The Role of the N-D1 Linker of the N-Ethylmaleimide-Sensitive Factor in the SNARE Disassembly

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Abstract

N-ethylmaleimide-sensitive factor (NSF) is a member of the type II AAA+ (ATPase associated with various cellular activities) family. It plays a critical role in intracellular membrane trafficking by disassembling soluble NSF attachment protein receptor (SNARE) complexes. Each NSF protomer consists of an N-terminal domain (N domain) followed by two AAA ATPase domains (D1 and D2) in tandem. The N domain is required for SNARE/α-SNAP binding and the D1 domain accounts for the majority of ATP hydrolysis. Little is known about the role of the N-D1 linker in the NSF function. This study presents detailed mutagenesis analyses of NSF N-D1 linker, dissecting its role in the SNARE disassembly, the SNARE/α-SNAP complex binding, the basal ATPase activity and the SNARE/α-SNAP stimulated ATPase activity. Our results show that the N-terminal region of the N-D1 linker associated mutants cause severe defect in SNARE complex disassembly, but little effects on the SNARE/α-SNAP complex binding, the basal and the SNARE/α-SNAP stimulated ATPase activity, suggesting this region may be involved in the motion transmission from D1 to N domain. Mutating the residues in middle and C-terminal region of the N-D1 linker increases the basal ATPase activity, indicating it may play a role in autoinhibiting NSF activity until it encounters SNARE/α-SNAP complex substrate. Moreover, mutations at the C-terminal sequence GIGG exhibit completely abolished or severely reduced activities of the substrate binding, suggesting that the flexibility of N-D1 linker is critical for the movement of the N domain that is required for the substrate binding. Taken together, these data suggest that the whole N-D1 linker is critical for the biological function of NSF to disassemble SNARE complex substrate with different regions responsible for different roles.

Introduction

N-ethylmaleimide-sensitive factor (NSF) is one of the first members of AAA+ (ATPase associated with various cellular activities) family and is identified to play a role in vesicular trafficking [1]. The soluble NSF attachment protein receptor (SNARE) proteins are the minimal machinery for membrane fusion [2], and they form parallel, four-helix, coiled-coil SNARE motifs that span the opposing membranes [3–5]. The assembly and disassembly of SNARE complex are required for constant vesicular transport. Though other diverse functions for NSF are found [6,7], the well-confirmed function of NSF is using energy from ATP hydrolysis to disassemble SNARE complex based on its interaction with an adaptor protein, soluble NSF attachment protein (α-SNAP) [8–10].

NSF is a ring-shaped homohexameric protein [11], with each protomer consisting of three domains: an N-terminal domain (N domain) and two highly conserved AAA+ domains (D1 and D2 domains) [12,13]. The N-domain is a positively charged domain in the surface that is important for SNARE/α-SNAP binding [14,15]. The D1 domain is the major ATPase domain providing the chemical energy by ATP hydrolysis required for the SNARE complex disassembly, while the D2 domain is a degenerate ATPase domain primarily responsible for maintaining NSF as a hexamer [12,16]. The NSF protein has a relatively very low intrinsic ATPase activity [13,17]. NSF-N domain has been proposed to exert some control over NSF ATPase activity because antibodies against to it cause a two-fold increase in hydrolytic activity [17,18]. NSF-D1 domain has a lower affinity for ATP (Kd = 15–20 μM) and accounts for the majority of basal and SNARE/α-SNAP stimulated ATPase activity [16,19,20]. As compared to NSF-D1 domain, NSF-D2 domain has a significantly higher affinity for ATP (Kd = 30–40 nM) but contributes no significant ATPase activity for NSF [19]. These data demonstrate the importance of the N and D1 domains for NSF ATPase activity and thus for their functions in SNARE complex disassembly.

There are two linker regions connecting the N and D1 domains, and the D1 and D2 domains, called the N-D1 linker and the D1-D2 linker, respectively. The NSF N-D1 linker is a highly conserved 20-residue linker, consisting of amino acid residues 203–222 (Fig. 1). Two conserved glycine residues (G221 and G222), present at the C-terminal region of the NSF N-D1 linker, can potentially provide the N-D1 linker with dynamic properties similar to the features seen in the linker of p97 [21,22]. Structurally the glycine residues are in close proximity to the nucleotide-binding site in p97-D1 domain and could be sensitive to the state of the bound nucleotide [21,23]. Structural studies of NSF in different nucleotide states showed that the protein undergoes conformation changes during nucleotide binding and hydrolysis, most notably with NSF-N domain changing its
disposition relative to the rest of the NSF hexamer [24,25]. Presumably, communication between the N and D1 domains couples complex assembly/disassembly with the ATP hydrolytic cycle. Despite the functions of the N and D1 domains are clearly demonstrated, little is known about the roles of N-D1 linker for NSF function.

In this study, we examined the role of NSF N-D1 linker using site-directed mutagenesis by monitoring four key activities as follows: the SNARE disassembly, the SNARE/\(\alpha\)-SNAP complex binding, the basal ATPase activity and the SNARE/\(\alpha\)-SNAP stimulated ATPase activity. Our results suggest that the whole NSF N-D1 linker actually plays a critical role in the biological function of NSF to disassemble SNARE complex with different regions responsible for different roles.

Results

Rationale for site-directed mutagenesis of the NSF N-D1 linker

The previous structural comparison and biochemical analysis suggested an important role for the linker between domains in AAA+ proteins [26–30]. To investigate the role of the N-D1 linker in NSF, we first compared the sequence similarity of this linker from several different eukaryotic species. As shown in Fig. 1, this linker is highly conserved from yeast to human. Strikingly, the “GIGG” sequence at the C-terminus of the linker is the most highly conserved, implying its functional importance for NSF. Mutating each residue to alanine, which is termed “alanine scanning”, is a useful strategy for identifying functionally important residues in proteins. Additionally, we mutated both glycine residues at the C-terminus to the less flexible proline residues, which might greatly impair the dynamic property of the N-D1 linker.

N-D1 linker mutants have the same oligomeric state as wild-type NSF

Plasmids expressing wild-type NSF and N-D1 linker mutants were introduced into E. coli BL21 (DE3). The proteins were initially purified by Ni-NTA superflow affinity chromatography, then further purified by gel filtration chromatography using a Superdex-200 column. The final purity was analyzed by the SDS-PAGE gel, showing the high purity of each protein (Fig. 2A).

The purified protein was then applied to the Superose-6 column to assess its oligomeric state. All of the mutants showed essentially the same fractionation profile as wild-type NSF, as was evidenced by the typical elution profiles of wild-type NSF and some N-D1 linker mutants shown in Fig. 2B. Excepting the small void volume peak, which contains irreversibly aggregated proteins, the main volume pick is hexameric NSF. This indicates that introduction of the mutations into NSF did not change their oligomeric states.

The N-terminal region mutants of the NSF N-D1 linker cause significant defects in the disassembly of the SNARE complex

We first assessed the effect of the mutations at the N-terminal region of the N-D1 linker on the biological function of NSF to disassemble SNARE complex. To assay for this disassembly activity, SNARE complexes were preincubated with wild-type NSF or the mutants and \(\alpha\)-SNAP on ice, following the addition of 2 mM Mg\(^{2+}\)-ATP. The reactions were performed at 37°C for the indicated times, then immediately stopped with the addition of the SDS-PAGE loading buffer and analyzed by SDS-PAGE. As expected, the amount of SNARE complexes progressively decreased with the increased incubation time (Fig. S1). But all of the mutants, E203A, N204A, Q206A, S207A, I208A, I209A and N210A, had severe defect. None of them were able to disassemble the SNARE complex to the extent of wild-type NSF (Fig. 3A&S1, Table 1). Among them, the N210A mutant shows the most...

Figure 1. Domain sketch of NSF protein and the amino acid conservation of the N-D1 linker. The diagram shows the structural domains of NSF: the N-terminal domain, D1 and D2 AAA+ modules. The position of the N-D1 linker and D1-D2 linker are indicated. Sequence alignment of the N-D1 linker from different eukaryotic species shows that the N-D1 linker is highly conserved. The numbers at the two ends of the sequence indicate the amino acid residues in NSF. Residues are colored by similarity. The letters under the sequence indicate the amino acid substitutions constructed in this study.

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seriously defective in disassembly efficiency, only 6% of that of wild-type NSF (Fig. 3A, Table 1).

Since the defect in the SNARE disassembly could be due to the defect in the SNARE/α-SNAP binding, we next examined the ability of these mutants to bind to the SNARE/α-SNAP complex in the ADP-AlFx (A transition state analog for the hydrolysis step) state. For this binding assay, equal amounts of wild-type or mutant NSF proteins were preincubated with SNARE complex, with or without α-SNAP, in the presence of ADP-AlFx. The binding complexes were recovered with amylose magnetic beads (MBP-VAMP), and the bound proteins were analyzed by SDS-PAGE as shown in Fig. 3B. These mutations showed either no effects (N204A and S207A) or partially reduced effects (E203A, Q206A, I208A, I209A, and N210A) on binding activity compared with wild-type NSF. This indicates that, when fixed at the ADP-AlFx state, the N-terminal region of N-D1 linker mutants can attain the conformation required for SNARE/α-SNAP complex binding.

Then, the basal and stimulated ATPase activity was investigated. Previous studies demonstrated that NSF has a low basal ATPase activity, which is stimulated 2–10-fold by the binding of SNARE/α-SNAP complex [16,31,32]. NSF D1 and N domains account for the majority of the basal and SNARE/α-SNAP stimulated ATPase activity [18,33]. But we do not know the influence of N-D1 linker in the ATPase activity. In our test condition, wild-type NSF produced inorganic phosphate at a rate of 0.53 nmol/μg/h, and SNARE/α-SNAP complex stimulated a 9.8-fold increase in the ATPase activity (Fig. 3C), which are consistent with the published data [13,31]. All of the mutations showed no or partial effects on the basal ATPase activities compared with wild-type NSF (Fig. 3C, Table 1), indicating that they can bind ATP. This is in agreement with their normal binding activities with the substrate (Fig. 3B), which requires the protein in the ATP-bound state. But surprisingly, these mutations also exhibit stimulated ATPase activities similar to that of wild type NSF. The apparently observation that all the mutations had normal ATPase activities suggests that the residues at the N-terminal region of N-D1 linker have no significant effect on ATP hydrolysis.

In summary, the mutations at the N-terminal region of N-D1 linker (E203A, N204A, Q206A, S207A, I208A, I209A and N210A) had limited effects on the substrate binding activity and ATPase activity compared with their effects on the SNARE disassembly.

The middle region mutants of the NSF N-D1 linker have increased basal ATPase activity

To determine the role of the middle region of the N-D1 linker in NSF activity, we prepared the individual mutations encompassing this region by mutating each residue from Pro211 to Met218 to alanine, and then examined the effects of these mutations on the
SNARE disassembly, the SNARE/α-SNAP complex binding, the basal and stimulated ATPase activity.

Similar to the mutations at the N-terminal region, all of the mutants showed reduced abilities to disassemble SNARE (Fig. 4A, Table 1), indicating the middle region is also important for the biological function of NSF. Except for E216A, all other mutants did clearly bind to the SNARE/α-SNAP complex, although to different extents (Fig. 4B). And all the mutants tested have no or marginal effects on the SNARE/α-SNAP stimulated ATPase activity compared with wild-type NSF (Fig. 4C, Table 1). However, all of these mutations result in the elevated basal ATPase activity (Fig. 4C, Table 1). In particular, the P211A,
D212A, W213A, F215A, E216A and M218A mutants showed the substantial increase, three times more than wild-type activity (Fig. 4C, Table 1). The basal ATPase activity of the M218A mutant even increased to the extent that is higher than the stimulated ATPase activity of wild-type NSF (Fig. 4C). These results suggest that the residues within the middle region linker may play an important role in helping maintain the conformation of NSF for the low basal ATPase activity.

The C-terminal region mutants of the NSF N-D1 linker exhibit severe defect in the SNARE/α-SNAP complex binding activity

As demonstrated above, the N-D1 linker of NSF protein is highly conserved from yeast to human (Fig. 1). Especially, the C-terminal region is almost identical with a 4-residue-long GIGG sequence containing a Gly-Gly motif (Fig. 1). From the structural studies of NSF, it is confirmed that the flexible nature of N domain is important for conformational changes of NSF during the ATP hydrolytic cycle [24,25]. The multiple glycines in the C-terminal region which could impart the conformational flexibility to the N-D1 linker suggest that C-terminal region of the N-D1 linker may modulate the mobility of the NSF-N domain, and affect the NSF function as a consequence. To test this hypothesis, we mutated the residues to Ala, and additionally two adjacent Gly residues to Pro to further decrease the flexibility (Fig. 1).

As shown in Fig. 5A & Table 1, all of the mutations abrogate the ability to disassemble SNARE complex. Similar to the mutations at the middle region of the N-D1 linker, the C-terminal region mutants exhibit an increase in the basal ATPase activity, and
marginal impact on the stimulated ATPase activity (Fig. 5C, Table 1). But different from the middle region mutants, these mutants showed severe defect in the SNARE/α-SNAP complex binding activity (Fig. 5B). Three mutants, G219A, I220A, and G221PG222P, fail to bind the substrate, and mutant G221AG222A retains only very weak binding activity compared with wild-type NSF (Fig. 5B), in the ADP-AlFx bound state. This is consistent with our data that the basal ATPase activities of these mutants are similar to their SNARE/α-SNAP stimulated ATPase activities, namely, there are no substrate stimulation effects on these mutants due to their defects in the substrate binding (Fig. 5B&C, Table 1).

**Discussion**

Although much research has been conducted since the NSF was first identified [1], the role of the N-D1 linker of NSF is still unclear due to the lack of the structural studies. Here, we present detailed mutagenesis analyses of the N-D1 linker by examining four key activities: the SNARE disassembly, the SNARE/α-SNAP complex binding, the basal ATPase activity and the SNARE/α-SNAP stimulated ATPase activity. Our results suggest that the whole N-D1 linker is critical for the biological function of NSF to disassemble SNARE complex substrate with different regions responsible for different roles.
The N-terminal region of the NSF N-D1 linker may be involved in the motion transmission from the D1 to the N Domain

Recent structural studies of NSF in different forms of nucleotides by electron microscopy suggested that upon the ATP hydrolysis, the NSF-D1 domain rotated anticlockwise relative to the NSF-D2 domain, which was coupled with the up-to-down movement of the NSF-N domain [24,25]. Since the NSF-D1 domain is the major ATPase domain [12], the conformational changes of the NSF-D1 domain induced by the ATP hydrolysis must be transmitted to the NSF-N domain. The N and D1 domains of NSF are connected by the highly conserved N-D1 linker which is implied to play an important role in the communication between these two domains despite the lack of experimental evidence. In this study, we present the biochemical evidence to support the hypothesis that the N-D1 linker, especially its N-terminal region, may play a vital role in the motion transmission from the D1 to the N domain.

First, we demonstrated the functional importance of the N-terminal region of the N-D1 linker by evaluating the mutational effects of the residues in this region on the SNARE disassembly. As shown in Fig. 3A & Table 1, all of the mutations, E203A, N204A, Q206A, S207A, I208A, I209A and N210A, markedly reduced their abilities to disassemble the SNARE complex substrate without exception, indicating the essential role of the N-terminal region of the N-D1 linker in maintaining the functional state of NSF. Second, we examined the activities accounting for the biological function of NSF, including the oligomeric structure, SNARE/α-SNAP complex binding, the basal and the SNARE/α-SNAP stimulated ATPase activity, of these mutants. Surprisingly, these mutants form hexamers (Fig. 2B) and bind to the substrate similar to wild-type NSF (Fig. 3B), and had only limited effects on the basal and the SNARE/α-SNAP stimulated ATPase activity (Fig. 3C, Table 1) compared with their effects on the SNARE disassembly (Fig. 3A, Table 1). Thus, it is reasonable to deduce that the dramatically diminished biological function of these mutants is due to the defect in the motion transmission from the D1 to the N domain.

Further structural and structure-based mutagenesis analyses will be required to elucidate the detailed mechanism of how the N-D1 linker, especially its N-terminal region, transmits the motion from the D1 to the N domain.

The middle and C-terminal region of the NSF N-D1 linker may play a role in autoinhibiting NSF ATPase activity until it encounters the SNARE/α-SNAP complex

One striking observation of the mutations in the middle and C-terminal region of the N-D1 linker, except for K217A, is the
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Mutations at the C-terminal sequence GIGG of the NSF N-D1 linker do have ATPase activities (Fig. 5C, Table 1), indicating that they can bind to ATP. Thus, their negative effects on the substrate binding are not because they cannot attain the ATP-bound state. Recent structural study shows that the NSF-N domain is flexible, which undergoes the nucleotide-dependent up-to-down movement during the ATP hydrolysis process, namely, it is in an up conformation in the ATP state and in a down conformation in the ADP state [24, 25]. Moreover, the up conformation in the ATP state may be a ready state for the substrate binding [25]. The two adjacent glycine residues are highly conserved in the C-terminus of both N-D1 linker and D1-D2 linker among the AAA+ proteins [21, 22, 27, 34]. Since glycine is the most flexible of the 20-aa residues, these two glycines could serve as a pivot point, allowing the movement of the N-domain [21, 22]. The mutation of both Gly residues to the less flexible residues may constrain the movement of the N-domain, affecting the position of the N-domain in the ATP state, ultimately resulting in the reduced substrate binding activity. Indeed, this hypothesis is supported by the result that the G221PG222P mutant is unable to bind the substrate, while the G221AG222A still retains some weak binding ability (Fig. 5B), since Pro is less flexible than Ala.

Materials and Methods

Plasmids and Mutagenesis

Plasmids encoding His6-NSF, GST-His6-α-SNAP, His6-VAMP 1-94-MBP, His6-VAMP 1-94, His6-SNAP25 1–90, His6-SNAP25 125–206 and GST-Syntaxin 2–253 have been previously described [25]. The site-directed mutants were generated with Easy Mutagenesis System (Transgen) using His6-NSF as the template. Eight mutants, P211A, D212A, F215A, E216A, K217A, M218A, G219A, I220A, were cloned into the pQE31 expression vector. Other mutants were cloned into the pQE31 expression vector. All of the mutations were confirmed by DNA sequencing.

Protein expression and purification

Mutated proteins were expressed in *E. coli* BL21 (DE3) cells at 16°C overnight after addition of 0.3 mM IPTG (isopropyl β-D-thiogalactopyranoside) at an *A*<sub>600</sub> of 0.6–0.8. Protein purification was performed as previously described [25], using the Ni-NTA superflow affinity chromatography (Qiagen) followed by the gel filtration chromatography using the Superdex-200 column (GE Healthcare). The purified proteins were analyzed by 13.5% SDS-PAGE gel containing 8 M urea and the gel was stained with Coomassie Blue. Protein quantification was performed by the Bradford method using the bovine serum albumin as the standard.

Analysis of Oligomeric State of NSF Mutants

The oligomeric analysis of NSF mutants was analyzed by gel filtration chromatography using a Superose-6 column (GE Healthcare) in the buffer containing ADP-AlFx.

SNARE disassembly assay

SNARE complex was assembled as previously described [25]. For SNARE disassembly assays, SNARE complexes were preincubated with wild-type NSF or the mutants and α-SNAP in 40 μl of a buffer (50 mM HEPES, pH 7.6, 100 mM NaCl) on ice, following the addition of 2 mM Mg<sup>2+</sup>-ATP. The reactions were performed at 37°C for the indicated times, then aliquots (10 μl) of each reaction were stopped with the addition of the SDS-PAGE loading buffer and immediately loaded onto 13.5% SDS-PAGE gels containing 8 M urea without boiling. The gels were then stained with Coomassie Blue. The SNARE proteins at 0 min and 60 min were quantified by densitometry using ImageJ.
MBP-SNARE/α-SNAP/NSF binding assay

MBP-SNARE complexes were assembled by mixing Syntaxin, SNAP25-N, SNAP25-C and MBP-VAMP in an equimolar concentration at 4°C overnight. The formed MBP-SNARE complexes were separated from unassembled individual SNAREs by gel filtration using Superdex-200 column (GE Healthcare). For the binding assay, 15 µg of MBP-SNARE complexes were incubated with excess wild-type or mutant NSF proteins (each 100 µg) and α-SNAP (30 µg) for 3 h at 4°C in the binding buffer (50 mM HEPES, pH 7.5, 100 mM NaCl, 1 mM DTT, 2 mM MgCl₂, and 10% glycerol with 2 mM ADP-AlFx). 40 µl of Amylose Magnetic Beads prewashed with the binding buffer were then added and incubated with rotation. After 4 h incubation at 4°C, the beads were washed three times (1 ml each time) with the binding buffer. The bound protein complexes were eluted with 40 µl of the elution buffer (the binding buffer containing 10 mM maltose) and then analyzed (5 µl of the eluted solution) by 8% SDS-PAGE. The reactions without α-SNAP were used to detect the background of the nonspecific binding. The gels were stained with Coomassie Blue, and the bound proteins were quantified by densitometry using ImageJ.

ATPase Activity Assay

ATPase assays measure released phosphate based on the formation of colored phosphomolybdate complexes with cationic dye malachite green [36]. Basal ATPase activity assays were carried out using 3 µg of wild-type or mutant NSF proteins in 50 µl of the ATPase assay buffer containing 20 mM HEPEs, pH 7.5, 100 mM NaCl, 2 mM MgCl₂ and 4 mM ATP. The reactions were incubated for 1 h at 25°C, and stopped by adding 20 mM EDTA. Then, 800 µl of the color reagent (0.7 mM malachite green, 8.5 mM ammonium molybdate, 0.1% NP-40, 1.75 M HCl) was added and mixed well. After 5 min incubation at room temperature (25°C), 100 µl of 34% citric acid was added and mixed well. This solution was incubated at room temperature for 15 min, and immediately the absorbance at 645 nm was measured using a spectrophotometer (Amersham). Values were corrected for non-enzymatic breakdown of ATP by running duplicate assays in the absence of NSF protein. The inorganic phosphate released was calculated based on the absorbance standard curve established using KH₂PO₄ standards. All assays were repeated at least three times, and the average activities with standard errors of measurement were presented.

To measure the SNARE/α-SNAP-stimulated ATPase activity, SNARE/α-SNAP complexes were assembled for 2 h at 4°C and then incubated with wild-type or mutant NSF proteins in the ATPase assay buffer. The following process was as above. The fold increase in ATPase activity was calculated by dividing the stimulated ATPase activity by the basal ATPase activity.

Supporting Information

Figure S1 SNARE disassembly by wild-type and mutant NSF proteins. SNARE complexes were incubated with wild-type or mutant NSF proteins, and α-SNAP in the presence of 2 mM Mg²⁺-ATP at 37°C for 0 min, 20 min and 60 min, followed by the addition of SDS-PAGE loading buffer and analyzed by SDS-PAGE. The gels were stained with Coomassie Blue.

(TIF)

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Author Contributions

Conceived and designed the experiments: CCL SS SFS. Performed the experiments: CCL. Analyzed the data: CCL SS SFS. Contributed reagents/materials/analysis tools: CCL SS SFS. Wrote the paper: CCL SS SFS.

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