

## STRUCTURE NOTE

# Structure of the C-terminal domain of *Saccharomyces cerevisiae* Nup133, a component of the nuclear pore complex

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## INTRODUCTION

Nuclear pore complexes (NPCs) are situated at the interface of the inner and outer nuclear membranes, where they are responsible for transport of biomolecules between nucleus and cytoplasm of eukaryotic cells.<sup>1–3</sup> *Saccharomyces cerevisiae* NPCs are ~50 MDa assemblies made up of at least 456 polypeptides of ~30 different sequences, called nucleoporins or Nups.<sup>4,5</sup> Several of these Nups, sharing similar structural motifs, assemble into stable subcomplexes that are arranged with eightfold symmetry to form a co-axial ring structure containing a membrane ring, two outer (nuclear and cytoplasmic) rings and two inner rings. One of these subcomplexes in the yeast NPC is the Nup84 complex that is situated within the nuclear and cytoplasmic peripheral rings.<sup>5</sup> The Nup84 complex is a heptameric assembly forming a Y-shaped structure composed of Nup84, Nup85, Nup120,

Nup133, Nup145C, Sec13, and Seh1.<sup>6</sup> The homologous metazoan complex is a nonameric assembly termed the Nup107–160 complex, carrying homologues of all seven yeast Nups plus Nup37 and Nup43 as additional components.<sup>7</sup> Nup133 is a ~1150 residue protein containing an N-terminal seven blade  $\beta$ -propeller domain followed by an ~80 kDa helical domain.<sup>8–10</sup> The C-terminal domain

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of human Nup133, in particular, residues 934–1156 [HsNup133(934–1156)], is essential for its binding to Nup107, together forming the stem of the Y-shaped complex. Crystal structures of the HsNup133(934–1156) have been determined in complex with Nup107 (residues 658–925). In this complex, 11 HsNup133(934–1156) helices are arranged as two helical blocks.<sup>11,12</sup>

We have determined the structure of the *S. cerevisiae* Nup133 (ScNup133) C-terminal helical domain at 1.9 Å resolution, corresponding to residues 944–1157 [ScNup133(944–1157)]. Our structure reveals a compact helical arrangement, differing in several respects from the structure of the homologous HsNup133(934–1156) determined in complex with Nup107. We also report small-angle X-ray scattering (SAXS) of ScNup133(944–1157) in solution that is consistent with our compact crystallographically observed structure.

## EXPERIMENTAL

### Cloning, expression, and purification of ScNup133(944–1157)

ScNup133 comprising residues 881–1157 was cloned from the genomic DNA of the yeast strain Sc2601D-5 (American Type Culture Collection, USA). The desired construct was PCR amplified using the forward, AAG TACGGTCATGTAGCATGGA and, reverse, CGTATTCTA CAGTGTGGTTTCATAG, primers and subsequently TOPO<sup>®</sup> (Invitrogen, USA) cloned into pSGX3, a derivative of pET26 b(+), giving rise to proteins with a noncleavable C-terminal hexa histidine tag. Plasmids were transformed into BL21 (DE3)-Condon + RIL (Invitrogen) cells for overexpression. Expression of selenomethionine-labeled proteins was carried out in 3 L of HY media at 22°C with 50 µg/mL of kanamycin and 35 µg/mL of chloramphenicol. Protein expression was induced at O.D.<sub>600</sub> ~1.0 by addition of 0.4 mM IPTG. Cells were harvested after 21 h by centrifugation at 4°C.

For purification, the *E. coli* cell pellet was resuspended in 30 mL of cold buffer containing 20 mM Tris–HCl pH 8.0, 500 mM NaCl, 25 mM imidazole, and 0.1% Tween20 and cells were lysed by sonication. Debris was removed by centrifugation at 4°C. The decanted supernatant was applied to a 5-mL HisTrapHP column (GE Health Care, USA) charged with nickel and pre-equilibrated with 20 mM Tris–HCl pH 8.0, 500 mM NaCl, 10% v/v glycerol, and 25 mM imidazole. The sample was washed with five column volumes (CVs) of 20 mM Tris–HCl pH 8.0, 500 mM NaCl, 10% (v/v) glycerol, and 40 mM imidazole and subsequently eluted with 2 CV of same buffer with an imidazole concentration of 250 mM. Eluted protein was passed over a 120-mL Superdex 200 column equilibrated with 10 mM HEPES pH 7.5, 150 mM NaCl, 10% (v/v) glycerol, and 5 mM DTT (protein storage buffer). SDS–PAGE analysis showed greater than 95% purity, and

protein fractions corresponding to the symmetric portion of the size exclusion chromatography profile were pooled for concentration using AMICON spin filters. Concentrated protein aliquots were frozen in liquid nitrogen and stored at –80°C.

### Crystallization, data collection, and structure determination

ScNup133(944–1157), at a concentration of 17.6 mg/mL, was subjected to crystallization screening with the Classics, Classics II, and PEGs kits (Qiagen, USA) using a Phoenix Liquid Handling Systems (Art Robins, USA) with equal volumes (0.3 µL) and reservoir solution in 96-well sitting drop format at 21°C. Diffracting quality crystals were obtained with 20% PEG3350 in the presence of 200 mM potassium thiocyanate (PEGs suite; condition number 62) and were cryoprotected with the addition of ~30% (v/v) glycerol before being flash frozen by immersion in liquid nitrogen. Diffraction data were recorded using the LRL-CAT 31-ID beamline at the advanced photon source (APS) and processed with MOSFLM<sup>13</sup> and SCALA (Collaborative Computing Project Number 4, 1994).<sup>14</sup> About 800 reflections were randomly selected for cross validation representing an *R*-free dataset. The structure of ScNup133(944–1157) was determined by the single-wavelength anomalous dispersion method, using data collected at a wavelength of 0.97929 Å, corresponding to the “Se peak” absorption edge. A single Se atom was located with SHELXD,<sup>15</sup> and phases were calculated using SHELXE as implemented in HKL2MAP.<sup>16</sup> Following numerous rounds of automated and manual model building with ARP/wARP<sup>17</sup> and COOT,<sup>18</sup> respectively, the atomic coordinates were refined using REFMAC5<sup>19</sup> as implemented in CCP4. The polypeptide backbone of ScNup133 could be traced continuously from residue Lys946 to Tyr1157 in the electron density maps. The final refined atomic model of ScNup133(944–1157) has excellent stereochemistry (Table I). Structural analyses were carried out using COOT and CCP4, and illustrations were prepared using PyMol (<http://pymol.sourceforge.net>).

### Small-angle X-ray scattering

Small-angle X-ray scattering (SAXS) measurements of ScNup133(944–1157) were carried out on the SIBYLS Beamline 12.3.1 at the advanced light source (ALS), Lawrence Berkeley National Laboratory. Samples were prepared on a 96-well plate and kept at 10°C until SAXS measurement. An automatic sample delivery system equipped with a Hamilton pipetting robot was used.<sup>22</sup> SAXS data were collected on a MAR165 area detector (Rayonix), placed at 1.5 m from the SAXS sample cell. Exposures of 2 sec/10 sec/2 sec were made in series for each protein sample maintained at 10°C. Each of the diffraction images was scaled using the transmitted beam intensity, azimuthally

**Table I**

Crystallographic Statistics

Data collection	ScNup133(944–1157)
PDB code	3KFO
Space group	$P2_12_12_1$
Unit-cell dimensions $a$ , $b$ , and $c$ (Å)	$a = 47.4$ , $b = 52.7$ , $c = 76.7$
Matthew's coefficient (Å <sup>3</sup> /Da)	1.89
Solvent content (%)	34.45
Resolution (Å)	31.00–1.90 (2.00–1.90)*
Number of unique reflections	15,683
Completeness (%)	99.7 (100.0)
$R_{\text{sym}}$ (%)	12.1 (41.4)
Multiplicity	7.7 (7.8)
$\langle I/\sigma(I) \rangle$	10.6 (4.6)
Refinement	
$R_{\text{cryst}}$ (%)	19.1
$R_{\text{free}}$ (%)	24.8
RMS deviations from ideal values	
Bond length (Å)	0.019
Bond angles (°)	1.63
Average B-factors (Å <sup>2</sup> )	
Protein	23.4
Water molecules	31.5
Cosolvent (glycerol)	45.7
Ramachandran Plot <sup>20</sup>	
MolProbity <sup>21</sup> residues in favored region (%)	99.5
Allowed region (%)	100.0

\*Values in parenthesis correspond to the highest-resolution shell.

integrated, and averaged to obtain fully processed data in the form of intensity versus  $q$  [ $q = 4\pi \sin(\theta)/\lambda$ ,  $\theta$  = one-half of the scattering angle;  $\lambda$  = X-ray wavelength]. The buffer profile was obtained in the same manner and subtracted from a protein profile. SAXS profiles of ScNup133(944–1157) were recorded at protein concentrations of 0.5, 1.0, 2.0, 5.0, and 17.6 mg/mL in protein storage buffer. Moreover, mild concentration dependence was removed by extrapolating to zero concentration. We merged the average of the low-scattering angle parts ( $q < 0.15 \text{ Å}^{-1}$ ) of the low concentration profiles (0.5–2.0 mg/mL) and the average of the high-scattering angle parts ( $q > 0.12 \text{ Å}^{-1}$ ) of the high concentration (5.0–17.6 mg/mL) profiles to obtain the final, merged experimental SAXS profile. SAXS profiles were calculated with IMP-FoXS<sup>23,24</sup> (<http://salilab.org/foxs/>) and CRY SOL<sup>25</sup> and compared to the merged experimental SAXS profile. A complete structural model of ScNup133(944–1157), including two N-terminal residues (Leu-Arg), C-terminal hexa-histidine tag (Gly-His-His-His-His-His), and 16 missing side chains, was generated using its crystal structure with the automatic PSF-based model building option of VMD<sup>26</sup> and the automodel function of MODELLER<sup>27</sup> (these missing atoms were not included in the final X-ray structure, because they appear disordered in the electron density map). Inclusion of the missing atoms improved the fit of the calculated SAXS profiles to our experimental observations ( $\chi^2$  value improved from 2.95 to 1.25), particularly at higher scattering angles ( $q > 0.2 \text{ Å}^{-1}$ ). The shape of ScNup133(944–1157) was calculated from the merged experimental SAXS

profile by running DAMMIF<sup>28</sup> and GASBOR<sup>29</sup> 20 times individually, followed by superposition and averaging with DAMAVER.<sup>30</sup>

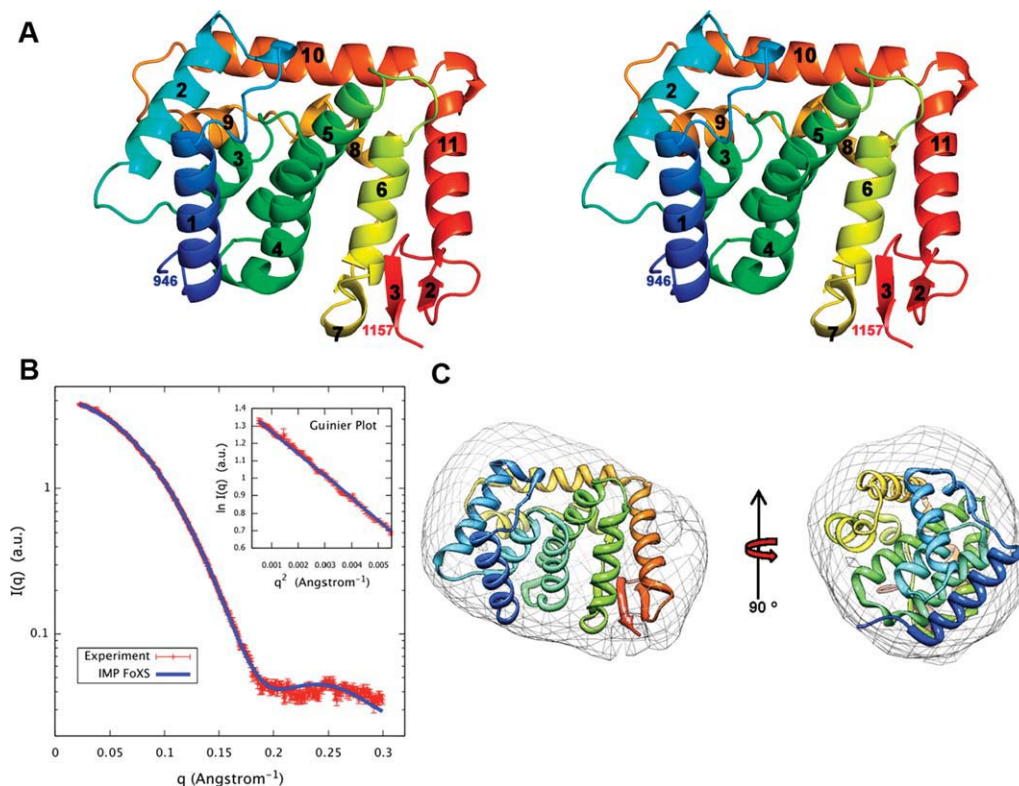
## RESULTS AND DISCUSSION

### Structure of ScNup133(944–1157)

The expected molecular weight of ScNup133 (residues 881–1157) is 33,025 Da. The measured molecular weight, using mass spectrometry, of the final purified selenomethionine form of ScNup133 was 25,871 Da (mass error < 0.007%). The difference in mass of –7154 Da was attributed to unintended N-terminal proteolysis during expression and/or purification (the presence of an intact C-terminus with hexa-histidine affinity tag was confirmed by binding the purified ScNup133 to Ni-NTA resin). N-terminal sequencing documented that the truncation occurred at the Thr943-Leu944 peptide bond. The resulting target protein, consisting of residues 944–1157 [ScNup133(944–1157)], yielded crystals diffracting to 1.9 Å resolution (orthorhombic  $P2_12_12_1$  space group, one molecule per asymmetric unit, Table I). Electron density maps phased with a single selenium atom allowed continuous tracing of the ScNup133 polypeptide backbone from Lys946 to Tyr1157.

The overall fold of ScNup133(944–1157) is composed of 11  $\alpha$ -helices with a short two-stranded  $\beta$ -sheet, formed by  $\beta 2$ – $\beta 3$ , at the C-terminus [Fig. 1(A)]. Helices  $\alpha 1$ – $\alpha 11$  form a compact structure. A four helical bundle (formed by the  $\alpha 3$ ,  $\alpha 4$ ,  $\alpha 5$ , and  $\alpha 6$  helices) at the center of the structure is surrounded by the  $\alpha 1$ ,  $\alpha 2$ ,  $\alpha 9$ ,  $\alpha 10$ , and  $\alpha 11$  helices. A search for structurally similar proteins using DALI<sup>34</sup> did not identify any known structures with Z-scores > 10. Highest Z-scores of 5.0 were obtained for PDB (<http://www.rcsb.org>) codes 3IHV and 3DWL, which correspond to a sugar-binding protein SusD homolog (sequence identity = 8%; root mean square deviation = 3.6 Å for 112 C $\alpha$  atomic pairs) and subunit 5 of the actin-related protein 2/3 complex (sequence identity = 12%; r.m.s.d. = 3.4 Å for 95 C $\alpha$  atomic pairs), respectively. It is likely that the structural similarities of these two proteins to ScNup133(944–1157) are due to geometrical restraints on the packing of helices rather than to a common ancestor in evolution. The absence of the human Nup133 structure among the top scoring DALI hits came as a surprise, prompting us to measure the SAXS profile of ScNup133(944–1157) to determine whether or not the arrangement of helices observed in our crystal structure is an artifact of crystallization.

The merged experimental SAXS profile closely resembles the SAXS profile calculated from the complete model of ScNup133(944–1157) obtained with the X-ray structure of [Fig. 1(B)]. The measured radius of gyration ( $R_g$ ) of  $18.8 \pm 0.5 \text{ Å}$ , obtained from AutoRg,<sup>35</sup> compares within the experimental errors with the  $R_g$  of 18.6 Å for

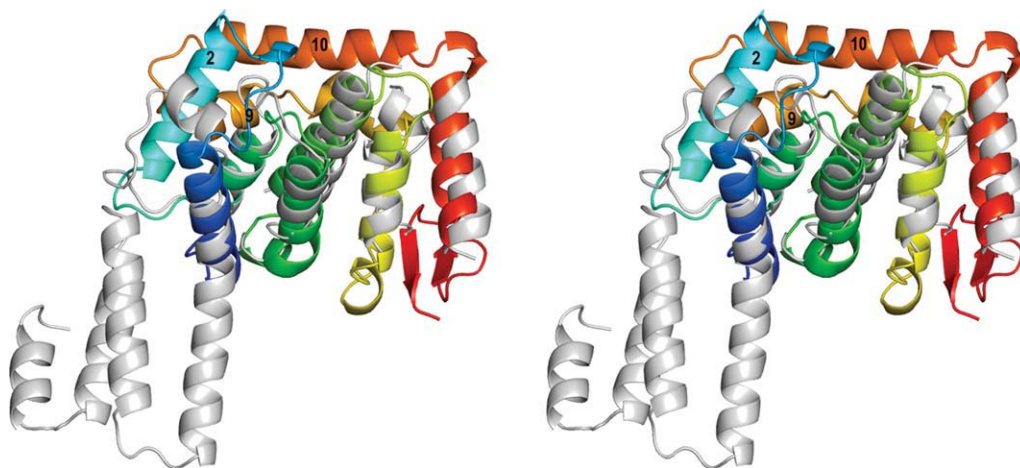


**Figure 1**

A: Stereo view of a ribbon representation of ScNup133(944–1157). Secondary structure elements were assigned by DSSP.<sup>31</sup> The fold is colored blue to red from the N- to C-termini. B: Comparison of the merged experimental SAXS profile (red) of ScNup133(944–1157) with SAXS profiles computed by IMP-FoXS<sup>32,33</sup> (blue) for the complete model of ScNup133(944–1157). Inset shows the SAXS profiles in the Guinier plot.  $D_{\text{max}}$  of radial distribution function,  $P(r)$  is 56.25 Å. C: The shape of ScNup133(944–1157) derived from the experimental SAXS profile, shown in two orthogonal orientations.

the crystal structure and  $R_g$  of 19.3 Å for the complete model. The ab initio shape reconstructed from the merged experimental SAXS profile with DAMMIF<sup>28</sup> [Fig

1(C)] and GASBOR<sup>29</sup> (not shown) matches the overall structure of ScNup133(944–1157). Therefore, SAXS analysis indicates that the compact fold of ScNup133



**Figure 2**

A: Structural superposition of the C-terminal domains of Sc and Hs Nup133. ScNup133(944–1157) is shown as in Figure 1(A) and HuNup133(934–1156) is shown in gray.



(944–1157) exists in solution and is not an artifact of crystallization.

### Comparison of ScNup133(944–1157) with human Nup133

Structures of human Nup133 residues 934–1156 [HsNup133(934–1156)] have only been determined in complexes with Nup107 (PDB Codes 3CQC,<sup>10</sup> 3CQG,<sup>10</sup> and 3I4R<sup>11</sup>). The amino acid sequence identity between the C-terminal domains of Sc and Hs Nup133 is rather low, ~12%. Structure-based alignments did not yield complete coverage of the two polypeptide chains. Structure-based alignment using Multiprot<sup>35</sup> resulted in a r.m.s.d. of 1.6 Å for 88 C $\alpha$  atomic pairs with sequence identity of ~9%. The r.m.s.d. for structure-based alignment using SSM<sup>32</sup> is 2.5 Å for 98 C $\alpha$  atomic pairs with sequence identity of ~8%. In the case of ScNup133, we have lost the putative Nup84 interaction surface due to the proteolysis. Therefore, the first helix  $\alpha$ 1 of ScNup133 aligns with the  $\alpha$ 4 of the HsNup133 structure [Fig. 2]. The four helical bundle present at the center of both ScNup133(944–1157) and HuNup133(934–1156) superpose well over each other along with the  $\alpha$ 11 helix. However, HsNup133(934–1156) lacks counterparts to ScNup133(944–1157) helices  $\alpha$ 2,  $\alpha$ 9, and  $\alpha$ 10; and  $\beta$ -strands. Therefore, two homologous Nups from two distantly related organisms show significant sequence and structural divergence in their C-termini. These differences show that a single example structure from each Nup obtained from just one organism does not represent the range of structures present throughout the Eukaryota among different Nups in different states. Thus, there is a pressing need for more structures of NPC components, from various organisms, both alone and in complexes with their respective binding partners.

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