in solution, and the kinetics compared. Indeed, the combination of Sec17 and NSF mediated slow and partial disassembly when used at high amounts. Using the same amounts afterwards to disassemble soluble complexes was not successful (Figure 4.48). It can thus be concluded that Sec17 efficacy and in analogy to αSNAP probably affinity are higher when the target complexes are incorporated into membranes.
Figure 4.47: All except one combination of the yeast and the mammalian disassembly machinery efficiently disassemble neuronal SNARE complexes. Approx. 40nM of SNARE complexes (H3/Sb^{28TMD}/SNAP25^{130OG}) were disassembled with the adaptor/enzyme-pairs as indicated. Concentrations used were 600nM for αSNAP, 6nM NSF and roughly estimated (from the gel) to be 350nM for Sec17 and 200nM for Sec18 after Ni/NTA-elution (hence still including the monomeric fraction). The reaction was triggered by MgCl$_2$ at 630s. The arrows indicate αSNAP addition which results in a small increase of donor signal. This rise is not observed for the addition of Sec17.
Figure 4.48: The ‘imperfect’ adaptor protein Sec17 also shows membrane dependence. Roughly 700nM Sec17 (as estimated from the gel) and 1.5nM NSF were used to disassemble 40nM of liposomal SNARE complexes (H3/Sb28TR/TMD/SNAP25130OG). The time point of Sec17-application as well as the start of the reaction by addition of MgCl2 are indicated.
4.6.3 Disassembling the Yeast SNARE Complex

As stated in the last section 4.6.2, recombinant Sec17 and Sec18 are less efficient in disassembling the neuronal SNARE complexes than NSF and αSNAP. Considering the fact that their activities are also less reproducible between preparations than those of the mammalian machinery, the lower efficiency is presumably at least partially caused by a lower intrinsic activity. This might be improvable by further optimization of the purification protocols. Nevertheless it is impressive that the yeast machinery disassembles the neuronal complex at all, considering that this do not represent its cognate target. It would hence be interesting to know, whether the yeast machinery performs better when disassembling yeast SNARE complexes. Likewise the question arises, whether the target recognition of the mammalian machinery is robust enough to disassemble yeast SNARE complexes as well. To address these questions, yeast SNARE complexes were assembled and subsequently disassembled by either the mammalian or the yeast disassembly machinery. The yeast SNAREs Sec9 (residues 403-651), Sec9<sup>cys<sup>587</sup>TR</sup>, Snc2 (residues 1-93) with and without TMD, Snc2<sup>cys<sup>24</sup></sup> as well as Sso1 (residues 179-264) with and without TMD were expressed, purified and kindly provided by Xiong Chen.

The machineries do not preferentially disassemble their cognate complexes

As illustrated in figure 4.49, both machineries are able to disassemble both complexes. Again, as has been observed for the neuronal complex earlier, the yeast machinery disassembles less efficiently than the mammalian, even when it comes to disassembling its cognate complex. Whether this is only due to a lower stability in the purification buffers or also has a physiological meaning remains unclear at this point. Interestingly, the kinetics do not vary significantly with respect to whether the respective cognate or non-cognate SNARE complex is disassembled. From this one can deduce that SNARE-complex recognition by the disassembly machinery indeed is either very robust or very conserved between yeast and mammals.

‘Component swapping’

Finally, the disassembly components were also ‘swapped’ as previously shown for the disassembly of the neuronal SNARE complexes. Knowing from section 4.6.2 that Sec17/NSF do not perform well together when disassembling neuronal complexes, using the yeast SNAREs opens up the possibility to further elucidate the reasons for this incompatibility. If Sec17 and NSF simply do not fit together, they would be expected to perform least efficiently of all combinations during yeast-complex disassembly as well. If on the other hand the reason for their imperfect performance is due to a synergistic defect caused by two non-cognate binding sites as discussed in section 4.6.2, they would be expected to disassemble yeast SNARE complexes more efficiently.
Figure 4.49: Disassembly of yeast vs. neuronal SNARE complexes. Approximately 40nM of either yeast (Sec9$^{578OG}$/Sec2$^{24TR}$/Sso1TMD) or neuronal (SNAP25$^{130OG}$Sb$^{28TR}$TMD/H3) SNARE complex were disassembled by the machinery as indicated. 90nM of $\alpha$SNAP and $\sim$3nM of NSF were used and the reaction triggered at 450s as marked. The amounts of yeast machinery used were identical between the reactions and roughly estimated (by appearance on the gel) to be 250nM for Sec17 and 100nM for Sec18 (still including monomeric fraction).

Figure 4.50: All except one combination of the yeast- and the mammalian disassembly machinery efficiently disassemble yeast SNARE complexes. (A) Anisotropy changes upon disassembly of $\sim$40nM of yeast SNARE complexes (Snc2TMD/Sec9$^{9TR}$/sso1p) with the adaptor/enzyme-pairs as indicated. Concentrations used were 120nM for $\alpha$SNAP, 6nM NSF and roughly estimated to be 350nM for Sec17 and 100nM for Sec18 from Ni elution (hence still including the monomeric fraction). The reaction was triggered by MgCl$_2$ at 220s. (B) $\sim$40nM of yeast SNARE complexes (sso1pTMD/Sec9/Snc2$^{TR}$) were disassembled by 120nM $\alpha$SNAP and either 6nM NSF or 100nM Sec18 as indicated.

As shown in figure 4.50, the second theory seems to be true. Sec17/NSF medi-
ated yeast disassembly is not severely hampered during yeast-complex disassembly. On the contrary, they rather efficiently disassemble the yeast SNARE complex like the combinations Sec17/Sec18 and αSNAP/NSF (4.50). Strikingly, the combination of αSNAP and Sec18 however does not mediate disassembly of the yeast complex under the same conditions used in the other experiments. Note that this combination (yeast<sub>complex</sub>/mammalian<sub>adaptor</sub>/yeast<sub>enzyme</sub>) correlates to the least efficient combination in the last section (mammalian<sub>complex</sub>/yeast<sub>adaptor</sub>/mammalian<sub>enzyme</sub>) in the sense that both contain two interfaces between non-cognate proteins. In summary it can be concluded that both of the protein interfaces between target, adaptor and enzyme seem to be conserved well enough between yeast and mammals to allow proteins of the respective other organism to mediate disassembly. Notwithstanding the fact that disassembly occurs even with exchanged machineries, the kinetics of disassembly may be slightly impeded by two non-cognate proteins at any of the interaction interfaces and are almost completely abolished when both interactions are mediated by non-cognate proteins.
Chapter 5

Discussion

The aim of this study was to investigate the molecular basis of SNARE disassembly including SNAP homologues, putative regulatory factors and the degree of functional conservation.

Since its discovery almost 30 years ago, surprisingly little progress has been made in deciphering the molecular details of NSF’s action in the cell. In part, this may be due to the fact that its function was originally misinterpreted to be the completion of fusion as such [68, 69, 70]. Only years later did it become evident that it is the SNARE assembly which leads to membrane merger, whereas the role of NSF is the recycling of dead-end SNARE-complexes after fusion instead. Nevertheless, its catalytic activities provide the fuel for intracellular membrane fusion by ATP-hydrolysis. NSF hence functions as the motor of membrane fusion, even though its activity is temporally uncoupled from membrane merger. The disassembly reaction thus bears a high degree of fascination, again leading to the question of why it has not been studied more extensively.

One reason probably is that the most efficient approach to get direct molecular insights into time-dependent protein interactions, such as enzymatic reactions, is to look at them using in vitro systems. A prerequisite for such studies is the availability of recombinant proteins. NSF however, being an enzyme, needs to retain its activity during isolation and handling to allow for its use in quantitative functional studies. Due to its fragility, its activity is easily lost under various buffering conditions. This difficulty adds onto the usual demands of protein purification, making it an especially challenging protein to be produced recombinantly.

A second difficulty when working with fragile enzymes, is to find easy ways of judging whether the enzyme is reproducibly active at the time of experiment or not. A read-out of disassembly which is simple enough to allow for direct conclusions about NSF-activity opens up this possibility. We therefore employed the strategy of developing time-resolved assays for NSF-mediated disassembly and used these for quality control of NSF already during various purification stages in order to help optimization of NSF purification. Since fluorescence spectroscopy of SNARE proteins is well established in our lab, it was the most obvious option to also employ this technique for the investigation of their disassembly. To this end, fluorescence-
Based experimental read-outs were established, which can directly monitor SNARE-complex interactions under disassembly conditions in a time-resolved manner.

During the time of my thesis, a second fluorescence-based in vitro assay using SNAREs fused to GFP-analoga as substrates was published [62, 67]. The authors successfully used the method to contradict previously described SNARE disassembly defects, which had been claimed to be caused by mutations in the SNARE-complex 0-layer [64, 65]. In the hands of Lauer et al., core SNARE complexes mutated in the 0-layer were disassembled equally well as wildtype SNARE core complexes in a time scale of seconds. Surprisingly, even though the amounts of NSF used were reasonably low for enzymatic reactions, the EC50 of αSNAP with respect to SNARE complex binding was 5µM, which is unexpectedly high considering that the complex concentration was only 100nM. Such low αSNAP-affinities have also been observed by other groups pursuing in vitro affinity-matrix based binding studies of αSNAP to SNARE complexes [71, 37]. Besides the fact that such high αSNAP concentrations are probably not physiological, their need is also difficult to reconcile with earlier experiments [72, 73], in which optimal functional reconstitution of αSNAP dependent processes was achieved using much lower amounts of αSNAP (≈0.6µM). Since, as mentioned above, some of the regulatory mechanisms might aim at the adaptor-level (namely αSNAP), these might stay hidden in a system requiring unnaturally high amounts of αSNAP for function.

Yet, in order to be able to also assess regulatory mechanisms and factors which do not act in an 'all-or-nothing' but rather a modulatory fashion, the conditions during the experiment should be as close to nature as possible. For this it is e.g. desirable, that none of the proteins involved is required in unreasonably high amounts for optimal function. As a prerequisite to addressing regulatory and mechanistic questions on the level of all proteins involved, including the SNAP-adaptor protein, this in vitro system should thus preferably lead to satisfactory reaction rates with lower amounts of αSNAP than those reported by Lauer et al. [62].

I hence sought to also understand, why the αSNAP affinity seems to be so low in some assays (and thus conditions), in order to get as close to physiological conditions in my in vitro studies as possible. The successful optimization of the assays finally made it possible to study the mechanisms of NSF-catalyzed SNARE-dissociation on a molecular basis.

5.1 Part 1 – Characterization of the SNARE Reaction Using Fluorescence Spectroscopy

The two main approaches used to characterize disassembly in this study, are FRET spectroscopy and fluorescence anisotropy. These fluorescence based read-outs depend on different physical properties. FRET spectroscopy exploits the fact that the amount of FRET (fluorescence resonance energy transfer) is proportional to the inverse power of six of the distance between two fluorophores. If two FRET-compatible fluorophores are attached to different proteins, the interaction of these
proteins can be seen in the fluorescence spectra of the fluorophores (as detailed in section 4.1.3).

The fluorescence anisotropy varies with the rotational motion of the fluorescence scattering of fluorophores in solution as detailed in section 4.1.3. This property may often be correlated to the size of the protein carrying the fluorophore, and conclusively serve as an indicator of protein binding.

To introduce the fluorescence into the system, we used small maleimide-conjugated fluorescent dyes, which specifically react with cysteine residues to form covalent bondages. An advantage of using these small fluorophores over the use of GFP-analoga is their small size and the possibility to specifically couple them to any cysteine in the molecule under study (provided that the mutation of the remaining cysteines does not impair the molecules' functionality). The spectral properties are therefore better defined and thus enable more defined distance measurements.

**Disassembly using FRET spectroscopy.** In short, for FRET spectroscopy the three neuronal SNAREs, two of them labelled, were pre-incubated to allow for FRET-complex formation. Subsequent addition of ΑSNAP and NSF under disassembly conditions then initiated the reaction (see figure 4.2). Several FRET pairs known to be suited for visualization of assembly were tested for their ability to also monitor disassembly. Of course, one would expect the same residues to get close during assembly which are taken apart again during disassembly and conclusively the FRET changes of these reaction should be the reverse of each other. Nevertheless, the labelling or the cysteine-mutations of certain residues might inhibit the disassembly reaction, which had to be ruled out first.

Doing this, two FRET pairs (Sb_{cys28}/SNAP25_{cys130} and Sb_{cys61}/H3_{cys225}) were soon found to be suited to monitor disassembly of ternary complexes (figure 4.2 and figure 4.13). It should be pointed out at this stage, that the Sb_{cys61OG}/H3_{cys225TR}/SNAP25-complex turned out to have strong disassembly defects in combination with a range of additional mutations (figure 4.14), which like the Sb_{cys61}/H3_{cys225} FRET pair did not noticeably inhibit disassembly on their own. Besides the putatively interesting mechanistic implications of these findings, which I will touch on again in the next section, they should also be seen as a reminder of how important it is to have several read-outs (e.g. different label positions) leading to identical results before any findings should be interpreted.

**Disassembly and ΑSNAP binding using fluorescence anisotropy.** Similarly, different labelling positions were tested for changes of fluorescence anisotropy during disassembly (figure 4.3). Again several labelling positions were observed to give rise to significant signal changes after addition of NSF, ΑSNAP and Mg^{2+}/ATP.

**Basic characteristics of the disassembly reaction.** The fluorescence changes indicating SNARE disassembly were shown to be NSF-, ΑSNAP-, Mg^{2+}- and ATP-dependent in both assays, making it very likely that they in fact represent ongoing
disassembly. Neither the mutant $\alpha$SNAP$^{L294A}$ nor a hydrolysis-mutant of NSF, NSF$^{E329Q}$ (data not shown), were able to promote disassembly. Furthermore, an instantaneous block of the reaction could be provoked by addition of AlF3, a compound known to freeze AAA ATPases in a transition state.

In line with reports mentioning a salt-sensitivity of the $\alpha$SNAP/SNARE-complex interaction [28, 71, 67], kinetics slowed down extremely in presence of as little as 150mM NaCl (data not shown). A salt combination of 120mM KGlu and 20mM KAc, which better mimics physiological conditions, however did not negatively influence disassembly, which is why I used this saltcomposition for all experiments shown in the thesis.

Detection of $\alpha$SNAP binding via fluorescence anisotropy. In addition to disassembly, some of the labelling positions can also be used to monitor $\alpha$SNAP binding to the SNARE complex via fluorescence anisotropy. This was demonstrated for the SNARE complex labelled at Syb$^{\text{Cy28}}$, where $\alpha$SNAP binding leads to a sharp increase of anisotropy (section 4.1.5).

The $\alpha$SNAP mutant $\alpha$SNAP$^{L294A}$, which has been reported to bind, but not to disassemble complexes, led to the same anisotropy increase upon addition to the fluorescent complex as wildtype $\alpha$SNAP. But in contrast to wildtype $\alpha$SNAP, this mutant did not induce the anisotropy decrease representing disassembly in this method, even when NSF, Mg$^{2+}$ and ATP were present. These are further evidences for the notion that the rise of anisotropy upon $\alpha$SNAP-addition actually represents binding, whereas the decrease upon addition in presence of magnesium, ATP and NSF indicates the disassembly.

Quality control of NSF. As mentioned above, the fluorescence read-outs were employed to optimize NSF purification procedures until reproducibly active NSF could be purified. NSF activity was monitored at various stages and different purification conditions, in order to get a better understanding of how NSF may be expressed and purified in a reproducibly active manner. The optimal procedure to produce recombinant NSF was found to be Ni/NTA- followed by gel-filtration chromatography as explained in section 4.1.1 in more detail. NSF purified according to these methods was found to disassemble $\sim$15 (+/-2) SNARE-complexes per min per $\mu$gNSF under conditions in which $\alpha$SNAP is saturating as calculated by the half times of the reaction. This is similar to the published activity found in a fluorescence based in vitro assay using SNAREs fused to GFP-analoga as substrates. [62, 67]

ATP, NSF and $\alpha$SNAP dependence. The ATP requirements for disassembly were shown to be in the range of 75$\mu$M (figure 4.5). This finding is in line with Matveeva et al. [74], who found that one NSF monomer contains two distinct ATP binding sites with a 100-fold difference in affinity. According to that study, the low affinity binding site exhibits a $K_d$ of 40nM and is not sufficient to stimulate ATPase activity, whereas the high affinity site with a $K_d$ of 15-20$\mu$M stimulates ATP hydrolysis.
NSF activity was constant when employed in a range of 1nM to 16nM (figure 4.4), which is in agreement with data published with the fluorescence based assay of Lauer et al. [62], where 1nM to 10nM were reportet to result in disassembly in a linear range.

As mentioned above, the αSNAP requirements for saturation of SNARE complex binding vary between different sources of literature, many of which report surprisingly low αSNAP affinities. I thus also determined the amounts of αSNAP needed for disassembly in the FRET and the anisotropy read-out and quantified αSNAP binding to SNARE complexes using anisotropy.

It soon became evident that in these assays the αSNAP efficiency is very low as well: More than a seven fold excess of αSNAP did not saturate the kinetics of disassembly (figure 4.6). This may even be an underestimate, because at higher αSNAP concentrations the disassembly speeds reach the time resolution limit of the assay. This is astonishing if one considers that only three αSNAPs are supposed to bind one complex according to previous findings. Later, using lower amounts of NSF and thereby slowing down reaction kinetics, I could demonstrate that even 1.1µM of αSNAP are still limiting when disassembling ~100nM SNARE complex. The titration experiment shown in section 4.16 indicated that more than 17 αSNAP molecules per SNARE complex are needed to saturate binding, when 70nM of SNARE complex are used.

Having observed this low αSNAP efficiency in the fluorescence assays, the question came up, whether there was a problem with the recombinant αSNAP used for the experiments.

On the one hand, as indicated above, the high αSNAP requirements in my assays are in line with Lauer et al.[62], who determined the EC50 for αSNAP binding to SNARE-complexes to be 5µM in their hands, at a complex concentration of only 100nM. Such apparently low αSNAP-affinities have also been observed by other groups persueing in vitro affinity-matrix based binding studies of αSNAP to SNARE complexes.[71, 37].

On the other hand, besides the fact that the high αSNAP concentrations needed to saturate SNARE complex binding are probably not physiological, their need is also difficult to reconcile with earlier experiments [72, 73], in which optimal functional reconstitution of αSNAP-dependent processes was achieved using much lower amounts of αSNAP (less than ~0.6µM). Furthermore, according to a semi-quantitative Western blot performed in our lab, the αSNAP concentration in the cell was estimated to be much lower (T. Lang, personal communication).

It was important to address this question, because excessive needs of one component for optimal function might also point to an imperfect interaction of this component under the conditions used. If this were to be the case for αSNAP, it would not be possible to study the reaction in a meaningful manner, because the reduced binding or the excess amounts of αSNAPs might shift equilibria of the remaining interaction parameters. This might lead to a scenario in which other bottle-necks of the reaction as well as regulatory mechanisms might be overlooked, especially if these act on the adaptor level.
5.1.1 The Membrane is a SNAP Receptor

Possible reasons for low $\alpha$SNAP efficiency In principle, the low $\alpha$SNAP efficiency could be inherent to the recombinant $\alpha$SNAP, possibly due to degradation of only a few residues which would not be visible on a gel or posttranslational modifications present in vivo but absent when expressed in E.coli. Degradation might for instance produce $\alpha$SNAP which is no longer able to bind SNARE complexes. Consequently, only a fraction of the presumably high amount of $\alpha$SNAP used in the experiments would be active. Alternatively, the low efficiency might be caused by non-optimal conditions in the fluorescence assays.

The recombinant $\alpha$SNAP binds SNARE complexes in a 3:1 stoichiometry.

As shown in section 4.3.1, I also investigated $\alpha$SNAP binding to the SNARE complex using isothermal titration calorimetry. The resulting enthalpies however were very low ($\sim 4$ kcal/mole of injectant), making the accurate determination of released energies and consequently of the affinity impossible. The stoichiometry of the SNARE complex could nevertheless be determined to be three $\alpha$SNAPs per complex in ITC experiments. Notably, this is exactly what is expected and hence allows us to conclude that the majority of the recombinant $\alpha$SNAP is able to bind SNARE complex, excluding the possibility that degradation is the reason for the low potency. The main difference between the ITC conditions and those during the fluorescence assays being the much higher protein concentrations during ITC measurements, two possible explanations remained: Either does the recombinant $\alpha$SNAP have a reduced affinity as opposed to endogenous, or a factor important for binding is missing in the fluorescence assay. Notwithstanding the low enthalpy values of the reaction, I still tried to fit the reaction in order to roughly estimate the $\alpha$SNAP affinity. The thermodynamic data could not be properly fitted using a binding model, which assumes that $\alpha$SNAP has the same affinity to all binding sites on the SNARE complex. Using a three-site sequential fit however, the curve could nicely be fitted (see figure 4.17). The affinities resulting from this fit were 10nM for the first, 50nM for the second and 380nM for the third binding site. Nevertheless, keeping in mind the low enthalpies and also considering that only a low number of data points per binding site were recorded, the affinities of $\alpha$SNAP to the SNARE complex determined might not be very reliable.

The recombinant $\alpha$SNAP has a high SNARE-affinity on membrane sheets.

To investigate the efficacy of the recombinant $\alpha$SNAP under the more physiological conditions of a different system, we performed disassembly measurements using the membrane sheet assay introduced in section 4.3.2. Surprisingly, in the membrane sheet assay $\sim 100nM$ of $\alpha$SNAP were sufficient to mediate disassembly equally well as $2\mu M$, indicating that the $\alpha$SNAP concentration of $100nM$ saturates the assay. Of course the amount of SNARE target in these assays is not known, not permitting a conclusion about the stoichiometry of $\alpha$SNAP and SNARE complexes in this assay. Nevertheless, together with the ITC measurements...
which indicated a molar ratio of 3:1, these findings strongly suggest that a factor missing in the FRET and anisotropy experiments but present on membrane sheets is responsible for the higher αSNAP-affinity.

The αSNAP efficiency is enhanced when disassembling liposomal complexes. The most obvious difference between the in vitro fluorescence spectroscopy and the membrane-sheet assay was the absence or presence of lipids, respectively. Since the disassembly reaction in the cell is most likely supposed to occur only after fusion has completed and all three SNAREs are located on one membrane, it is of course tempting to speculate that the membrane is of some importance during the disassembly reaction. One possible way of communication between the disassembly machinery and the membrane would be an interaction of αSNAP with the membrane in addition to its interaction with the SNARE-complex, which consequently might increase αSNAP/SNARE affinity. Such a scenario is especially feasible, since αSNAP is an amphiphilic protein which has been reported to bind to plastic-surfaces [51]. Furthermore, αSNAP was shown to bind lipids in a SNARE-independent manner [50].

To test the hypothesis that αSNAP might be more efficient in the presence of membranes, FRET-spectroscopy and anisotropy measurements were carried out using SNARE complex as a target which was incorporated into liposomes prior to the experiment. Indeed, the αSNAP efficacy was found to be increased by a factor of ~10 to 20 compared to its efficiency in solution (figure 4.22). As little as 45nM of αSNAP mediated disassembly of ~50nM liposomal SNARE complex and 120nM were sufficient to saturate the assay (see figure 4.21). The reasons for the improved αSNAP efficacy in the presence of liposomes were not known at this stage.

NSF activity is not enhanced on liposomes. To investigate, whether the membrane boost is only due to a higher αSNAP affinity or whether the incorporation of complexes into liposomes also influences NSF activity, NSF-action on liposomal and soluble SNARE complexes was compared at αSNAP concentrations promoting optimal disassembly, respectively. Doing so, NSF proved to be as effective on liposomes as in solution, indicating that its activity or SNARE/SNAP affinity are not influenced by membrane incorporation of the SNARE targets. Furthermore, the same amount of an antibody against the N-terminal domain of NSF was needed to block NSF activity on liposomes as in solution, further strengthening the conclusion.

Neither of the SNARE transmembrane domains provides the additional αSNAP binding site. Next, the reasons for the membrane caused potentiation of αSNAP efficiency were elucidated. Alternatively to the lipids now present in the reaction, one of the transmembrane domains used to anchor the SNARE complexes to the liposomes might also be responsible for the improved αSNAP potency. To exclude this possibility, reactions were performed using either only the Synaptobrevin or only the Syntaxin transmembrane domain (figures 4.21 and 4.26). Both showed
identical αSNAP affinities, demonstrating that neither of the transmembrane domains is responsible for the improved disassembly.

The mere presence of lipids does not suffice. γSNAP and Sec17 have been reported to have highly flexible C-termini, which is why for both of these proteins the very C-terminal residues are not included in the solved crystal structures [30, 29]. One could now argue that this apparent flexibility might hamper function and that the lipids present during the liposome experiments might simply stabilize the αSNAP C-terminus and thereby improve its affinity. The observed potentiation effect would thus not be caused by the membrane anchorage but only result from an optimized stabilization of the recombinant αSNAP. To exclude this possibility, I disassembled SNARE complexes in a mixed micelle solution, where a fraction of SNAREs would be expected to reside on liposomes, whereas the remainder would be in solution. Here, part of the disassembly reaction required only low amounts of αSNAP to proceed, while the remainder only occurred at a high αSNAP concentration (see figure 4.29). The fraction of disassembly at low and high αSNAP concentrations correlated with the fractions expected to be liposomal or solubilized, respectively (figure 4.30).

The membrane boost is independent of the lipid composition. Various protein/lipid interactions are known to depend on a specific lipid component. I therefore tested, whether the improved αSNAP function on liposomes is dependent on the lipid composition. To do so, I reduced the heterogeneity of the liposomes, which had been constituted of PC, PE, PI, PS and cholesterol to closely resemble neuronal plasma membrane content in the earlier experiments. Interestingly, disassembly of SNARE complexes on liposomes consisting of only PC and PE showed a similar αSNAP dependence as those on the more heterogeneous liposomes used previously (figure 4.28). Even liposomes consisting of PC only were disassembled at αSNAP concentrations of less than 100nM (data not shown).

The N-terminus of αSNAP mediates lipid binding. These data made it tempting to speculate that a direct interaction between αSNAP and the membrane lipids is responsible for the increased efficacy on liposomes. If this were true, it should be possible to map the lipid-binding property to a certain region of αSNAP. Deletion of this region should then abolish the membrane interaction and consequently the observed boost of αSNAP effectivity on membranes. Since no crystal structure of αSNAP has been solved so far, we used the Sec17 crystal structure as a reference to predict the putative αSNAP region responsible for lipid binding (figure 5.1A). The C-terminus of αSNAP is believed to mediate the interaction with NSF, which is believed to bind to the SNAP/SNARE acceptor-complex from the cytosolic site. According to the current state of evidence, αSNAP and the SNARE complex are thus likely to align in an anti-parallel fashion, which places the N-terminus of αSNAP close to the membrane. Based on the Sec17 structure and
αSNAP/SNARE-complex interaction studies using various point-mutated αSNAPs, Marz et al. have constructed a model of αSNAP bound to the SNARE complex [67]. An adapted version of this model is shown in figure 5.1 B. Here one can see an arm-like structure at the N-terminus of αSNAP pointing away from the complex, the most far-out region of which (residues 27-32 in αSNAP) includes mostly hydrophobic amino acids. This region corresponds to the loop between helices α1 and α2, designated α1’. If this region were the interaction site of αSNAP with the membrane, its deletion should abolish membrane binding and hence the higher αSNAP efficiency on membranes. I therefore cloned an αSNAP-mutant in which I deleted the first 32 residues (designated αdelSNAP), in which the critical region would be abrogated.
Figure 5.1: The hypothetical lipid binding site of αSNAP. (A) Structure of Sec17. Helix α1 and loop α1’ are missing in the αSNAP mutant αdelSNAP as indicated by the pair of scissors, (adapted from Rice et al. [30]) (B) Model of αSNAP/SNARE-interaction adapted from [67]. The SNARE complex is depicted in grey and the αSNAPs modelled on top are shown in red, orange and yellow. The α1’ loop of αSNAP points away from the SNARE complex towards the membrane. This putative αSNAP/lipid interaction site is highlighted in green. (C) Sequence alignment of several αSNAP homologues. Dark gray shading indicates strong conservation, gray shading conservation. The hydrophobic residues F27 and F32 are highlighted in green and conserved throughout all species.
First of all it was shown that αdelSNAP is able to disassemble, even though at a lower efficiency than αSNAP. This reduced overall efficiency is in line with findings by Hayashi et al [71], who found a 75% reduction of αSNAP binding to SNARE complexes after deletion of the first 28 amino acids using a GST-binding assay.

As opposed to αSNAP, the kinetics of αdelSNAP-mediated disassembly in solution and on liposomes were identical, as we had predicted which is shown in section 4.4.4. As a matter of fact, deleting the first 32 amino acids hence abolished the potentiation of αSNAP efficiency at the membrane.

Secondly, I directly compared αSNAP and αdelSNAP efficiencies on liposomes. Theoretically, it should be predictable at this stage, how much more efficient the wildtype αSNAP should be on liposomes. Now knowing that αdelSNAP has an approximately six fold lower efficiency than αSNAP in solution, and from my additional findings (figure 4.22) that the efficiency of wildtype αSNAP increases by a factor of 10-20 in the presence of membranes, the efficiency difference on liposomes can be estimated. Accordingly, an efficacy factor in the range of 60-120 (roughly six times ten to twenty), would be expected. As a matter of fact, experiments carried out to confirm this prediction (and hence also confirm the hypothesis that αdelSNAP has indeed lost the ability to be boosted by membranes) indicated a 80-fold higher efficiency for αSNAP.

Taken together, these findings strongly suggest that the N-terminal arm-like structure observable in the crystal structure of αSNAP is essential for a protein/lipid-interaction which mediates the potentiation of complex disassembly at low αSNAP concentrations. An additional αSNAP binding site is hence provided by the membrane lipids, presumably leading to a drastic enhancement of the overall αSNAP affinity to its acceptor site. This might be a mechanism of cooperative binding of the SNARE complex and the membrane, which stabilizes αSNAP at the site of catalysis.

**Strategy to further confine the αSNAP residues constituting the lipid binding site.** According to the above-mentioned SNAP/SNARE binding model proposed by Lauer et al. [62], there is a loop close to the N-terminus of αSNAP, which points towards the direction of the membrane. This loop α1’ contains hydrophobic residues at the positions homologous to F27, F28, L31 and F32 in αSNAP throughout all sequences shown in the alignment in figure 5.1 C, two of which are strictly conserved (F27 and F32). Lauer et al. state [62] that the C-terminal globular bundle of αSNAP was left out of their model in order to allow for the tight binding of αSNAP along the whole SNARE-complex surface. They argue that the globular bundle, if fitted into the model according to the bend observed in the Sec17 structure, would have crashed with the SNARE complex. They justify the removal with the proposal that this C-terminal globular bundle might flexibly bend away upon interaction with the complex. There is however no evidence for this notion apart from the fact that the two structures would collide if the bundle would not have been omitted. Considering the low enthalpy-changes during my ITC measurements
of αSNAP-binding to the SNARE complex, the interaction between SNAP and the SNARE complex might on the other hand also be weak, so that the two proteins might not be bound as tightly as indicated in the model. Now imagining that the C-terminal bundle of αSNAP were present and would not bend away, one can picture the upper αSNAP portion shown in the model to tilt outwards. This would bring the loop α1’ even closer to the membrane (which I schematically introduced into the picture) and hence make a interaction of the SNAP and the membrane at this site highly probable (as depicted by arrows). In order to test, whether the lipid binding site of αSNAP is as actually confined to this loop constituted of 6 residues, an αSNAP mutant could be constructed accordingly and tested as previously described for αdelSNAP.

**Putative mechanism of the 'membrane boost'.** Having shown that an additional binding site between αSNAP and the membrane strongly increases the affinity of αSNAP to the SNARE acceptor, the question still remains, which mechanistic implications the increased αSNAP affinity has for SNARE disassembly. Spontaneously, one can think of two possibilities. For one, the lipids might simply function as a kind of SNAP collector, binding to αSNAP and thereby increasing its local concentration on the membrane and thus near the complex. The increased local concentration would then support SNAP binding. Alternatively, the SNAP/lipid interaction might actually support the strength of αSNAP binding and maybe even increase the stability of SNAP/SNARE-binding during disassembly. A combination of both scenarios is also conceivable.

**The SNAP isoforms α- and β-SNAP are exchangeable – Even the membrane boost is conserved.** Since α- and β-SNAP have 83% sequence identity [34], one possibility is that they are simply functionally similar iso-forms. This view is supported by the finding of a single SNAP in Drosophila as well as squid [75] with sequences, which are equally related to those of both mammalian α- and β-SNAPs.

There have been contradictory findings with respect to βSNAP’s interchangeability with αSNAP [33, 32, 31]. None of these groups, however, have directly compared βSNAP and αSNAP disassembly in a time-resolved experiment. During this thesis, I compared α- and βSNAP-mediated SNARE disassembly in solution as well as on liposomes. Due to ongoing degradation problems during purification, the efficiencies of both proteins can not be compared in an absolute manner, but βSNAP could be shown to efficiently disassemble neuronal SNARE complexes. In addition I found that the relative potentiation of efficiency is identical from solution to liposomes for both proteins. It can thus be concluded that the membrane boost found for αSNAP is conserved for its isoform βSNAP.
5.1.2 Summary – Advantages of the Fluorescence-Based Experiments over Conventional Disassembly Assays

The fluorescence assays presented here have major advantages over other published in vitro assays of SNARE disassembly.

Quantitative monitoring of disassembly with high time resolution. First, the efficiency of disassembly can directly be related to the changes during SNARE-assembly, providing a direct possibility to quantify the amount of SNARE complex disassembly. Second, signal changes can be recorded online, making the time needed for efficient mixing (approx. 5s, depending on the viscosity of the buffer) the limiting parameter. The reaction can thus be monitored on-line, which additionally allows for the accurate quantification of reaction rates. At the same time, the 'real-time' recordings also make the observation of reversible changes possible. If, for example, there were a point in a reaction under study, at which one of the disassembling factors got limiting and reassembly took over, this would be visible in the spectrum. Adding excess amounts of an unlabelled SNARE of the same type as one of the labelled SNAREs in the complex to the solution can be used to make the reassembly invisible. If Synaptobrevin, which does not bind αSNAP [71] and should thus not interfere with the disassembly reaction, is used for this purpose, different kinetics in the presence or absence of unlabelled Synaptobrevin point towards ongoing reassembly. As opposed to read-outs with a limited time resolution, this approach can therefore be employed to differentiate between defects in disassembly and an enhanced rate of assembly. Additionally, the use of FRET pairs at different positions in the complex also allows to see, whether the SNARE helices or certain parts of these significantly change their position already upon αSNAP or NSF binding. Major structural rearrangements upon docking of the disassembly machinery should hence be observable. However, I did neither observe drastic changes in FRET intensities upon addition of αSNAP and NSF nor when I employed AlF₃ which supposedly freezes NSF in a transition state, indicating that (at least with respect to the distance between residue 130 in SNAP25 and residue 28 in Synaptobrevin) no major structural rearrangements take place before the actual dissociation in presence of Mg²⁺ occurs.

The incorporation of SNAREs into liposomes dramatically increases αSNAP efficiency. The use of liposomes for anchorage of the SNARE complexes is a major improvement of the assay, also with respect to the fluorescence based assay published by Lauer et al. [62]. The additional αSNAP binding site provided by the membrane strongly increases αSNAP affinity to the SNARE-complex compared to the affinity found in membrane-free experimental set-ups. The fluorescence systems established during this thesis hence are the first fully reconstituted in vitro assays to directly monitor SNARE-complex disassembly, in which the complete high-affinity αSNAP binding site is present. We can thus use stoichiometric amounts of αSNAP for our studies, which is essential to solve questions concerning the αSNAPs/SNARE complex-interactions during disassembly. This is especially interesting, since only
little is known about the precise mechanism of SNAP/SNARE-interactions during disassembly. Even though three αSNAPs have been determined to bind to one SNARE-complex in purified 20S complexes as determined by quantitative amino acid analysis [61], it is not known whether three αSNAP actually have to be bound for functional disassembly. It would for instance also be possible that one or two αSNAPs suffice to promote disassembly, maybe at a lower speed. All results collected in this study are in line with such a scenario, as will be explained in the next section.

5.2 Part 2 – Modulating the Reaction

5.2.1 Robustness of the Reaction – Bottlenecks and Putative Regulatory Targets

The second part will focus on different approaches to increase the current understanding of the molecular interactions of NSF, the SNAPs and the SNARE complex during the disassembly reaction. To this end, I applied several strategies to target different components of the reaction, most of which aim for a better understanding of the SNAP/SNARE interface. As mentioned above, one major insight into the molecular details of this interface was the additional αSNAP binding site provided by the membrane lipids, which leads to a dramatically increased SNAP/SNARE affinity.

However, it as yet remains unclear, which parts of the SNARE complex are important for disassembly. Likewise, with respect to the assumed three sites of αSNAP/SNARE- and even six sites of NSF/αSNAP-interaction, it is not known to which degree the machinery is robust in the sense that the specific interruption of single interaction-sites only decreases reaction rates or leads to a complete abolishment of disassembly.

Obviously, one could argue that it should be easy to address this question, especially now that we are able to apply stoichiometric amounts of αSNAP. For instance, one might simply use less than three αSNAPs per complex and quantify the reduction of disassembly. Unfortunately, this is not as easy as it sounds, because so little is known about the characteristics of the SNAP binding site. Neither is it known, whether the SNAP affinity to all sites is equally high or whether they display two or three unequally affine binding sites. Nor do we know, whether there is cooperativity between the different sites, which might lead to the occupation of two or all three sites on one complex before the first site of the next complex will be occupied. When using limiting amounts of αSNAP we would thus never be sure about whether only part of the complexes is occupied or whether every complex is occupied incompletely. Kinetically, this can not be distinguished either, because due to the enzymatic properties of the reaction, the complexes would simply be disassembled one after another in the first scenario, whereas the second scenario might lead to a reduction of the speeds of the disassembly as such - both would hence result in
slower total disassembly kinetics.

The ITC measurements performed in section 4.3.1 pointed to a sequential binding (figure 4.17), but as the enthalpies of αSNAP/SNARE complex binding were very low, it is difficult to interpret the measurements and the affinities for the different sites could not be reliably determined and cooperativity thus not be investigated. The anisotropy approaches can not be used to differentiate between different sites of binding either. Even though the anisotropy read-out of Sb\textsubscript{cys}\textsuperscript{28}-labelled complexes serves an indicator of αSNAP binding, we do not know whether the signal increase seen results from the binding of only one αSNAP in close vicinity to the label position or whether all three SNAPs contribute to the signal. The collective term of 'the αSNAP affinity' used so far is hence an average property of the amount of αSNAPs needed for optimal disassembly, which does not differentiate between the distinct αSNAP/SNARE interaction sites.

Due to the lack of direct possibilities to address the properties of the distinct αSNAP binding sites, indirect approaches may help. If, for example, certain deletions in the SNARE complex led to inhibited disassembly, this might result from an ablation of certain αSNAP binding sites. It should nevertheless be kept in mind that SNARE deletions leading to inhibited disassembly do not necessarily always result from an interrupted interaction site but may as well directly (e.g. sterically) inhibit the mechanics of the disassembly reaction. Anyways, a better knowledge about such coherences might provide clues about the relation between αSNAP binding sites and SNARE-disassembly.

Hence, NSF and the SNARE-complex were selectively altered in order to see, whether the alterations would affect disassembly. It should be pointed out that the experiments discussed in the following section were performed before the discovery of the αSNAP/lipid interaction and were thus carried out in solution. Since in these experiments either NSF or the SNARE complex, but not αSNAP were modified, αSNAP was always present in excess during the reactions. Alterations completely abrogating the interaction with αSNAP should lead to inhibited disassembly even in the presence of excess αSNAP. Slight alterations of SNAP/SNARE- or NSF/SNAP-binding on the other hand might be overlooked taking this approach in solution.

**Inhibition on the level of NSF - Disturbing the NSF/SNAP interaction?**

First, the interaction interface between NSF and the SNAPs was addressed. Experiments performed with the monoclonal anti-N antibody (type IgG2) generated in the context of this study led to a complete block of disassembly. It is likely that the antibody interferes with NSF binding to αSNAP, which has been reported to depend on the NSF N-terminal domain [22]. Since all αSNAP/NSF connection sites presumably are structurally identical, a structural change in one of the proteins is likely to affect all SNAP/NSF interactions equally, and consequently the complete abolishment of SNARE disassembly is not surprising. It should nevertheless be noted, that the antibody might also inhibit the reaction by different means, e.g. steric hindrance of the NSF molecule during disassembly. The antibody was shown to specifically recognize only NSF of all the proteins present in cytosol, and might
hence also serve useful as a tool to specifically block disassembly in \textit{in vivo} studies in the future.

\textbf{Alterations on the target-level.} The minimal SNARE regions required for SNARE disassembly \cite{6} as well as various mutants which may or may not inhibit SNARE disassembly have been described \cite{71, 64, 62, 65}. SNARE assembly has been shown to exhibit an N-terminal nucleation making the N-terminal residues obligatory for efficient assembly. It is possible that for disassembly analogous obligatory residues exist. For example, the conservation of the '0'-layer residues has originally been suggested to be essential for disassembly. This led to the hypothesis, that the '0'-layer might be the area of attack of the disassembly machinery \cite{64, 65}. In the scope of this thesis, I re-investigated some of these findings as well as some additional SNARE complex alterations, exploiting the increased time resolution of the fluorescence assays as compared to the approaches used in the original publications.

As pointed out earlier, due to the asymmetric structure of the SNARE complex, the interaction between the SNAPs and the SNARE-complex target is mediated by at least two, and most likely three, structurally different sites. Losses of interaction at any of these three sites as well as their consequences can thus be addressed independently.

\textbf{The disassembly reaction tolerates ‘single-locus’ target alterations.} First, I ensured that the complexes containing only the H3 domain of Syntaxin are equally prone to disassembly as full-length Syntaxin, which supports earlier findings from our lab gathered with the gel-based disassembly assay \cite{6}. Knowing this, I used only the H3 domain for all remaining experiments performed in solution. To elucidate whether certain regions of the SNARE complex are essential for disassembly, I next investigated various SNARE complex mutants.

Amongst these were the Synaptobrevin '0'layer mutant (Sb\textsubscript{R56P}) as well as the Syntaxin '0'layer mutant which have both led to controversial results between different approaches in the literature. In my hands, both mutants did not inhibit disassembly in the fluorescence assays as demonstrated in figure 4.9, which is in line with the findings of Lauer \textit{et al.} \cite{62} that were published during this work. Both mutants have been reported to inhibit disassembly before \cite{64, 65}, but these findings might also stem from an enhanced re-assembly capacity of the mutant SNAREs, which would not be differentiated in their gel-based disassembly assays. Alternatively, a slightly reduced \(\alpha\text{SNAP}\) affinity to the mutated complexes might be the reason for the disassembly defects observed by Scales \textit{et al.} and Martinz-Arca \textit{et al.}, which might not be apparent in my fluorescence measurements performed in the presence of excess \(\alpha\text{SNAP}\). Indeed, a slightly reduced affinity of \(\alpha\text{SNAP}\) to the Syntaxin\textsuperscript{R226Q} containing complex was reported by Lauer \textit{et al.} \cite{62}.

Furthermore, C-terminal deletions of SNAP25 as well as point mutations exchanging the two amino acids which SNAP25 contributes to the '5'-layer of the neuronal SNARE-complex (SNAP25\textsuperscript{M71A,F192A}) did not inhibit SNARE-complex disassembly as shown in figure 4.10. Likewise, a C-terminal Synaptobrevin deletion (Sb1-
70) did not impair disassembly at all, whereas a C-terminal Sx deletion (Sx183-240) led to a slight inhibition (figure 4.11). The Sx mutant however did not reduce total amounts of disassembly but only slightly impaired the reaction rate at the later stages. Even when both, Sb and H3, were concertedly deleted, the inhibitory effect did not increase.

In summary, none of these mutants lead to efficient inhibition of SNARE disassembly. These findings lead to the conclusion, that the interaction interface between the disassembly machinery and the SNARE complex is not confined to a single site (as e.g. the '0'-layer or the C-terminus) on the SNARE complex but probably covers a larger portion of the SNARE surface. Partial interruption of the interface might then weaken the interactions but not abolish it completely.

Even though it is interesting that the disassembly reaction seems to be robust enough to tolerate these various target alterations, the experiments pursued this far did not lead to any mechanistic insights concerning the putative bottlenecks or protein/protein interactions critical for the performance of disassembly. When searching for efficient ways to disturb disassembly, as an alternative to selectively deleting target-regions one can add factors which might impede either target/adapter-recognition or hinder the reaction sterically. Good candidates for proteins whose attachment might influence function are antibodies. To this end, I tested three anti-SNARE complex antibodies for their effectiveness to block disassembly. Two of these antibodies (132.1 and 132.2, see figure 4.2.3) indeed partially inhibited disassembly. However, as also observed for the deletion mutants, both of these antibodies did not inhibit disassembly completely. According to the SNAP/SNARE interaction-model proposed by Lauer et al., the αSNAPs cover almost the whole surface of the complex when all three are bound. If this model is correct, it is highly unlikely that none of the many complex alterations or antibodies tested interfered with any of the SNAP/SNARE binding sites. This might indicate that disassembly may be able to proceed, even when less than three αSNAPs are bound. At the same time it is also possible, that the single mutations do not completely abrogate SNAP/SNARE binding at one site but only reduce the interface of binding and thus destabilize the interaction. The two αSNAPs bound to the remaining SNAP/SNARE interfaces might then be sufficient to compensate for certain destabilizations at one site by a mechanism of cooperativity. The only conclusion that can thus be drawn from these experiments is that the disassembly reaction is robust in the sense that deletions at single interaction interfaces can be tolerated.

Interference at two distant sites can result in severe inhibition. To further strengthen the notion that less than three SNAPs may be able to disassemble one SNARE complex, dual target alterations at non-proximal sites were investigated next, aiming at a possible disruption of more than one SNAP/SNARE binding site which might lead to more pronounced disassembly defects.

As a matter of fact, even though none of the mutations or antibodies discussed in the last paragraph led to severe impairment of disassembly, the reaction could ultimately be blocked in several cases, when two of these mutants or antibodies were
combined. For one, in the presence of both of the two partially inhibitory antibodies, 132.1 and 132.2, the reaction came to a complete halt. Similarly, the combination of the FRET-pair Sb<sub>cys</sub>61/H3<sub>225</sub> and any of the C-terminal or 5-layer SNAP mutants abrogated disassembly almost completely. Conclusively, one defect on the target level does not seem to be enough to abrogate disassembly, whereas the combination of two mutations can strongly interfere with disassembly, even for deletions which are completely silent with respect to disassembly when present independently.

**Conclusion:** The disassembly mechanism might tolerate partial impairment of αSNAP binding. Interestingly, none of the various single-site mutations of the SNARE complex described above severely inhibited SNARE disassembly, most did not even disturb disassembly at all. Even the combination of two C-terminal SNARE deletion mutants (Sb and H3) did not increase the slight inhibition witnessed for H3 alone. Only when two different loci on the SNARE complex were disturbed, severe inhibition of the reaction could be observed.

It can not be excluded that slight impairments on the level of αSNAP binding, which might lead to less disassembly, are overseen in these experiments, because they were performed using excess amounts of αSNAP. Alterations which strongly interfere with αSNAP binding should be observable, but we would not know if mutations led to a slightly changed αSNAP affinity. The single-site mutants found to function like wildtype here, can hence not be excluded to have slightly changed αSNAP binding-characteristics compared to the wildtype complex. To elucidate whether this is the case for several of the single-site mutants and if so, whether this has an impact on the kinetics of disassembly when more physiological amounts of αSNAP are used, one could carry out these experiments again, this time on liposomes which allow for the use of stoichiometric and substoichiometric amounts of αSNAP.

What can nevertheless be seen from these experiments, is that even in the presence of excess amounts of αSNAP, the complex can be disrupted severely enough to abolish disassembly. Strikingly, these severe inhibitions are brought upon only when several mutations are introduced together, the mutations can thus have synergistic effects. Interestingly, only when two different loci on the SNARE complex were disturbed, severe inhibition of the reaction could be observed, whereas for instance two nearby deletions at the C-terminus (shortened Syntaxin and Synaptobrevin) did not show synergistic effects. It is tempting to speculate that the defects result from ablations of αSNAP/SNARE complex binding. If one thinks along these lines, the observations suggest that the NSF machinery indeed might be robust enough to function even though one of the interaction sites between substrate and machinery is weakened or blocked. Considering that the antibodies have been mapped to different regions of the complex, it is possible that the two synergistic clones 131.1 and 131.2 interfere with different SNAP/SNARE interaction sites. Similarly, the other structural alterations of the neuronal SNARE complex which also showed synergistic effects, are located such that they might interfere with different SNAP binding sites. This might indicate that even if one site is weakened or blocked, other sites might be able to compensate for the defect and suffice to mediate disassembly.
Part 2 – Modulating the Reaction

Notwithstanding the fascination of this hypothesis, it might as well be, that the SNARE-complex alterations do not interfere with αSNAP binding but rather change the target structure in such a way that the conformational changes required for efficient disassembly can not occur sufficiently. This possibility is easy to picture when thinking of bound antibodies to the SNARE complex, which might lead to steric hindrance. For the defects caused in the Sb610G H322STRU complexes combined with the SNAP25-mutations this possibility is albeit less plausible, because the strong defects occurred when an additional deletion as opposed to an attachment was introduced.

To further elucidate the question, whether less than three αSNAPs per SNARE complex can mediate disassembly, I will now switch perspectives, away from the target- to the adapter-level. Whereas the previous experiments were aiming at interrupting SNARE/SNAP interactions via complex-alterations, I now searched for a way to directly hinder αSNAP-binding without the need of destroying the SNARE complex.

5.2.2 Inhibition on the αSNAP Level

To specifically inhibit the SNARE/SNAP interaction face via changes on the level of αSNAP, several possibilities exist. Obviously, in analogy to the experiments using altered SNAREs described in the last section, one could delete or mutate specific regions or residues of αSNAP and afterwards look at these mutants’ effects on disassembly. This strategy has for instance been employed by Marz et al [67], who mutated various acidic and basic αSNAP residues and could thereby show that the SNARE/SNAP interaction is mediated by mostly ionic interactions. The model constructed hereafter is illustrated in figure 5.1 B. These studies furthermore showed that the SNAP/SNARE interaction involves residues distributed along almost the whole stretch of the nine N-terminal αSNAP helices rather than being restricted to a small region. Even though these are interesting findings, this strategy is not useful when trying to distinguish between the different SNAP/SNARE interaction sites. A mutation in the αSNAPs itself might, but most probably will not, selectively inhibit one specific SNAP/SNARE interaction but would rather affect all three of them. Even if, theoretically, a mutant would interfere with binding at one specific SNARE site only, we would not be able to tell from the spectral data. A better strategy to ablate binding of αSNAP at only one or two sites, would therefore be to employ a mixture of αSNAP together with an αSNAP competitor, which competes for only one or two of the SNAP binding sites.

Complexins inhibit SNARE disassembly in vitro. One putative candidate for such a protein is Complexin. Complexins are small soluble proteins, which bind as an α-helix to the SNARE complex four-helix-bundle in an anti parallel fashion [39, 38, 41, 76, 42]. Additionally, single molecule studies have recently shown that Complexin may bind to and thereby stabilize Syntaxin/SNAP25 acceptor complexes [77]. However, it does not bind the individual SNAREs or the 2:1 SNARE complex
consisting of two Syntaxins and one SNAP25 as has been shown by isothermal titration calorimetry [41]. It is therefore highly probable that their functional target in the cell is the assembled SNARE complex and/or the Syntaxin/SNAP25 1:1 acceptor complex, but the precise role of Complexins is not understood up to date.

Complexins have originally been reported to displace αSNAP from the SNARE complex when added in excessive amounts, leading to speculations that they might have inhibitory properties with respect to SNARE disassembly [39]. Notwithstanding these findings, the gel-based in vitro assay monitoring SNARE disassembly introduced in section 4.1.2, did not show any negative influence of Complexin1 or Complexin2 on SNARE-complex disassembly [40]. However, due to a time resolution in the range of several minutes, these experiments are preferably suited for comparing ‘all-or-nothing’ disassembly defects rather than minor kinetic differences which could hence not be ruled out.

Recently, three different studies resulted in findings which all led to an interpretation according to which Complexin might act as a so-called fusion clamp, binding the SNARE complex in a partially zippered state prior to complete fusion and thereby ‘clamping’ it until the signal for the final fusion step is received [43, 44, 45]. If this were so, Complexin might preferentially act on ‘trans’ rather than ‘cis’ complexes and might conclusively act at a different stage in the SNARE cycle than αSNAP. Hence, even though Complexin seems to be able to replace αSNAP from SNARE complexes [39], a resulting inhibition of the disassembly in in vitro reaction would not necessarily imply that the two competing proteins ever meet in the cell and it would hence be difficult to draw physiological conclusions.

Anyhow, as pointed out above, in vitro experiments using Complexin might serve as a tool to address the question, whether less than three αSNAPs suffice to mediate efficient SNARE disassembly. I therefore tested SNARE disassembly in the presence of Complexin1 (see figure 4.39) using the fluorescence assays and found that the rate of disassembly indeed decreased. The inhibition dramatically increased under conditions in which αSNAP is limiting (figure 4.39 B and 4.40), and the ratio of αSNAP to Complexin was crucial for the degree of inhibition. These findings strongly suggest that the disassembly defect witnessed in presence of Complexin indeed results from a direct competition with αSNAP.

Evidence for SNARE disassembly mediated by less than three αSNAPs?

Conveniently, the structure of Complexin bound to the neuronal SNARE complex has been solved [42], enabling for speculations about the nature of the possible αSNAP/Complexin1 competition with respect to the different binding sites. The Complexin/SNARE-complex binding site overlaps with only one of the three hypothesized SNAP binding sites, according to the SNAP/SNARE model by Marz et al. discussed above. This of course does not necessarily mean that the other two SNAP binding sites are unaffected by Complexin binding, because mechanisms of cooperativity might complicate the picture of the molecular interactions at the SNAP binding sites.

Complexin was found to bind the SNARE complex with a very high affinity in
solution (less than 10nM) [41], suggesting that it should be dominant over αSNAP with respect to SNARE complex binding at the one overlapping binding site in solution. On membranes however, the affinity of Complexin to SNARE complexes might be much lower (∼680nM), as has been reported by Bowen et al [78] who performed single molecule FRET measurements on deposited lipid bilayers.

The fact that Complexin does not abolish disassembly completely, even though its SNARE binding site overlaps with one of the αSNAP’s, may be seen as a further evidence for the considerations that less than three αSNAPs per complex can mediate disassembly. On the other hand, the partial disassembly could also be due to Complexin binding to only a part of the complexes, the remainder of the complexes carrying three αSNAPs and thus being dissociated. Along these lines, a similar amount of inhibition would relate to a similar ratio of Complexin-containing to Complexin-free SNARE complexes. If the reasons for an intermediate inhibition were due to an equilibrium of αSNAP and Complexin at one site, one should hence be able to shift the equilibrium towards Complexin-bound complexes by increasing the Complexin concentration.

One general observation I made during the experiments however was that the degree of disassembly does not seem to be continuous, but rather seems to be restricted to discrete levels. For example, as shown in figure 4.41, even a five fold increase of Complexin (75nM to 370nM) at a constant αSNAP concentration (45nM) only slightly increases the degree of inhibition. Both of these different Complexin concentrations led to an intermediate block of SNARE disassembly. Likewise, lowering the αSNAP concentration to 22.5nM at 75nM Complexin results in identical speeds as 45.5nM αSNAP and 370nM of Complexin.

A stepwise dependence of disassembly speeds in presence of varying amounts of αSNAP and Complexin would match nicely with the idea, that discrete rates of disassembly might reflect a different number of αSNAPs bound. Depending on how close the affinities between the sites are to each other, the transitions might be more or less discrete at different occupation levels. Discrete levels however are difficult to reconcile with a view in which the reaction rate only relates to the portion of complexes occupied by Complexin.

These considerations might hence suggest that αSNAP is still able to mediate disassembly even though one SNAP/SNARE interaction site is blocked by Complexin. On the other hand, as already discussed for the SNARE deletion experiments, the site of Complexin binding might, even though it overlaps with one of the proposed αSNAP binding sites, not abrogate SNAP binding to this site completely but only partially. The partial binding of SNAP to the SNARE complex might still suffice for disassembly, maybe through a stabilizing effect mediated by the remaining SNAPs.

Since we do not know whether αSNAP binding to the different sites is cooperative, we do not know either, whether the physical ablation of one SNAP/SNARE interaction mechanistically influences the other SNAP/SNARE interaction. It is for instance conceivable that the affinities to the remaining αSNAP/SNARE sites increases after the first αSNAP has bound. If now this first site were occupied by Complexin, the two additional sites might display a lower affinity. The fact that
the impediment of disassembly is only partial however indicates that at least one αSNAP is still able to bind to the complex even when Complexin is present.

Interestingly, when very low amounts of αSNAP (22.5nM) are used, which in the absence of Complexin promote only slow disassembly, Complexin can even abolish disassembly completely. This may either indicate that the affinities as a matter of fact differ between the three SNAP binding sites, the Complexin binding site at the same time being the one displaying the highest affinity for αSNAP. If then this site were occupied by Complexin, αSNAP might not be able to bind the complex at all, when present only at concentrations sufficient for binding to the first binding site.

In summary, it is hence possible that the reduced rate of disassembly caused by Complexin reflects the kinetics of a reaction where only one or two as opposed to the possible three SNAPs are bound. This would, in accord with the SNARE complex mutation experiments discussed earlier, point to a possible existence of fine-tuning mechanisms of SNARE-complex disassembly, which act via a modulation of the number of αSNAPs bound per complex. Such a mechanism, which allows for the independent action of one or two SNAPs rather than depending on a concerted action of all three SNAP proteins, would resemble a feature recently discovered for another triple A ATPase called ClpX [27]. Here the authors could show that the single subunits of one ClpX-hexamer were capable to exert their catalytic function independently from each other, whereas a concerted action of several subunits apparently increased the enzymatic efficiency in a proportional dependence. If NSF also were to act in such an ‘asymmetric’ fashion, there might be a correspondent dependence of NSF activity on the number of SNAPs bound per SNARE complex.

However, all the inhibitory effects described might as well be explained by an influence of Complexin on αSNAP activity, which is not caused by the mere occupation of an αSNAP interaction site, which can not be excluded at this stage. In order to further address the model of potential discrete levels of αSNAP modulation depending on the amount of functional SNARE/SNAP interfaces, further Complexin- and αSNAP-titration experiments are necessary.

A modulatory rather than dominant negative function of Complexin as suggested by the experiments shown, might also help to abate some of the confusion amongst Complexin investigators. For a long time, seemingly contradictory outcomes of knock-out experiments and over-expression studies in various animals and cell-types have complicated the determination of Complexin function, leaving a range of proposed models in the field up to now. Most of these studies report a reduction of exocytosis in presence of excess Complexin, which would be in line with an negative impact on fusion. However, at the same time the absence of Complexin appears to result in impaired exocytosis as well. These apparently contradictory results could in principle also be explained by a fine-tuning capability which for instance might depend on a physiological Complexin concentration. If this is the case, too little as well as too much Complexin would harm the balanced fusion process and might lead to similar phenotypes.
αSNAP has a selective advantage over Complexin on liposomes as opposed to solution. Recently, three different studies resulted in findings which led to a similar interpretation, according to which Complexin might act as a so-called fusion clamp, binding the SNARE complex in a partially zippered state prior to complete fusion and thereby ‘clamping’ it until the signal for the final fusion step is received [43, 44, 45]. Even though this scenario primarily envisions Complexin to function as an inhibitor of the last step of SNARE assembly, it does not exclude the possibility that the ‘Complexin clamp’ serves as a protection against unwanted disassembly at the same time. If for instance, SNARE-complex assembly has almost completed, αSNAP might already recognize its binding sites and induce NSF-mediated disassembly of ‘trans’ complexes. At this stage, Complexin might have to efficiently compete with αSNAP. After fusion however, it would be logical if αSNAP were able to overcome the Complexin mediated inhibition of disassembly in order to allow for fast recycling of cis SNARE complexes.

In line with the observation that the αSNAP affinity is strongly enhanced on membranes compared to solution, whereas complexin apparently displays a higher affinity to SNARE complexes in solution than on membranes [41, 79], the ratio of Complexin to αSNAP needed to efficiently block SNARE disassembly in the fluorescence assays, differs significantly between experiments pursued in solution and on liposomes. In solution, sub-molar amounts of Complexin compared to αSNAP already impair disassembly. On liposomes, the amount of Complexin needed to inhibit disassembly is in the same range as in solution, whereas the power of αSNAP increases by a factor of ~ twenty due to the membrane potentiation. As opposed to αSNAP, complexin function hence does not seem to be boosted by the membrane anchorage SNARE complex.

Complexin vs. αSNAP at physiological αSNAP concentrations. According to my knowledge, there are no published data about the physiological concentration of Complexin1 and αSNAP in the cell, or more precisely, at the sites of SNARE action. According to semi-quantitative Western blotting carried out in our lab, the cellular αSNAP concentration was determined to be in the range of ~20-50nM (T. Lang, personal communication). This of course does not imply the αSNAP concentration at the membrane is identical. Yet it is likely that the low amounts (~100nM or less), seen to promote optimal disassembly on membranes but not in solution, are closer to physiological concentrations than the high amounts required for optimal disassembly in solution (~1µM or more). At 100nM αSNAP, as much as 370nM Complexin did not suffice to block disassembly on liposomes, whereas the same amount of Complexin even impairs disassembly mediated by more than 1µM αSNAP in solution. One might of course argue that the characteristics of disassembly in solution are anyways of no physiological significance. Still, what we do not know so far, is whether the membrane boost observed for αSNAP is only applicable when the complex resides in ‘cis’ and all three αSNAPs can thus attach to the same membrane, or whether the boost would also occur with regard to binding of complexes docked in a ‘trans’ state. If the anchorage of all SNAREs to
the same membrane were a prerequisite for the high αSNAP efficiency, Complexin would be dominant over αSNAP before, whereas in turn αSNAP would be superior after fusion. However, since we do neither know the physiological concentration of Complexin nor whether these two proteins ever meet during the SNARE cycle, this at the moment is mere speculation and requires further investigation.

**Summary of Complexin function in SNARE disassembly.** Taken together, the data show that Complexin inhibits SNARE disassembly and the amount of inhibition is dependent on the αSNAP concentration. Furthermore, the Complexin block is only partial at intermediate αSNAP concentrations but can be complete, when αSNAP is reduced to a minimum. At this point, no definite conclusions can be drawn with regard to whether Complexin binding completely abolishes one or more αSNAP/SNARE interaction sites. Finally, as opposed to αSNAP, Complexin does not show an increased affinity to liposomal ‘cis’ complexes over ones in solution, leading to an inversion of masteries between Complexin and αSNAP, depending on whether the target complex is on liposomes or in solution. Yet, soluble SNARE complexes do not represent a physiological target, so that a physiological relevance of possible interpretations of this finding can not be concluded.

5.2.3 Phosphorylation of NSF as a Regulatory Mechanism?

Another level of regulation might be modifications of the enzyme itself. Several reversible modification mechanisms have been proposed, which might have an impact on NSF activity. For instance, one NSF modification which has been suggested to regulate the disassembly reaction is phosphorylation at residue Y83. According to Huynh et al. [36], the defect is caused by a reduced affinity to αSNAP. To validate the notion that phosphorylation of NSF at this site alters SNARE disassembly, I employed *in vitro* phosphorylated NSF and a phosphomimetic NSF mutant NSF(Y83E) for disassembly in the *in vitro* assays. Indeed, when I tested the mutant NSF_Y83E in the gel based disassembly assay, the mutant seemed to perform less efficiently than the wild type (figure 4.1.2). Since however, as pointed out earlier, the gel experiment is only semi-quantitative and the reasons for a changed outcome are more difficult to assess, such a difference in band intensities does not suffice for mechanistic interpretations.

In this context, I disassembled different complexes, incorporated into the membrane either *via* the Syntaxin or the Synaptobrevin domain and containing either full length Syntaxin or only the Syntaxin H3 domain, to ensure that no defect only evident for one of these complexes would be overseen (figures 4.45 and 4.44).

**NSF_Y83E rapidly disassembles on liposomes.** Even though there were slight kinetic differences between wild type and NSF_Y83E on liposomes, increasing the mutant’s concentration by 50% completely abrogated the differences so that disassembly proceeded with the same kinetics as wild type NSF. A difference as subtle as that is not significant enough to definitely assign a defect to NSF phosphorylated
at position 83. The fact that such a low increase of the mutant concentration can overcome the differences, might for instance be due to slightly less active enzyme in the mutant’s preparation. This for instance might result from reduced stability of the mutant. At this stage it should again be pointed out that NSF is known to be a fragile enzyme and it is difficult to compare the preparations of the wild type and the point mutation quantitatively. It can never be excluded that differences between the enzymes stem from buffering conditions which even though optimized for NSF as described in section 4.1.3, do not mimic the enzymes natural environment. Furthermore, the conditions used in the fluorescence assays were optimized for NSF which does not necessarily imply that they are also the best possible for modified NSF variants, as NSF\textsuperscript{Y83E}. It can hence not be excluded that the purification procedures differentially affect the active state of the two enzymes, e.g. by changing the ratio of the hexameric to the monomeric pool and the subtle differences observed do therefore not necessarily represent physiological modulations. These experiments thus do not prove a significant change of disassembly kinetics upon phosphorylation of NSF.

\textbf{NSF\textsuperscript{Y83E} disassembly might be slightly impaired under certain conditions in solution.} In solution, phosphorylated NSF once more disassembled slower and less efficiently than wildtype when used at identical concentrations at 1,1\(\mu\)M \(\alpha\)SNAP (figure 4.46). As opposed to the liposomal findings however, this deficiency could not be overcome by more phosphorylated NSF. Instead, the defect was almost completely abrogated by increasing the \(\alpha\)SNAP concentration from 1,1\(\mu\)M to 2,2\(\mu\)M. In the gel-based disassembly assay, NSF\textsuperscript{Y83E} also performed less efficiently than wildtype NSF. I next examined, whether this is due to a reduced \(\alpha\)SNAP affinity of NSF\textsuperscript{Y83E} in solution, as has been suggested by Huynh et al. [36]. To do so, I performed disassembly experiments using NSF wt and NSF\textsuperscript{Y83E}, respectively, at very low \(\alpha\)SNAP concentrations. However, the kinetics of NSF and NSF\textsuperscript{Y83E}-disassembly in these experiments were similar (data not shown). Similarly, the use of very low enzyme concentrations (NSF or phosphorylated NSF, respectively) led to comparable kinetics between wildtype and mutant. A general lower \(\alpha\)SNAP affinity can thus be excluded, because this would have been evident under conditions in which \(\alpha\)SNAP is limiting.

\textbf{How may this be interpreted?} If there is a significant difference between the two mutants, it thus only seems to come to the fore in a specific range of \(\alpha\)SNAP concentrations. If one keeps in mind that the disassembly reaction using wildtype NSF is quite robust, phosphorylated NSF might for instance be more fragile with respect to alterations or inhibitory factors at the SNARE- or \(\alpha\)SNAP interaction site. One could speculate that the level of NSF-activity induced by \(\alpha\)SNAP might depend on the number or tightness of the \(\alpha\)SNAP molecules bound to the complex and this induction pattern might for instance be different after phosphorylation. It is for instance possible that complexes occupied by one or three \(\alpha\)SNAPs stimulate phosphorylated NSF as efficiently as wildtype NSF, whereas complexes occupied by
two αSNAPs induce the activity of phosphorylated to a smaller extent than that of wildtype NSF. Anyhow, since at this point I have not even been able to prove that wildtype disassembly is indeed possible with less than three αSNAPs, these are mere speculations and further experiments, covering a whole range of αSNAP concentrations, need to be performed in order to finally elucidate the function of NSF phosphorylation. Alternatively to a disassembly defect, kinetic differences of disassembly recordings in solution may always as well be caused by different speeds of reassembly. Instead of the proposed disassembly defect of phosphorylated NSF, it might also be less efficient than NSF in keeping the freed SNAREs from re-assembly or only disassemble up to a state of partial disassembly, from which re-assembly may occur faster. Since high αSNAP levels negatively influence SNARE complex assembly kinetics, the apparently better performance of NSF
\textsuperscript{Y83E} in the presence of high amounts of αSNAP might as well be caused by a reduction of re-assembly upon raising the αSNAP concentration. Since all these thoughts are speculative, further studies are essential to gain insights into whether the phosphorylation indeed leads to a mechanistic difference of the mutant or whether the mutant is simply less stable in my hands than the wildtype.

5.2.4 Conservation of the Reaction Mechanism in Yeast

The fact that αSNAP and NSF are ubiquitously expressed in all tissues and mediate the recycling of most (if not all) SNARE proteins involved in intracellular fusion processes points at a strong mechanistic conservation of the different disassembly reactions throughout the cell. Considering that the four-helix-bundle structures of the different SNARE complexes show a high degree of similarity, the overall sequence homology between the different complexes studied thus far is surprisingly low. It is hence possible that the SNAP/SNARE machinery does not care about conservation of specific residues but rather recognizes its target proteins on the basis of their quaternary structure. Several studies have already focused on the question, of whether even between two organisms as distant as yeast and mammals the degree of conservation is high enough to provide functionality of one disassembly machinery in the other organism. Some of these have come to the conclusion that single components of the machineries seem to be interchangeable [20], but others have come to apparently contradictory results [28, 49].

The homologues of αSNAP and NSF in yeast are Sec17 and Sec18, respectively. The structures of Sec17 as well as the Sec18 N-terminal domain have been solved crystallographically [30, 55]. In analogy to the neuronal SNARE complex, \textit{in vitro} binding studies of Sec17 yeast complex coupled to GST-beads revealed that the stoichiometry of Sec17 to SNARE complex is 3:1 [53]. The crystal structures of the yeast and the neuronal SNARE complex as well as of NSF and the Sec18 N-terminal domains reveal a remarkable degree of similarity [54, 55, 56, 29], suggesting that the mode of interaction with their respective SNAP adapters might also be conserved. NSF and Sec18p may however not be directly interchangeable, because NSF is unable to complement the Sec18-1 mutation in yeast [49]. Nevertheless,
functional homology between NSF and Sec18p has been proposed after it was shown that yeast cytosols can reconstitute NSF activity to NEM-treated mammalian \textit{in vitro} transport assays \cite{57,80}. Yeast cytosol yet might include additional factors to Sec18, so that the recovery here can not be reliably be ascribed to Sec18 function. When studying the interchangeability of the adapter proteins, yeast cytosol was unable to bind NSF to mammalian Golgi-membranes \cite{28}, but Sec17 could replace SNAP activity in mammalian intra-Golgi assays \cite{49}. NSF efficiently bound to plastic-coupled Sec17 and Sec18 to $\alpha$SNAP \cite{20}. Some functional homology thus seems to exist, but to which extent and whether for all of the proteins involved in the mechanism, remains unclear. No time-resolved experiments have yet been made, which have directly assessed the influence of exchanging yeast and mammalian proteins during SNARE complex disassembly.

I hence disassembled yeast SNARE complexes using components of the mammalian machinery and \textit{vice versa}. Additionally, I used various combinations of the yeast and the mammalian components, to determine which of the parts or protein interfaces are especially conserved and whether the mechanisms found for the neuronal SNARE complex (as e.g. the ‘membrane boost’) are conserved in yeast.

A quantitative comparison of the actual complex turnover by the different combinations of disassembly machineries could not be made, because Sec18 as well as Sec17 turned out to be less stable under the purification conditions used than $\alpha$SNAP and NSF. It soon became evident that both, Sec17 as well as Sec18, tend to loose activity very fast (in a day or two from purification). The experiments using the yeast machinery, therefore do not provide absolute activities or rates of disassembly. Nevertheless, qualitative interpretations can be made and some questions regarding the grade of conservation of the disassembly mechanism could be resolved merely on the basis of comparative measurements.

**Target recognition is highly conserved.** Strikingly, both the yeast and the disassembly machinery were able to disassemble both SNARE complexes (figures 4.49 and 4.47 A). For both complexes, the yeast machinery disassembles less efficiently than the mammalian. Whether this is only due to a lower stability in the purification buffers or also has a physiological meaning remains unclear at this point.

Interestingly, both machineries dissociated their non-cognate complexes comparably fast and efficiently as their respective cognate complexes. This finding is in line with the hypothesis that rather the quaternary structure than the sequence conservation may be important for target recognition.

**The adapter/enzyme interface is highly conserved.** Disassembly of neuronal complexes by the yeast machinery (Sec17 and Sec18) and the combination of Sec18 and $\alpha$SNAP worked equally efficient. The efficient disassembly mediated by $\alpha$SNAP and Sec18 indicates that the interface and basic mechanism on the level of the adapter-enzyme interplay seem to be conserved (figure 4.47). In line with
that proposal, Sec17 and NSF also efficiently disassembled the yeast complex, again pointing to a conservation of the adapter/enzyme interface.

**Two non-cognate protein interactions strongly impair disassembly.** Interestingly, the combination of Sec17 and NSF did not promote disassembly of the neuronal complex when used at the same concentrations as in the other combinations which all led to efficient disassembly (figure 4.47 B). Similarly, the combination of αSNAP and Sec18 did not disassemble the yeast complex under the same conditions used in the other experiments (figure 4.50 B).

It can be excluded that the interfaces between Sec17 and NSF as well as between αSNAP and Sec18 have diverged so much during evolution, that enzyme and adapter cannot productively interact any longer, because these combined machineries should not be able to disassemble any SNARE complexes then. Both combinations however were able to promote disassembly of the respective other SNARE complex, the neuronal in the case of Sec18/αSNAP and the yeast in the case of NSF/Sec17 as described in the last paragraph.

If one pictures the two connections between SNARE complex, SNAP adapter and enzyme, one notices that in both of these inefficient combinations both of the protein/protein interfaces are between non-cognate proteins, whereas all the other combinations discussed so far included only one non-cognate interaction interface. All these findings indicate that each of the protein/protein interfaces involved in the disassembly process is conserved enough to allow for exchange of proteins between yeast and mammals at one interface, but exchanges at two interfaces lead to synergistic defects of disassembly. The exchange of only the adapter protein, which brings about two non-cognate interfaces, therefore strongly impedes disassembly.

The membrane boost is conserved. The membrane boost observed for αSNAP was found to be conserved for the yeast adapter protein Sec17. An amount of Sec17 and NSF, which did not at all disassemble SNARE-complexes in solution, did mediate disassembly on liposomes (4.48). It can thus be concluded that like for αSNAP, Sec17-efficacy is higher when the target complexes are incorporated into membranes. Similarly, the high efficiency of αSNAP when disassembling membrane-bound complexes can as well be observed for the disassembly of yeast complexes. Here, like for the neuronal complexes, 120nM of αSNAP suffice for efficient disassembly (see figure 4.49), when the yeast complex is incorporated into liposomes. This implies that the potentiation of αSNAP efficacy on membranes is independent of the neuronal SNARE complex as substrate.

Summary. It can be concluded that both of the protein interfaces between target, adapter and enzyme seem to be conserved well enough between yeast and mammals to allow proteins of the respective other organism to mediate disassembly. Notwithstanding the fact that disassembly occurs even with exchanged machineries, the kinetics of disassembly may be slightly impeded by two non-cognate proteins
at any of the interaction interfaces, whereas they are strongly hampered when both interactions are mediated by non-cognate proteins. Additionally, the high αSNAP efficiency on membranes described earlier is independent of the nature of the SNARE target and the membrane boost conserved for the yeast adapter protein Sec17.
Chapter 6

6.1 Summary

SNAREs (SNAP receptors) are small membrane-bound proteins which are key mediators of intracellular membrane fusion. The actual fusion step is believed to occur through complex formation of SNAREs sitting on opposing membranes, first bringing the membranes into close apposition and finally enabling them to fuse. SNARE complex spontaneously form upon mixing of the single SNAREs in solution and are extremely stable. After membrane merger, the complexed SNAREs need to be dissociated and recycled again to allow for further rounds of fusion. Since the dissociation of SNARE complexes requires energy, this so-called SNARE disassembly is mediated by an enzyme, the AAA ATPase NSF (N-ethyl maleimide sensitive fusion protein) and its cofactors αSNAP. Presumably three αSNAPs bind to one SNARE complex to form a binding site for the hexameric NSF. This then binds to the αSNAPs and catalyzes the reaction via hydrolysis of ATP.

The aim of this study was to investigate the molecular basis of SNARE disassembly including SNAP isoforms, putative regulatory factors and the degree of functional conservation. To this end, I established two fluorescence assays (based on FRET and fluorescence anisotropy) which allow for monitoring of the disassembly reaction in vitro. During the characterization of the disassembly reaction, it soon became evident that the recombinant αSNAP performed far less efficient when disassembling soluble SNARE complexes using these methods, than it did during disassembly of ‘ex vivo’ SNARE complexes using so-called plasma membrane sheets. The use of liposome-incorporated SNARE complex for disassembly sufficed to annihilate the observed discrepancy between the ex vivo and the in vitro preparation, indicating that the lipid membrane increases the potency of αSNAP during SNARE complex disassembly. I could further demonstrate, that an N-terminal deletion in the αSNAP molecule abolishes the membrane caused potentiation of efficacy observed for αSNAP. This finding suggests, that the N-terminus of αSNAP interacts with the membrane to increase the affinity or effectivity of αSNAP during disassembly. In addition to the long established interaction between αSNAP and the membrane-bound SNARE complex, the membrane thus serves as a second SNAP receptor which increases αSNAP efficiency. Additionally I demonstrated that the potentiation of SNAP efficiency in the presence of membranes is conserved in the
SNAP isoform βSNAP and in yeast.

Furthermore I performed disassembly reactions using different conditions, mutations and putative inhibitory proteins in order to gain molecular insights into the protein/protein interactions involved and putative regulatory mechanisms of SNARE disassembly. An antibody against the NSF N-terminal domain raised in the scope of this study eliminated SNARE disassembly completely and might serve as a useful tool to specifically block NSF action in in vivo experiments in the future.

Three anti SNARE complex antibodies blocked disassembly to a certain extent but a complete block was possible only when two antibodies, one recognizing the N-terminal end and the other recognizing the C-terminal end, were used in concert. Similarly, other SNARE complex mutants only abrogated SNARE disassembly when two alterations were introduced at non-proximal sites. These data indicate that the disassembly process is very robust and the disassembly machinery does not have a preference for either end of the complex.

The SNARE-complex interacting protein Complexin1 significantly inhibited SNARE-complex disassembly in the fluorescence assays in an αSNAP concentration-dependent manner. This indicates that Complexin and αSNAP compete with each other for SNARE complex binding under in vitro conditions. The inhibition of disassembly by Complexin was more pronounced in solution than on liposomes, suggesting that Complexin affinity to the SNARE complex in the presence of membranes is not increased to the same extent as witnessed for αSNAP. A phosphorylation mutant of NSF, which has previously been suggested to have a reduced affinity to αSNAP, could not be definitely shown to exert a significant defect with regard to the disassembly reaction.

I further showed, that between yeast and mammals, both of the protein interfaces between target, adapter and enzyme seem to be conserved well enough to allow proteins of the respective other organism to mediate disassembly. Notwithstanding the fact that disassembly occurs even with exchanged machineries, the kinetics of disassembly may be slightly impeded by two non-cognate proteins at any of the interaction interfaces, whereas they are strongly hampered when both interactions are mediated by non-cognate proteins. Additionally, the high αSNAP efficiency on membranes described earlier is independent of the nature of the SNARE target and the membrane boost conserved for the yeast adapter protein Sec17.
6.2 Zusammenfassung

Für die Fusion intrazellulärer biologischer Membranen sind Vertreter der konservierten Familie der SNARE-Proteine von essentieller Bedeutung. Man vermutet, dass gerichtete Komplexbildung zwischen SNAREs auf gegenüberliegenden Membranen dazu führt, dass die zu fusionierenden Membranen in räumliche Nähe gezogen werden und schließlich verschmelzen. SNARE-Komplexe bilden sich spontan, wenn die einzelnen SNARE-Komponenten in Lösung vermengt werden und sind äußerst stabil. Es muss daher Energie aufgewendet werden, um die SNARE-Komplexe nach der Membranfusion wieder in ihre Einzelteile zu dissozieren um damit die einzelnen SNAREs für weitere Fusionereignisse zur Verfügung zu stellen. Deshalb wird die Dissoziation der SNARE-Komplexe durch ein Enzym, die AAA ATPase NSF sowie ihren Kofaktor αSNAP, vermittelt. Dazu binden vermutlich drei αSNAP-Moleküle einen SNARE-Komplex und bilden so ein Podest, das als Angriffsstelle für das hexamere, ringförmige NSF dient.


Außerdem habe ich mithilfe verschiedener Mutanten und Inhibitoren mögliche regulatorische Angriffspunkte charakterisiert. Ein Antikörper gegen die N-terminal Domäne von NSF, der auch im Rahmen dieser Arbeit hergestellt wurde, vermochte die Dissoziationsreaktion komplett zu blockieren und könnte zukünftig für in vivo-Experimente eingesetzt werden. Drei Antikörper gegen den SNARE-Komplex hemmten die Dissoziationsreaktion teilweise, aber eine komplette Inhibition gelang nur, wenn zwei Antikörper gemeinsam eingesetzt wurden, einer gegen den N-Terminus und
der andere gegen den C-Terminus von αSNAP. Analog dazu blockierten SNARE-Komplex-Mutanten die Dissoziationsreaktion nur, wenn zwei entfernte Positionen mutiert worden waren. All dies lässt darauf schließen, dass die Reaktion sehr robust ist.

Das SNARE-Komplex-bindende Protein Complexin1 inhibierte die Reaktion in Abhängigkeit von der αSNAP-Konzentration. Die Ergebnisse zeigen, dass Complexin und αSNAP unter in vitro-Bedingungen miteinander um die SNARE-Komplexbindung konkurrieren. Der inhibitorische Effekt von Complexin war in Lösung stärker als auf Liposomen, sodass davon ausgegangen werden kann, dass die Complexin-Affinität zum SNARE-Komplex sich in Gegenwart von Membranen nicht im selben Maße steigert wie die von αSNAP.

Schließlich konnte ich zeigen, dass beide Protein-Schnittstellen zwischen Substrat, Adaptor und Enzym zwischen Hefen und Säugern stark genug konserviert sind, um den Austausch einzelner Komponenten zu ermöglichen, ohne dass die Reaktion zum Erliegen kommt. Dabei führt der Austausch des Adaptors zu einem weitaus stärkeren Defekt als der Austausch des Enyzms oder des Komplexes. Offenbar wird die Reaktion demnach nur dann schwer gestört, wenn sich an beiden Schnittstellen Proteine aus verschiedenen Organismen treffen.
Bibliography


Chapter 7

Appendix

List of Publications

Marcin Barszczewski, John J. Chua, Alexander Stein, Ulrike Winter, Rainer Heintzmann, Felipe E. Zilly, Dirk Fasshauer, Thorsten Lang, and Reinhard Jahn:  
A Novel Site of Action for \( \alpha \text{SNAP} \) in the SNARE Conformational Cycle Controlling Membrane Fusion  

Ulrike Winter and Dirk Fasshauer:  
Mechanism of SNARE complex assembly and disassembly  
Chapter in ‘SNARE proteins’, Landes Bioscience, in press

Dana Bar-On*, Ulrike Winter*, Esther Nachliel, Menachem Gutman, Uri Ashery, Dirk Fasshauer and Thorsten Lang:  
Cortical membrane sheets  An ‘ex vivo’ platform for imaging kinetics of plasmalemmal protein interactions  
submitted.

* These authors contributed equally
7.1 Abbreviations and Symbols

<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Description</th>
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<tbody>
<tr>
<td>aa</td>
<td>Amino acid</td>
</tr>
<tr>
<td>APS</td>
<td>Ammonium persulfate</td>
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<tr>
<td>ATP</td>
<td>Adenosine triphosphate</td>
</tr>
<tr>
<td>BSA</td>
<td>Bovine serum albumine</td>
</tr>
<tr>
<td>cmc</td>
<td>Critical micelle concentration</td>
</tr>
<tr>
<td>DOM</td>
<td>Dodecylmaltoside</td>
</tr>
<tr>
<td>DTT</td>
<td>Dithiothreitol</td>
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<tr>
<td>DNA</td>
<td>Deoxyribonucleic acid</td>
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<tr>
<td>E. coli</td>
<td>Escherichia coli</td>
</tr>
<tr>
<td>EDTA</td>
<td>Ethylenediaminetetraacetic acid</td>
</tr>
<tr>
<td>FRET</td>
<td>Fluorescence resonance energy transfer</td>
</tr>
<tr>
<td>GST</td>
<td>Glutathion-S-transferase</td>
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<tr>
<td>GTP</td>
<td>Guanosine triphosphate</td>
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<tr>
<td>IgG</td>
<td>Immunoglobulin G</td>
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<tr>
<td>IAA</td>
<td>Indole acryl acetic acid</td>
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<tr>
<td>IPTG</td>
<td>Isopropyl-β-D-thiogalactoside</td>
</tr>
<tr>
<td>kDa</td>
<td>Kilo Dalton</td>
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<tr>
<td>LB</td>
<td>Luria Bertani medium</td>
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<tr>
<td>mS</td>
<td>Milli Siemens</td>
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<tr>
<td>NEM</td>
<td>N-ethyl-maleimide</td>
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<tr>
<td>Ni-NTA</td>
<td>Nickel-nitrilo-triacetate</td>
</tr>
<tr>
<td>NSF</td>
<td>N-ethylmaleimide-sensitive factor</td>
</tr>
<tr>
<td>PAGE</td>
<td>Polyacrylamide gel electrophoresis</td>
</tr>
<tr>
<td>PBS</td>
<td>Phosphate buffered saline</td>
</tr>
<tr>
<td>PCR</td>
<td>Polymerase chain reaction</td>
</tr>
<tr>
<td>PC</td>
<td>Phosphatidylcholine</td>
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<tr>
<td>PE</td>
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<tr>
<td>PMSF</td>
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<tr>
<td>PS</td>
<td>Phosphatidylserine</td>
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<tr>
<td>pH</td>
<td>Negative logarithm of H⁺ concentration</td>
</tr>
<tr>
<td>rpm</td>
<td>Revolutions per minute</td>
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<tr>
<td>Abbreviation</td>
<td>Description</td>
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<td>RT</td>
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<td>Sb</td>
<td>Synaptobrevin</td>
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<td>Sx</td>
<td>Syntaxin</td>
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<tr>
<td>SDS</td>
<td>Sodium dodecyl sulfate</td>
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<tr>
<td>SEM</td>
<td>Standard error of the mean</td>
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<tr>
<td>SNAP</td>
<td>Soluble-NSF-attachment protein</td>
</tr>
<tr>
<td>SNAP25</td>
<td>Synaptosome-associated protein of 25 kDa</td>
</tr>
<tr>
<td>SNARE</td>
<td>SNAP-receptor</td>
</tr>
<tr>
<td>TB</td>
<td>Terrific broth</td>
</tr>
<tr>
<td>TEMED</td>
<td>N,N,N’,N’-tetramethylethylene diamine</td>
</tr>
<tr>
<td>TMD</td>
<td>Transmembrane domain</td>
</tr>
<tr>
<td>Tris</td>
<td>Trisand-aminomethane</td>
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<tr>
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I hereby declare that the Ph.D. thesis entitled ‘Biophysical Characterization of SNARE complex Disassembly Catalyzed by NSF and αSNAP’ has been written independently and with no other sources and aids than quoted.

Hiermit erkläre ich an Eides statt, dass ich die vorliegende Arbeit selbst und ohne weitere als die durch Zitate angegebene Hilfe angefertigt habe.

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