Chapter 4

The 5 kDa Protein NdhP Is essential for Stable NDH-1L Assembly in *Thermosynechococcus elongatus*


Abstract

The cyanobacterial NADPH:plastoquinone oxidoreductase complex (NDH-1), that is related to Complex I of eubacteria and mitochondria, plays a pivotal role in respiration as well as in cyclic electron transfer (CET) around PSI and is involved in a unique carbon concentration mechanism (CCM). Despite many achievements in the past, the complex protein composition and the specific function of many subunits of the different NDH-1 species remain elusive. We have recently discovered in a NDH-1 preparation from *Thermosynechococcus elongatus* two novel single transmembrane peptides (NdhP, NdhQ) with molecular weights below 5 kDa. Here we show that NdhP is a unique component of the 450 kDa NDH-1L complex, that is involved in respiration and CET at high CO$_2$ concentration, and not detectable in the NDH-1MS and NDH-1MS’ complexes that play a role in carbon concentration. C-terminal fusion of NdhP with his-tagged superfolder Green Fluorescence Protein (GFP) and the subsequent analysis of the purified complex by electron microscopy and single particle averaging revealed its location in the NDH-1L specific distal unit of the NDH-1 complex, that is formed by the subunits NdhD1 and NdhF1. Moreover, NdhP is essential for NDH-1L formation, as this type of NDH-1 was not detectable in a ΔndhP::Km mutant.

**Keywords:** NDH-1, *T. elongatus*, NdhP, electron microscopy, negative stain particle averaging.

**Author contributions:** Conceived and designed the experiments: HW and MMN; molecular biology and cloning: HW; cultivation and harvest of *T. elongatus*, BN-PAGE/SDS-PAGE and mass spectrometry: HW and TW; analysis of mass spectrometry: HW and MMN; Electron Microscopy: LEF; Electron microscopy analysis: LEF and EJB; Manuscript writing: HW, MMN, LEF and EJB; Figures and tables: HW and LEF.
1. Introduction

The cyanobacterial and chloroplast type I NADPH dehydrogenase (NDH-1) complex is structurally and functionally related to the energy-converting NAD(P)H:Quinone oxidoreductase (Complex I) – one key-enzyme of the energy metabolism in eubacteria (like Escherichia coli) and the respiratory chain of mitochondria (Friedrich and Scheide, 2000; Brandt, 2006; Hirst, 2009). This large membrane protein complex is composed of up to ~45 subunits in mammals with a total molecular weight approaching 1 MDa (Carroll et al., 2006). Remarkable improvements in the X-ray structural analysis of eubacterial Complex I revealed detailed insights into the electron transfer dependent proton-pumping mechanism on the molecular level (Sazanov and Hinchliffe, 2006; Baradaran et al., 2013).

Cyanobacterial NDH-1 and the closely related chloroplast NDH complex are located in the thylakoid membrane and play a pivotal role in respiration (chlororespiration in chloroplasts) as well as in cyclic electron transfer (CET) around PSI (Shikanai, 2014; Kouřil et al., 2014). They share the so called oxygenic photosynthesis-specific domain of unknown function, which is composed of NdhL, -M, -N and -O (Birungi et al., 2010) and the basic subunits NdhA-NdhK, which are homologous to subunits of the eubacterial complex. Chloroplast NDH has a more intricate structure and includes several additional proteins compared to cyanobacteria (Ifuku et al., 2011). Most strikingly, the cyanobacterial and chloroplast type NDH-1 complex seem to lack homologues to NuoE/F/G of the eubacterial complex that are responsible for NADH oxidation. This led to the recent idea that electron transfer via ferredoxin might be a possible pathway to chloroplast NDH and cyanobacterial NDH-1 (Yamamoto et al., 2011; Hu et al., 2013).

Based on reverse genetics and proteomic studies, significant progress has been made in resolving the subunit composition and function of the cyanobacterial complex (Ogawa and Mi, 2007; Battchikova et al., 2011a). At least four different NDH-1 subtypes (NDH-1L, NDH-1L’, NDH-1MS, NDH-1MS’) have been identified or postulated for cyanobacteria so far. NDH-1L and NDH-1L’ are involved in respiration whereas NDH1-MS and NDH-1MS’ link the Complex I specific and electron flow dependent proton-pumping activity to unique carbon concentration mechanisms (CCM) in an unknown process. All NDH-1 variations have been shown to play a role in CET, in particular under...
stress conditions (Bernat et al., 2011). They all share a common NDH-1M core unit but they differ in the presence of subunits that are unique for each complex type like NdhF1/D1 (NDH-1L), NdhF1/D2 (NDH-1L’), NdhF3/D3/CupA/CupS (NDH-1MS) and NdhF4/D4/CupB (NDH-1MS’). Two more isoforms of the NdhD subunit (NdhD5, NdhD6) were identified on genome level but they have not been assigned to specific NDH-1 complexes yet. Although reverse genetic and proteomic studies have already revealed a detailed picture of cyanobacterial NDH-1 complexes, it is still incomplete in many aspects. We have recently identified two novel peptides – NdhP and NdhQ – in purified NDH-1 complexes from T. elongatus (Nowaczyk et al., 2011). NdhP plays a role in NDH-1 mediated electron flow in Synechocystis sp. PCC 6803 (Schwarz et al., 2013) and shares a weak similarity with NDF6, which is important for NDH-1 activity in Arabidopsis thaliana (Ishikawa et al., 2008). Here we show by isolation and structural characterization of individual NDH-1 complexes from T. elongatus that the 5 kDa NdhP peptide is a unique component of the NDH-1L subtype and essential for its assembly.

2. Materials and Methods

2.1 Construction of T. elongatus mutants
To delete the ndhP gene (genomic region: 1189596–1189465) the coding region was replaced by a kanamycin resistance cassette. Genomic DNA from T. elongatus was used as a template for the amplification of the upstream region of ndhP by PCR using the specific oligonucleotide primers ndhP_up_for and ndhP_up_ KO_rev (Table S1 (Wulfhorst et al., 2014)). The ndhP downstream region was amplified using the primer pair ndhP_down_for and ndhP_down_rev. The DNA fragments were restricted with SacI and XbaI (upstream) and PstI and Xhol (downstream) and ligated into the vector pBluescript SK (+) (Stratagene). A kanamycin resistance cassette was introduced via the XbaI and PstI restriction sites and the resulting plasmid pNdhP_KO was used for transformation of T. elongatus according to (Iwai et al., 2004). Complete segregation of the mutant allele was confirmed by PCR with the primers SegCheck NdhP_for and SegCheck NdhP_rev.
The NdhP-sfGFP-His₆ mutant was generated based on a modified pNdhP_KO plasmid. NdhP and the corresponding upstream region were replaced by a PCR product (primers: ndhP_up_for and ndhP_up_sfGFP_rev) that introduces an EcoRV site upstream of the ndhP stop codon. The coding region for superfolder GFP (Pédelacq et al., 2006) with a c-terminal His-Tag (sfGFP-His₆) was cloned as a synthetic DNA construct (Life Technologies) and introduced into the final plasmid pNdhP-sfGFP-His₆ via the EcoRV restriction site. After transformation of T. elongatus with pNdhP-sfGFP-His₆, segregation was checked by PCR (primers: SegCheck NdhP_for and SegCheck NdhP_rev).

To generate the NdhL-TS (TwinStrep-tag) mutant, ndhL (tsr0706) and the corresponding upstream region were amplified by PCR (primers: ndhL_up_strep_for and OneSTrEP_rev1) and the product was used as template for a second PCR (primers: ndhL_up_strep_for and OneSTrEP_XbaI_rev2). The fragment was cloned into pBluescript SK (+) via SacI and XbaI restriction sites. The ndhL downstream region was amplified by PCR (primers: ndhL_down_for and ndhL_down_rev) and cloned via EcoRI and Apal. Finally, the kanamycin resistance cassette was introduced via XbaI and EcoRI and the resulting plasmid was used for transformation of T. elongatus. Segregation was checked via PCR (primers: SegCheck_NdhL-strep_for and Seg-Check_NdhL-strep_rev).

2.2 Culture conditions – T. elongatus
T. elongatus wild type strain and mutants (NdhP-sfGFP-His₆, ΔndhP::Km and NdhL-TS) were grown in BG-11 liquid medium (Rippka et al., 1979) at 45°C, bubbled with 5% CO₂ under illumination of increasing intensity of 50–200 mMol photons (dependent on cell density). The medium used for the cultivation of the mutants was supplemented with 80 mg/ml kanamycin.

2.3 Cell harvest and solubilization of thylakoid membranes
The cells were harvested and thylakoids were prepared as described earlier (Kuhl et al., 2000). The buffer composition for n-dodecyl-beta-D-maltoside (DDM, Glycon Biochemicals) solubilization of thylakoid membranes depended on the following purification method. For subsequent Ni²⁺ affinity chromatography the thylakoids were suspended in MES-Buffer (20 mM MES pH 6.5; 25 mM MgCl₂; 1% (w/v) DDM; 1 mM 4-(2-aminoethyl) benzenesulfonyl 133
fluoride) and for streptavidin affinity chromatography m-buffer (100 mM Tris pH 8.0; 25 mM MgCl₂; 1% (w/v) DDM; 1 mM 4-(2-aminoethyl) benzenesulfonyl fluoride) was used. The thylakoid membranes were suspended in the respective buffer at a final chlorophyll concentration of 1 mg/ml and incubated under gentle agitation at 20°C for 60 min. To adjust the final DDM concentration to 0.5% an equal volume of dilution buffer (same composition but without DDM) was added. Insoluble material was removed by centrifugation at 45,000 g.

### 2.4 Purification of NDH-1 complexes

Solubilized thylakoid membranes were filtered through a 0.45 mm membrane filter and samples containing His-tagged NDH-1 complexes were applied to a 5 ml FF crude IMAC column (GE Healthcare) at a flow rate of 1 ml/min after equilibration with IMAC-equilibration buffer (20 mM MES pH 6.5; 0.5 M mannitol; 150 mM NaCl; 0.03% (w/v) DDM). The column was washed with 5 column volumes (CV) IMAC-equilibration buffer containing 10 mM imidazole and His-tagged complexes were eluted with a 10–500 mM imidazole gradient (4 CV) in IMAC-equilibration buffer.

Samples containing TwinStrep-tagged NDH-1 complexes were applied to a 1 ml StrepTactin high capacity column (IBA Biotechnologies) at a flow rate of 1 ml/min after equilibration with Strep-equilibration buffer (100 mM Tris pH 8.0; 0.5 M mannitol; 150 mM NaCl; 0.03% (w/v) DDM). Unbound material was removed by washing with 7 CV Strep-equilibration buffer. Strep-tagged protein complexes were eluted with buffer containing 2.5 mM desthiobiotin. The eluted proteins were concentrated using a spin concentrator with 100 kDa cut off (Millipore) and stored at -80°C.

### 2.5 Electrophoresis

Blue native (BN)-PAGE was performed according to Battchikova et al. (2011b). Purified proteins (5–10 mg) were mixed with 1/10 volume of sample buffer, loaded on a blue native gradient gel (5–12.5% acrylamide) and electrophoresis was carried out at 4°C at increasing voltage (50–200 V). For electrophoresis in the second dimension, the respective BN gel-lane was cut out, incubated in buffer containing 5% β-mercaptoethanol and 6 M urea and loaded on a 1 mm-
thick 14% SDS-PAGE gel with 6 M urea. After electrophoresis (4°C, 13 mA) the proteins were visualized by silver staining according to Blum et al. (1987).

2.6 Identification of proteins by 1D-nLC-ESI-MS/MS and MALDI-ToF MS
Protein spots from BN-gels and silver-stained 2D-gels were excised and digested with trypsin as described earlier (Nowaczyk et al., 2011). The digests were desalted by ZipTips (Millipore), resuspended in 0.1% formic acid in water and analyzed by 1D-nLC-ESI-MS/MS as described in Nowaczyk et al. (2011). Intact NDH-1 subunits were examined by MALDI-ToF MS according to Nowaczyk et al. (2012).

2.7 Transmission electron microscopy and single particle analysis
Purified samples were prepared for negative staining with 2% uranyl acetate on glow-discharged carbon-coated quantifoil grids. Electron microscopy was performed on a Tecnai G2 20 Twin electron microscope (FEI, Eindhoven, The Netherlands) equipped with a LaB6 cathode, operated at 200 kV. Images were recorded at 330 nm defocus with an UltraScan 4000 UHS CCD camera (Gatan, Pleasanton, CA, USA) at 100,000-fold nominal magnification with a pixel size of 0.224 nm at the specimen level. GRACE software (Oostergetel et al., 1998) was used for semi-automated data acquisition. Particle picking was done manually in Eman2 (Tang et al., 2007). Output particles were generated in image format and imported into Groningen Image Processing software (GRIP), which was used for further analysis.

A total of 9000 wild type and 15000 NdhP-GFP-labeled NDH-1 complexes were collected and processed using standard procedures (Boekema et al., 1999). Images were pretreated using a low frequency cut-off filter based on the maximum size of the particle and a high frequency cut-off filter based on the maximum resolution available in negative stain (10 Å). The presented images were optimized by application of conditional summing with the correlation coefficient of the final alignment step as a quality parameter to select the most homogeneous images in each class (correlation decay was between 0.1 and 0.2, depending on the number of particles in the class). Resolution of the final
results was determined with a Fourier-ring correlation with a 3 σ threshold criterion.

Comparison of the X-ray structure of NDH-1 from \textit{T. thermophilus} (Baradaran \textit{et al.}, 2013) (PDB code: 4HEA) to the density map from NDH-1L that was retrieved in this study was done by using UCSF Chimera (Pettersen \textit{et al.}, 2004) and Adobe Photoshop. All proteins were fitted separately.

\textbf{2.8 Bioinformatic tools}

Sequence alignment was performed using the Clustal algorithm in the program JalView (Thompson \textit{et al.}, 1997; Waterhouse \textit{et al.}, 2009), visualized with the ClustalX residue color code. Modelling of cyanobacterial subunits was performed using the SwissModel server (http://swissmodel.expasy.org, (Guex \textit{et al.}, 2009)) with the structure of NDH-1 from \textit{T. thermophilus} (Baradaran \textit{et al.}, 2013) (PDB: 4HEA) as template.

\textbf{3. Results and Discussion}

\textbf{3.1 NdhP is a unique component of the NDH-1L complex}

We have shown previously that the single-transmembrane protein NdhP co-purifies with NDH-1L complexes, which were isolated via the histidine-rich region of NdhF1 (Nowaczyk \textit{et al.}, 2011) but it remained unclear whether NdhP is also a component of the NDH-1MS or NDH-1MS’ complex and the corresponding subcomplexes NDH-1S or NDH-1S’. To answer this question an NdhL-TwinStrep-tag (NdhL-TS) mutant was constructed that enables purification of the other NDH-1 variants. The NdhL subunit is part of the basic NDH-1 unit called NDH-1M and it was shown by Zhang \textit{et al.} that purification of NDH-1 complexes via NdhL-His resulted in a mixture of NDH-1M, NDH-1L, NDH-1MS and NDH-1S complexes (Zhang \textit{et al.}, 2005). We used the Strep-tag/Streptactin affinity chromatography system due to its superior purity compared to His-tag affinity chromatography (Lichty \textit{et al.}, 2005).

The TwinStrep-tag was fused to the C-terminus of NdhL and the complete segregation of the NdhL-TS mutant was confirmed by PCR analysis (Figure 4-1). Solubilized membranes of 5 L culture were subjected to StrepTactin affinity
chromatography and the concentrated elution fractions were analyzed by BN- and 2D-gelelectrophoresis (Figure 4-2). Four different NDH-1 complexes were separated by BN-PAGE and the identity of NDH-1L, NDH-1M and NDH-1S was confirmed by high resolution LC-MS/MS (Tables S2, S3 and S4 (Wulfhorst et al., 2014)) and MALDI-ToF MS analysis (Figure S1 (Wulfhorst et al., 2014)). Moreover, all specific subunits of NDH-1S’ (NdhD4, NdhF4, CupB) were identified by MS analysis (Table S5 (Wulfhorst et al., 2014)) which confirms – for the first time to our knowledge – its existence on protein level.

The NdhP subunit was solely detected in the NDH-1L complex by identification of a specific peptide (Table S6, Figure S2 (Wulfhorst et al., 2014)) that was missing in NDH-1M, NDH-1S and NDH-1S’ samples. Although no corresponding bands of the intact NDH-1MS or NDH-1MS’ complex are visible in the BN-PAGE analysis, NdhP can be assigned to the NDH-1L complex – most likely inside the NdhD1/F1 unit - as it is not present in any of the corresponding subcomplexes.

### 3.2 Electron microscopy localizes NdhP in the distal NdhD1/F1 unit of NDH-1L

It was previously shown that fusion of several NDH-1 subunits with YFP and the subsequent analysis by electron microscopy and single particle averaging revealed the position of the tagged subunits inside the NDH-1 complex of *Synechocystis* sp. (Birungi et al., 2010). To investigate the distinct localization of NdhP within the NDH-1L complex, we fused the 27 kDa superfolder GFP (sfGFP) protein (Pédelacq et al., 2006) with an additional His-tag to the NdhP C-terminus. A C-terminal fusion was applied to ensure cytoplasmic localization of the sfGFP tag, as the N-terminus of the single transmembrane helix protein NdhP was predicted to be oriented towards the lumen (Nowaczyk et al., 2011). This construct should avoid misfolding as well as impaired membrane insertion of NdhP caused by the sfGFP fusion. The thermophilic target strain *T. elongatus* was cultured at elevated temperature (45°–50°C) for optimal growth. Superfolder GFP was used for the tagging approach, as it shows enhanced thermostability compared to other GFP variants (Cava et al., 2008).
Figure 4-1: Construction of T. elongatus NdhL-TS and segregation check. Schematic representation of the ndhL mutant locus (left). The Twin-StrepTag was fused to the NdhL C-terminus and a kanamycin resistance marker was used to force segregation of the mutant allele. Complete segregation was confirmed by PCR analysis (right).

Figure 4-2: Isolation of NDH-1 complexes via NdhL-TwinStrep-Tag and 2D-PAGE analysis of purified samples. A) 2D-PAGE analysis isolated NDH-1 complexes. The identity of NDH-1L and NDH-1M as well as NdhP was confirmed by MS/MS analysis (see Table S2, S3 and S6 (Wulfhorst et al., 2014)). B) Identification of NDH-1S and NDH-1S' by 2D-PAGE analysis and mass spectrometry (see Table S4 and S5 (Wulfhorst et al., 2014)). C) Tagged NDH-1 complexes were purified by StrepTactin affinity chromatography.
The complete segregation of the *T. elongatus* NdhP-sfGFP-His mutant was confirmed by PCR analysis (Figure 4-3). His-sfGFP-tagged NDH-1 complexes were isolated via Ni-affinity chromatography (Figure 4-4A) and analyzed by BN-PAGE (Figure 4-4B). As expected, only a single protein complex with a molecular weight of approximately 450 kDa was isolated and mass spectrometry analysis revealed the presence of the NDH-1L specific subunits NdhD1 and NdhF1 (Table S7 (Wulfhorst et al., 2014)). The NdhP-sfGFP-fusion was verified by mass spectrometry, as a specific peptide of the linker region that connects NdhP and sfGFP, was identified in the sample (Table S8, Figure S3 (Wulfhorst et al., 2014)).

**Figure 4-3: Construction of *T. elongatus* NdhP-sfGFP-His and segregation check.** Schematic representation of the ndhP mutant locus (left). The sfGFP-His6-Tag was fused to the NdhP c-terminus and a kanamycin resistance marker was used to force segregation of the mutant allele. Complete segregation was confirmed by PCR analysis (right).

**Figure 4-4: Isolation of NDH-1L via NdhP-sfGFP-His by Ni-affinity chromatography.** A) His-tagged NDH-1L complexes were eluted from a Ni-NTA column with a linear gradient of 10–500 mM imidazole. The absorption of sfGFP at 485 nm was used to follow the purification. B) BN-PAGE of eluted proteins. The band with an apparent molecular weight of approximately 450 kDa was identified as NDH-1L by MS/MS analysis (see Table S7 and S8 (Wulfhorst et al., 2014)).
Affinity purified NDH-1 complexes from both *T. elongatus* NdhP-sfGFP-His and NdhL-TS mutant were studied by electron microscopy. A total of 15000 NdhP-sfGFP-labeled and 9000 NdhL-TS-labeled NDH-1L complexes were collected (see Materials and Methods) and the selected single particle projections were analyzed by single particle averaging and assigned to homogenous classes, followed by subsequent averaging of class members into 2D maps. From NDH-1L of *T. elongatus* NdhP-sfGFP-His two classes of differently oriented complexes were generated, resulting in a left- and right-handed side view (Figures 4-5A and B). Compared to the projection maps of NDH-1L of *T. elongatus* NdhL-TS (Figures 4-5C and D) an extra density is visible in both side views of NDH-1L from NdhP-sfGFP-His (red arrows), which has to be the sfGFP-His-tag. In conclusion, the previously predicted location of NdhP (Nowaczyk *et al.*, 2011) is clearly confirmed by the results of the structural analysis. Based on the asymmetric shape of NDH-1L, one can assign the c-terminally fused sfGFP-tag to the cytoplasmic side of the membrane.

Moreover, the size of the tag is optically different between side view A and B. This is best explained by the tag pointing towards the viewer into the stain-layer in side view A and away from the viewer out of the stain-layer in side view B (Figure 4-5 insets). This indicates that the tag is not exactly in the middle. By fitting the 2D projection map with the homologous part of the X-ray structure of NDH-1 from *T. thermophilus* (Baradaran *et al.*, 2013) (Figure 4-6A) the position of the sfGFP-His-tag could be specified to be “above” the subunits NdhB (Nqo13, red) and NdhD1 (Nqo14, yellow), possibly with its C-terminus pointing more into the direction of NdhD1. Interestingly, the structures of Nqo7 (NdhC, light blue), Nqo8 (NdhA, orange) and the hydrophilic domain (Nqo4/NdhH (dark green), Nqo9/Ndhl (cyan), Nqo5/NdhJ (light purple), Nqo6/NdhK (light green) had to be rotated to make them fit to the 2D projection map. This changes the direction of these subunits from pointing to the right to pointing away from the viewer. A more detailed structural comparison of the Nqo13 and Nqo14 subunits from eubacterial NDH-1 (Baradaran *et al.*, 2013) with models of NdhB and NdhD1 from *T. elongatus* (Figures 4-6B, C and S4 (Wulfhorst *et al.*, 2014)) revealed that the general structure (position of transmembrane helices) seems to be consistent. One striking difference between both complexes might be an additional helix at the cytoplasmic side of NdhB. A tentative localization of NdhP in this part of the complex would be in good agreement with our data.
Figure 4-5: Projection maps of NdhP-sfGFP-His-tagged (A and B) and NdhL-TS-tagged NDH-1L complexes (C and D). A) A conditional sum of 9158 images (correlation decay: 0.2) at 16 Å resolution. B) Average of 1326 images (corr. d.: 0.15) at 25 Å resolution. C) Sum of 4409 images (corr. d.: 0.15) at 26 Å. D) Sum of 1758 images (corr. d. 0.2) to a 27 Å resolution map. The position of the sfGFP-tag is indicated by red arrows. The scale bar represents 10 nm. The insets represent schematic side views of A and B to clarify the tag-position, where the black arrows indicate the viewing direction (towards the carbon) and the blue brackets show the perceived size of the tag due to its asymmetric placing.

3.3 Isolation of NDH-1L via NdhF1 is impaired in a ΔndhP::Km mutant

To investigate the role of NdhP in the NDH-1L complex, a comparative purification from *T. elongatus* wild type and a ΔndhP::Km mutant was conducted by Ni-affinity chromatography via the histidine rich region of the NdhF1 subunit (Zhang *et al.*, 2005). The mutant was generated by replacement of *ndhP* with a kanamycin resistance cassette and complete segregation was confirmed by PCR analysis (Figure 4-7). Equivalent amounts (10 mg Chl) of thylakoid membranes from wild type and mutant were solubilized and applied to Ni-affinity chromatography. BN-PAGE analysis (Figure 4-8A) of purified protein complexes revealed the presence of a putative NDH-1L complex (red asterisk) in the wild type sample that is missing in the ΔndhP::Km mutant. Both samples contain equal amounts of co-eluted trimeric PSI complexes that are typical contaminants for this type of purification (Arteni *et al.*, 2006). The characteristic protein spot pattern that is visible in the SDS-PAGE dimension (Figure 4-8B, red asterisk) confirms the presence of NDH-1L in the wild type sample and its absence in the ΔndhP::Km mutant.

In a recent study Schwarz *et al.* analyzed a ΔndhP::Km mutant of *Synechocystis* sp. (Schwarz *et al.*, 2013). It showed reduced NDH-1 mediated cyclic electron transport (CET) around PSI under high CO₂ (HC) conditions, whereas under low CO₂ (LC) no significant change was observed. Interestingly, NDH-1L is the dominant complex under HC conditions, whereas NDH-1MS is strongly induced under LC conditions. Since the latter complex also contributes to CET, the authors speculate that NdhP is a NDH-1L specific subunit that mediates efficient CET under HC conditions. Here we can clearly show that NdhP is a unique and essential component of the NDH-1L complex. NDH-1L is not detectable anymore in the ΔndhP::Km mutant and - as a direct consequence - this mutant shows the same phenotype (Schwarz *et al.*, 2013) as ΔndhD1 and ΔndhF1, which also lack the NDH-1L complex (Bernat *et al.*, 2011; Battchikova *et al.*, 2011b).
Figure 4-7: Construction of *T. elongatus* ΔndhP and segregation check. Schematic representation of the ΔndhP mutant locus (left). A kanamycin resistance marker was used to replace the entire ndhP gene. Complete segregation was confirmed by PCR analysis (right).

Figure 4-8: Comparative purification of NDH-1 complexes via NdhF1 from wild type and ΔndhP::Km mutant cells by Ni-affinity chromatography. A) BN-PAGE analysis of purified complexes. The 450 kDa complex in the wild type (WT) lane is indicated by a red asterisk. B) 2D-PAGE analysis. The band pattern of the respective complex (red asterisk) is characteristic for NDH-1L.

In conclusion, the present study corroborated the important role of NdhP within the cyanobacterial NDH-1 complex. NdhP is located in the distal unit of NDH-1 that is formed by the NdhB and NdhD1 subunits. It was allocated specifically to the NDH-1L complex, which is involved in respiration and CET under high CO₂ concentration (Figure 4-9). And obviously NdhP is essential for the stable assembly of this multisubunit membrane protein complex, as it is not detectable anymore in the ΔndhP::Km mutant. A similar observation was published very recently for the NDH-1L complex of *Synechocystis sp.* PCC 6803
(Zhang et al., 2014). Although, they are challenging to identify and sometimes overlooked, single transmembrane domain proteins seem to play a central role in the organization of membrane protein complexes in general (Zickermann et al., 2010) and the 5 kDa NdhP single-transmembrane NDH-1 subunit is one striking example for this often undervalued class of proteins.

![Diagram showing NDH-1 complexes](image)

**Figure 4-9:** Model of NDH-1 complexes that have been verified on protein level. The basic complex NDH-1M is combined with specific domains to assemble the functional complexes NDH-1L, NDH-1MS', and NDH-1MS.

### 5. Acknowledgments

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6. References


