Chapter 3

Distinctive properties of the Nuclear Localization Signals of Inner Nuclear Membrane proteins Heh1 and Heh2

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Experiments presented in Figures 3.5 except D, 3.7, 3.8A and S5 were performed by RAH. Figures 3.5 D was performed by M.v.R with help from RAH. This chapter has been published in:

Abstract

Targeting of Endoplasmic Reticulum (ER)-synthesized membrane proteins to the Inner Nuclear Membrane (INM) has long been explained by the 'diffusion-retention model'. However, several INM proteins contain non-classical Nuclear Localization Signal (NLS) sequences that, in a few instances, have been shown to promote importin α/β- and Ran-dependent translocation to the INM. Here, using structural and biochemical methods, we show that yeast INM proteins Heh2 and Src1/Heh1 contain bipartite import sequences that associate intimately with the minor NLS-binding pocket of yeast importinα and, unlike classical NLSs, efficiently displace the IBB-domain in the absence of importin β. In vivo, the interactions at the minor NLS-binding pocket make the h2NLS highly efficient in recruiting importin α at the ER and drive INM localization of endogenous Heh2. Thus, h1/h2NLSs delineate a novel class of super-potent, IBB-like membrane protein NLSs, distinct from classical NLSs found in soluble cargos, and of general interest in biology.
Introduction

Transport of soluble cargos through the nuclear pore complex (NPC) is typically an active, signal-mediated, and highly regulated process, which requires soluble transport factors of the importin β superfamily (also known as β-karyopherins) and the small GTPase Ran (Bednenko et al., 2003; Cook et al., 2007; Nardozzi et al., 2010; Stewart, 2007). Transport factor-cargo complexes move through the NPC interior by interaction with phenylalanine-glycine-rich repeats present on disordered NPC proteins, the FG-Nups. Import complexes usually assemble in the cytoplasm upon recognition of a cargo nuclear localizationsignal (NLS) by β-karyopherins (Cingolani et al., 1999). This interaction can be direct (Cingolani et al., 2002) or mediated by transport adaptors such as importinα and snurportin (Lott et al., 2010). Importin α is made up of ten stacked Armadillo (Arm) repeats, each formed by three α helices (Goldfarb et al., 2004; Pumroy and Cingolani, 2015), and binds classical NLS (cNLS) substrates, exemplified by the SV40 T-large antigen monpartite NLS and the nucleoplasmin bipartite NLS. The basic side chains of an NLS occupy a shallow groove within the Arm repeats 2–4 of importin α, known as the major binding site, as well as a minor binding site between Arm repeats 7–8. At each site, as many as five points of contact between NLS and importin α have been identified (referred to as P1–P5 and P10–P50 at major and minor binding site, respectively) (Chang et al., 2012, 2013; Chen et al., 2005; Conti and Kuriyan, 2000; Conti et al., 1998; Fontes et al., 2000, 2003; Giesecke and Stewart, 2011; Lott et al., 2011; Marfori et al., 2012; Roman et al., 2013).

Unlike soluble cargos, significantly less is known about trafficking of membrane-embedded cargos to the nuclear envelope (NE) (Antonin et al., 2011; Burns and Wente, 2012; Laba et al., 2014; Zuleger et al., 2012). Proteomic approaches have identified close to 100 NE transmembrane proteins (NETs) (Schirmer et al., 2003), many linked to genetic diseases known as laminopathies (Capell and Collins, 2006), but to date specific localization at the inner nuclear membrane (INM) has been proven for only a few proteins. Morphologically, the NE is composed of an outer and an inner membrane, which have distinct protein composition. The outer nuclear membrane (ONM) is contiguous with the ER so that membrane proteins destined for the INM and synthesized in the ER can diffuse laterally through the ER membrane system and the ONM until they encounter NPCs. At the NPCs, the INM and ONM are continuous with the...
pore membrane, and so the transmembrane (TM) domain of a membrane protein can pass from the ONM to the pore membrane. Hence, the proteins can reach the INM via a continuous membrane system (Powell and Burke, 1990), where it is finally retained upon binding to other NE components (also known as the diffusion-retention model [Ellenberg et al., 1997; Smith and Blobel, 1993; Soullam and Worman, 1993]).

Over the past decade, several lines of evidence have suggested that, in addition to diffusion-retention, other mechanisms must exist whereby the NPC plays an active role in trafficking membrane proteins to the INM (Ohba et al., 2004). In higher eukaryotes, several important INM-localized membrane proteins such as POM121, UNC-84, and Sun2 were shown to use importin α-dependent NLSs (Funakoshi et al., 2011; Tapley et al., 2011; Turgay et al., 2010; Yavuz et al., 2010). In Saccharomyces cerevisiae, INM proteins Src1/Heh1 and Heh2 (orthologs of mammalian MAN1 and LEM2) have NLSs that, like soluble proteins, bind importin-α/β (named Kap60/Kap95 in yeast) to promote nuclear translocation; deletion of such NLSs or lack of functional Kap60, Kap95, or Ran hydrolysis results in mislocalization (King et al., 2006). More recently, it was found (Meinema et al., 2011) that the NLS together with an intrinsically disordered (ID) linker of ~180 to 230 amino acids long in the extra-luminal surface of Heh1 and Heh2 is essential and sufficient for INM targeting. The long ID linkers are proposed to facilitate recruitments of importin-α/β and project the highly basic NLSs inside the NPC, allowing for importin α/β-mediated nuclear import. As for cNLS-bearing cargos, nuclear targeting depends on importin β interaction with FG repeats inside the NPC and RanGTP hydrolysis (Meinema et al., 2011). Truncated isoforms of Kap60 lacking the importin-β binding (IBB) domain have been implicated in nuclear import of Heh2 (Liu et al., 2010), although it is unclear how these isoforms can promote passage through the NPC in the absence of Kap95, since Kap95 is absolutely essential for Heh2 localization to the INM (King et al., 2006; Meinema et al., 2011, 2013). Thus, there is mounting evidence in the literature for the existence of a dedicated import pathway for INM proteins that requires importin α/β binding to a special NLS exposed on the extra-luminal domain of INM proteins. To obtain a quantitative description of the structure, recognition, and potency of a membrane protein NLS, we have carried out a structural biochemical analysis of Heh2 and Heh1 NLS sequences (abbreviated as h1NLS and h2NLS) complemented by an in vivo study of h2NLS karyophilic properties.
Results

Crystallization of Heh1 and Heh2 NLS Sequences with Kap60

S. cerevisiae INM proteins Heh1 and Heh2 contain long NLSs characterized by highly basic NLS boxes and a variable intra-NLS sequence, of 8–12 residues or even longer and commonly found in classical bipartite NLSs (Jans et al., 2000) (Figure 1A), but falling well within a more recent description (Lange et al., 2010). In vitro, peptides encoding h1NLS and h2NLS are prone to aggregation and highly susceptible to proteolysis, which hampers structural analysis. To study the interaction with Kap60, we co-expressed plasmids encoding Kap60 lacking the IBB (ΔIBB-Kap60) and GST-tagged h1NLS (residues 171–221) or h2NLS (residues 100–137). We then performed a one-step affinity purification of homogeneous ΔIBB-Kap60:NLS complexes. Co-expression was effective in preventing proteolytic degradation of the highly basic NLSs, essential to obtain well-ordered crystals. The structures of ΔIBB-Kap60 bound to h2NLS and h1NLS were solved by molecular replacement and refined to an Rwork/free of 18.9%/22.7% at 2.50 Å resolution and 19.6%/21.5% at 2.25 Å, respectively (Table 1). Both crystal structures revealed strong S-shaped electron density running along the Kap60 concave surface, mainly localized at the major and minor NLS-binding boxes and weak density between these two boxes. We will first describe the structure of h2NLS that has continuous density between the two boxes, and then that of h1NLS.

h2NLS Binds the Arm-Core of Kap60 Like an IBB domain

The structure of h2NLS bound to Kap60 can be divided in three regions, which make over 50 close contacts with the Kap60 Arm-core, burying 3,510 Å² of solvent-accessible surface area (Figures 1B and 1C; Figure S1). The first region includes h2NLS residues 100–105, which bind within (and downstream of) Kap60 minor NLS-binding site (Arm 7–8). This region has the lowest refined B-factor (~36.2 Å²) in the h2NLS model. It is superimposable to the smaller NLS box of NP-NLS (Conti and Kuriyan, 2000) and to other non-classical NLSs that bind exclusively (or preferentially) to the importin α minor NLS site (Chang et al., 2012, 2013; Giesecke and Stewart, 2011; Lott et al., 2011) (Table S1). Unlike cNLSs that usually have only two basic residues at the minor NLS-binding site, four basic amino acids in h2NLS (102-KRKR-105) insert their side chains deeply inside Kap60 groove, making ~15 close contacts (Figure 1C), of which
R103 occupies position P2’. The second region starts after R105, where the NLS backbone makes a 90° turn to form a 3/10 helix, H1 (105-REQ-107), which connects via a short linker (108-ISTDNE-113) to a second helix, H2 (114-AKMQL-118), followed by a short stretch (119-IEEKS-123) (Figures 1B and 1C). Both helices and linker have weak electron density (Figure S1) and high B-factor in our final model (~108.9 Å²). This region of h2NLS makes minor contacts with the Kap60 surface and is highly variable in other putative membrane protein NLSs (Lusk et al., 2007). The third structural region of the h2NLS contains seven consecutive basic residues (124-PKKRRKKRS-132), which span within (and upstream of) the major NLS-binding site of Kap60 (Arms 1–4) (Figures 1B and 1C). The average refined B-factor of this region is ~68 Å², higher than at the minor NLS box: only residues 125-KKKR-128 at position P1–P4 (Table S1) have clear side-chain density (Figure S1), while only main-chain atoms are visible for the residues 129-KKR-131. Thus, h2NLS binds Kap60 like a classical bipartite NLS but makes more extensive contacts at the minor NLS box than seen in the structure of Kap60 bound to NP- NLS (Conti and Kuriyan, 2000).
Table 1
Data Collection and Refinement Statistic

<table>
<thead>
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<th>Data Collection</th>
<th>ΔIBB-Kap60:h2NLS</th>
<th>ΔIBB-Kap60:h1NLS</th>
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<td>C2</td>
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<tr>
<td>Redundancy</td>
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<td>3.8 (3.7)</td>
</tr>
</tbody>
</table>

Refinement

| Resolution (Å) | 30-2.50 | 30-2.25 |
| No. Reflections| 40,407  | 25,9    |
| Rwork/Rfree    | 18.9/227| 19.6/21.5|
| No. atoms      |         |         |
| Protein        | 6,556   | 3,285   |
| Ligand (h1/h2NLS) | 449   | 148    |
| Water          | 277     | 113     |
| B-factors (Å)  |         |         |
| Protein        | 45.7    | 62.6    |
| Ligand/ion     | 72.2    | 85.9    |
| Water          | 40.0    | 54.0    |
| Rmsd           |         |         |
| Bond lengths (Å)| 0.008 | 0.003 |
| Bond angles (°)| 1.00  | 0.8    |

Value in parentheses are for highest-resolution shell.
The Rfree was calculated using 5% of randomly selected reflections.

Heh1 NLS Makes Strong Contacts at the Minor NLS-Binding Box

The exact boundaries of the Heh1 NLS were unknown before this study, although it was shown that a region between residues 173 and 220, encoding several basic patches similar to a cNLS, and a ~200-residue unfolded linker were required and sufficient for nuclear import (Meinema et al., 2011). The crystal structure of ΔIBB-Kap60 crystallized in complex with a 50-mer spanning Heh1 residues 171–221 (Figure 1A) has density only for the first half of the Heh1 construct (residues 173–195) (Figure 2A), while no discernable electron density was observed for residues 196–221. h1NLS resembles h2NLS closely and occupies both minor and major NLS-binding sites. The first basic box (173- RKKRK-177) of h1NLS binds intimately the minor NLS pocket, while, unexpectedly, a minimal basic stretch of residues (189-SKEND-195)
occupies the major NLS box. The canonical intra-NLS spacer of 11 residues has poor density and some of its residues (180-DSDDWSES-187) were not modeled in the final structure. Noticeably, the five basic amino acids in h1NLS that bind the minor NLS pocket engage in nearly 20 close contacts with Kap60 Arm 6–8 (Figure 2B), of which R176 occupies position P2’ (Table S1). Instead, at the major NLS-binding pocket, only P2 and P5 are occupied by lysines (Figure 2B), whereas non-basic side chains interact at P1, P3, and P4, as previously seen for PLSCR1 NLS (Chen et al., 2005) (Table S1). Overall, Kap60 recognizes the NLS of Heh1 and Heh2 using a combination of electrostatic and hydrophobic contacts with main- and side-chain atoms. Arrays of Asn (Conti et al., 1998) projecting from Kap60 Arm-core stabilize the NLS backbone, while conserved Trps (Figures 1C and 2B) engage in hydrophobic and cation-π interactions (Koerner et al., 2003) with the critical side chains of R103/176 and K126/190, which occupy P2’ and P2 positions at minor and major NLS-binding boxes, respectively.

Figure 3. 2 Mapping crystallographically h1NLS in complex with Arm-core of Kap60. Crystal structure of ∆IBB-Kap60 (gray surface) in complex with h1NLS (red ribbon). The dotted line indicated residues in the intra-NLS linker that is poorly visible in the electron density and that was not included in the final model. (B) Schematic diagram of the interactions between h1NLS (in red) and Kap60 residues (in gray) in a distance range of 2.5-4.5 Å.
h1NLS and h2NLS Bind DIBB-Kap60 with Nanomolar Affinity

The intimate association of h1NLSs and h2NLS with Kap60 observed crystallographically prompted us to measure their binding affinity for Kap60. Using nano-isothermal titration calorimetry, we measured the heat released upon titration of increasing concentrations of maltose-binding protein-tagged h1NLS (MBP-h1NLS) or h2NLS (MBP-h2NLS) into a cell containing ΔIBB-Kap60 (Figure 3A). This analysis yielded an equilibrium dissociation constant ($K_D$) of 27.3±8 nM for h2NLS and 30.5±10 nM for h1NLS, which is somewhat lower than the $K_D$ of a control NP-NLS for ΔIBB-Kap60 measured under identical experimental conditions ($K_D = 46.0 \pm 14$ nM) (Figure S2). The observation that the two membrane protein NLSs bind Kap60 with similar affinity, although h1NLS has only two basic residues at the major NLS-binding box (Figure 2B) versus seven in h2NLS (Figure 1C), suggests a minimal contribution of this moiety in the overall binding affinity for Kap60. This is clearly not the case for cNLSs, which are disrupted by a single point mutation at P2 in the major NLS-binding box (Colledge et al., 1986; Kalderon et al., 1984). To test this idea, we introduced Ala mutations at position P2' and P2 of h1/h2NLSs and measured their effect on the overall equilibrium binding affinity for ΔIBB-Kap60. A mutation at position P2 reduced moderately (~2-fold) the h1NLS affinity for ΔIBB-Kap60 ($K_D = 68.6 \pm 14$ nM), consistent with the small number of contacts made at the major NLS-binding site (Figure 3B), whereas a 4-fold drop in affinity was caused by an Ala substitution at P2' ($K_D = 123.0 \pm 8.6$ nM) (Figure 3C). A similar effect was seen in h2NLS, where a mutation at P2 yielded a 4-fold drop in binding affinity for ΔIBB-Kap60 ($K_D = 106.4 \pm 15$ nM) (Figure 3B), while a 5-fold destabilization was caused by an Ala substitution at P2' ($K_D = 131.5 \pm 27$ nM) (Figure 3C). Combining mutations at P2' and P2 did not significantly aggravate loss of binding affinity for ΔIBB-Kap60 ($K_D = 139.5 \pm 26$ nM and 167.7 ± 32 nM for h1NLS and h2NLS, respectively) (Figure 3D) as compared with single point mutants at P2', confirming that the overall affinity of membrane protein NLSs for Kap60 depends primarily on structural determinants at P2', in the minor NLS-binding box.
h1NLS and h2NLS Compete off the IBB Domain in the Absence of Importin β

Superimposition of ΔIBB-Kap60 bound to Heh2 or Heh1 NLSs with FL-Kap60, previously solved as part of an export complex (Matsuura and Stewart, 2004), revealed a striking structural resemblance between the membrane protein NLSs and the IBB domain (rmsd 1.1 Å) (Figure 4A). The h2NLS, which has a continuous trace between NLS boxes, the h1NLS, and the IBB adopt a nearly identical conformation at the minor and major NLS-binding pockets of Kap60 with a striking conserved lysine at position P2 (IBB-54/h2NLS-126/h1NLS-190) and an arginine at P2’ (IBB-34/h2NLS-103/h1NLS-176). In contrast, the intra-NLS regions are partially helical in h2NLS (residues 106-120), not visible in the structure with h1NLS (residues 182-187) and random coiled in
IBB (residues 37–49), suggesting this region makes non-essential contacts with Kap60. However, despite the structural similarity to an IBB, h1/h2NLSs do not associate directly with Kap95 (data not shown), suggesting these NLSs mimic only the importin α-bound conformation of IBB, which is mainly unstructured, but cannot adopt the helical conformation of IBB induced upon binding to importin β (Cingolani et al., 2000; Mitrousis et al., 2008).

As it was previously shown that the h2NLS can bind FL-Kap60 in the absence of Kap95 (King et al., 2006), we hypothesized that a mutation at P2’ affects the way h2NLS competes off the IBB domain of Kap60. To test this hypothesis, we tried to measure association of h1/h2NLSs with FL-Kap60 using isothermal titration calorimetry (ITC) but obtained uninterpretable binding data, likely due to the concomitant presence of two binding events, namely the intra-molecular dissociation of IBB from Kap60 Arm-core and the intermolecular association of h1/h2NLS with Kap60. To overcome this problem, we turned to an on-bead binding assay (Pumroy et al., 2015), where GST-tagged FL-Kap60 (GST-FL-Kap60) and GST-ΔIBB-Kap60 were immobilized on glutathione beads and incubated with a 2-fold molar excess of h1/h2NLSs or control NP-NLS. In the absence of importin β, the IBB domain binds the Arm-core preventing association of cNLS cargos (Kobe, 1999) (Figure 4B). Instead, h2NLS bound stoichiometrically both to FL-Kap60 and to Kap60 Arm-core, confirming this NLS can efficiently bypass IBB autoinhibition. The h1NLS was also able to overcome autoinhibition, yet to a lesser extent compared with h2NLS and as much as 40% of MBP-h1NLS was recovered bound to beads after 15 min incubation (Figure 4B). Mutation at P2’, but not P2 (Figure 4C), completely disrupted the interaction of h2NLS with FL-Kap60, rendering h2NLS indistinguishable from NP-NLS. Similar results were obtained for h1NLS, which, although less effective at displacing the IBB, was disrupted by a single point mutation at P2’ but not P2 (Figure 4D). Thus, the membrane protein NLSs of Heh1 and Heh2 adopt an IBB-like structure that combines binding determinants seen in the recognition of cNLSs, as well as a deeper interaction at the minor NLS-binding site and particularly the P2’ position, which make these NLSs able to bypass IBB autoinhibition.
In Vivo Potency of h2nls Depends on P2’ Position

To complement our in vitro studies we sought to confirm the importance of the interaction with the minor NLS-binding site, particularly at the P2’ position, for transport of membrane proteins in vivo. We focused on h2NLS, which was previously characterized in detail in live cells, both in the context of the full-length protein and in reporter proteins (King et al., 2006; Meinema et al., 2011, 2013). The advantage of using Heh2-derived reporter proteins is that they are mobile within the network of the NE and ER (Meinema et al., 2011, 2013) because they lack domains that contribute to
nuclear retention, such as the LEM domains found in Heh1 and Heh2 (Heh2 domain composition is schematically illustrated in Figure 5A). Being mobile, their distribution in the network of the NE and ER reflects their nuclear import rates: for a Heh2-reporter with h2NLS and ID linker, we find a higher fluorescence at the NE than at the peripheral ER, while a reporter lacking the NLS or ID linker shows similar levels of fluorescence in the entire NE-ER network (Meinema et al., 2011). The nuclear location is completely dependent on Kap95, as demonstrated by conditionally tethering Kap95-FRB to Pma1-FKBP at the plasma membrane, which results in gradual decrease in the NE/ER ratio (Figure 5B) (Meinema et al., 2011, 2013), arguing against a recent report that Kap60 isoforms lacking the IBB, and therefore unable to heterodimerize with Kap95, are responsible for Heh2 translocation to the INM (Liu et al., 2010).

To test the importance of position P2’in vivo, we introduced Ala substitutions at position P2’, P2, and P2’/P2 in a soluble GFP-h2NLS fusion (Figure 5C) and in a membrane-embedded Heh2-based reporter (Figure 5D) and imaged their subcellular localization. As shown previously (Meinema et al., 2011), the karyophilic properties of the h2NLS are so strong that many cells showed no cytosolic h2NLS-GFP and the N/C ratio is very high (N/C ~75 is likely an underestimation). The reduction in nuclear accumulation of soluble (GFP-NLS, Figure 5C) and TM (GFP-h2NLS-L-TM, Figure 5D) reporter proteins was most severely affected by substitution at position P2’ in the minor NLS box. In the case of the TM reporter protein, the NE/ER ratio was moderately reduced when introducing the P2 mutation (NE/ER ratio 21.6 ± 2.7 and 31.5 ± 2.5 for P2 and wild-type [WT]), but mutation at P2’ resulted in complete loss of nuclear accumulation (NE/ER ratio 2.8 ± 0.6), comparable with a ΔNLS mutant (NE/ER ratio 2.3 ± 0.2 in Meinema et al. (2011). Likewise, the double mutant (P2’/P2) localized similar as the P2’ mutant, confirming the dominant negative role of the P2’ mutation.

Knowing now that the interactions at the minor binding site of Kap60 are critical, we compared the karyophilic properties of the h2NLS with known NLSs. In the context of the TM reporter proteins, the h2NLS led to NE/ER ratios that were approximately 3-fold higher compared with a single partite variant of the h2NLS (lacking 102-KRKR105) or 8-fold higher compared with a cNLS (Meinema et al., 2011). Complementing these studies we also replaced the h2NLS with the NP-NLS and observed that GFP-NP-L-TM is accumulated approximately 2.5-fold lower than with h2NLS (NE/ER ratio 14.7 ± 1.2) (Figure 5E), which is consistent with the reduced NE targeting of full-length Heh2.
carrying the NP-NLS (King et al., 2006). The high-affinity NLS of Cdc6 (Hahn et al., 2008) behaved similarly to the NP-NLS with NE/ER ratio 13.7 ± 1.3 (Figure 5E). Reinforcing the specific role for Kap60 and Kap95 in nuclear import of Heh2 (King et al., 2006), the high-affinity Kap104-dependent NLS of Nab2 had a similar NE/ER ratio (NE/ER ratio 3.2 ± 0.4) (Figure 5D) as observed without an NLS (NE/ER ratio 2.3 ± 0.2) (Meinema et al., 2011). Similarly, NP-NLS fused to a soluble import cargo (GFP) was much less efficient than h2NLS in promoting nuclear translocation (N/C ratio 3.2 ± 0.2) (Figure 5C). We conclude that, in vivo, h2NLS is an exceptionally potent import signal.

Figure 3.5 Quantitative analysis of h2NLS karyophilic properties. (A) Cartoon showing the domain composition of Heh2 where GFP is in green, NLS in red, Heh2’s ID linker represents a curved line and the TM domain is in black quantification of average N/C-ratios. (B) Deconvolved wide-field images of the Heh2-based transmembrane reporter protein expressed in the Kap95AA strain (Haruki et al., 2008) (No RAP) and when Kap95-FRB is conditionally trapped at Pma1-FKBP at the plasma membrane upon addition of rapamycin (RAP). (C) Confocal fluorescent images of yeast expressing GFP fused to indicated NLSs: WT h2NLS, h2NLS mutants at position P2’ and P2 and NP-NLS and average NE/ER-ratios over n cells. (D) Confocal fluorescent images of yeast expressing GFP-h2NLS-L-TM with mutation at position P2’, P2, P2’/P2 and quantification of average NE/ER-ratios over n cells (E) Same as (D) but with different indicated NLSs. Scale bar is 5 μm and SEM is indicated.

R103 at Position P2’ Is Critical for Heh2 Function and Translocation to the INM

To test the importance of the interaction at the minor binding site P2’ position, we sought to determine the localization of full-length N-terminal GFP-tagged Heh2 expressed from the chromosome from its endogenous promoter (Figure S4). This is more physiological than the Heh2 reporter, although this protein can engage in protein-protein interactions at the INM that retain it in the nucleus. Indeed, the sole mutation R103A at the P2’ resulted in a complete loss of the NE specific localization,
indistinguishable from that of a ΔNLS mutant (Figure 6A).

To gain further insight into the \textit{in vivo} relevance of the P2’ mutation, the mutation was introduced in a strain lacking NUP84. Previously, it was shown that the double mutant \textit{nup84Δheh2Δ} (Yewdell \textit{et al.}, 2011), in contrast to the single mutants, fails to grow, and the double mutant \textit{nup84Δheh2Δh2NLS} is synthetic sick compared with the single mutants. Consistent with the complete loss of accumulation of the P2’ mutant, and the \textit{in vivo} relevance of this accumulation, the double mutant \textit{nup84Δheh2P2’} is also synthetic sick, indistinguishable from \textit{nup84Δheh2ΔNLS} (Figure 6B). Thus mutation of the P2’ position in h2NLS correlates with loss of function \textit{in vivo} both on the level of cellular localization and cell fitness.

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{figure3}
\caption{6 Mutation of R103 at position P2’ abolishes NE accumulation of Heh2 and the double mutant with \textit{nup84Δ} is synthetic sick. (A) Deconvolved wide-field images of yeast expressing native levels of GFP-Heh2 with WT NLS (h2NLS), without the NLS (ΔNLS), and Heh2 with the P2’ mutation. Scale bar: 5 μm. (B) Synthetic sick/lethal interaction using tetrad dissection of \textit{nup84Δ} expressing WT (h2NLS) and mutant variants of Heh2 (ΔNLS, P2’) or no Heh2 (Heh2Δ). Each tetrad is oriented vertically and represents the meiotic progeny of a heterozygous diploid between GFP-\textit{HEH2-NAT}/\textit{NUP84} and \textit{HEH2}/\textit{nup84::KANMX}. Two representative tetrads for each double mutant are shown. The genetic background of each spore is identified by the presence of the NAT and KAN marker, respectively. The double mutant spore colonies are enclosed in circles whereas single mutants are enclosed in squares or diamonds, and WT strains are not enclosed. \textit{See also Figure S4.}}
\end{figure}

\section*{Position P2’ Is Critical to Retain NLS-Bound Kap60 at the ER}

Next, we aimed to confirm our \textit{in vitro} data by showing that the binding to Kap60 depends on the interaction at the P2’ position. We thus assessed the binding of Kap60 (and Kap95) to the h2NLS and the P2’ mutant \textit{in vivo}, using an assay in which we monitor co-enrichment of Kap60-GFP with the membrane re- porters (Meinema \textit{et al.}, 2013). While Kap60-GFP normally does not enrich at the peripheral ER, it did so in 45% of the cells (n = 86) expressing an ER-localized h2NLS-containing reporter protein (mCherry-h2NLS-L(37)-TM) (Figure 7). This protein lacks a functional linker domain so that it remains ER localized (and does not accumulate at the INM) (Meinema \textit{et al.}, 2013).
Co-enrichment of the reporter protein and Kap60-GFP reflects binding of Kap60 to the h2NLS, because cells expressing a reporter that lacks an NLS (mCherry-L-TM) did not show Kap60-GFP at the peripheral ER (n = 47) (Figure 7). Interestingly, cells that expressed the reporter protein with the mutant NLS (mCherry- h2NLSP2’-L-TM) also did not show Kap60 enriched at the peripheral (n = 54) (Figure 7), consistent with the dramatic reduction in import efficiency to the INM. Thus, h2NLS recruits Kap60 by making a crucial contact with the minor NLS-binding box that is critically dependent on R103 at position P2’.

Figure 3. 7 In vivo analysis of h2NLS interaction with Kap60. Deconvolved wide-field images of cells co-expressing Kap60-GFP with mCherry-tagged reporter proteins mCh-h2NLS-L(37)-TM, mCh-h2NLS P2’-L-TM or mCh-L-TM. Scale bar is 5 µm and SEM is indicated. See also Figure S5 and Supplemental Experimental Procedures.

Nup2 and h2NLS compete at the Minor NLS-Binding Site

Next, we asked whether Nup2, a mobile nucleoporin that also binds the minor NLS-binding site of Kap60 (Matsuura et al., 2003; Pumroy et al., 2012), plays a role in disassembly of h2NLS from Kap60, as proposed for cNLS cargos (Dilworth et al., 2001; Hood et al., 2000; Solsbacher et al., 2000). Accumulation of GFP-h2NLS-L-TM in a nup2Δ strain was approximately 2.5-fold decreased compared with a WT strain, consistent with measurements on full-length Heh2 (King et al., 2006) (Figure 8A). Since Nup2 also functions in the recycling of Kap60 back to the cytoplasm (Solsbacher et al., 2000), and the knockout suffers from other cellular effects like an mRNA export defect (Casolari et al., 2004; Dilworth et al., 2005), the results were not readily interpreted. However, the localization of a similar reporter protein containing NP-NLS (GFP-NP-L-TM) was not dramatically affected, pointing to a specific role for Nup2 in import of the h2NLS cargo.
Figure 3. Role of Nup2 in displacement of Heh2 from Kap60. (A) Confocal fluorescence images of a wild type yeast strain (BY4742) and a nup2Δ knockout strain expressing GFP-h2NLS- L-TM and the reporter with mutations at position P2’, P2, and NP-NLS, as well as quantification of average NE/ER-ratios (average of ~30 cells). SEM, and 5 μm scale bar are indicated. (B) Nup2-mediated displacement of ΔIBB-Kap60 from GST-h2NLS (and its mutants at P2’, P2 and P2'/P2) coupled to glutathione beads. The complex was challenged with 1,25-10 fold molar excess of MBP-Nup2 (res1-51) and ΔIBB-Kap60 left on beads is quantified on the right panel (error bars from averaging three independent experiments). (C) Model for recognition and association of a membrane protein NLS to auto-inhibited FL-Kap60. From left to right are schematic illustration of auto-inhibited FL-Kap60, an ER-synthesized membrane protein (like Heh2) projecting an h2NLS-like import sequence in the cytoplasm and two putative snapshots of FL-Kap60 partially and fully bound to the membrane protein NLS.
(Figure 8A). To determine if the N-terminal 51 residues of Nup2 were sufficient to dissociate h2NLS from ΔIBB-Kap60, we immobilized a stoichiometric complex of ΔIBB-Kap60:h2NLS on glutathione beads and challenged it with increasing molar excess (from 1.25 to 10x) of purified Nup2 (residues 1–51), followed by SDS-PAGE and quantification (Figure 8B). Notably, a 10-fold excess of Nup2 dissociated as much as 60% of the otherwise very stable ΔIBB-Kap60:h2NLS complex. Ala substitution at position P2’ enhanced Nup2-mediated displacement of ΔIBB-Kap60 from h2NLS more markedly than the mutant at P2 (Figure 8B). Thus, Heh2 association to Kap60 is affected by Nup2, consistent with an intimate interaction of both proteins with the minor NLS-binding site of Kap60.

**Distinctive Features of Membrane Protein NLSs**

In this study, we have characterized the NLS of yeast INM protein Heh2 and Heh1 and defined a set of molecular properties that we propose are distinctive of these membrane protein NLSs. First, h2NLS resembles the IBB domain of importin α in the autoinhibited conformation, as opposed to a bipartite NLS. Analogous to known IBBs (Lott and Cingolani, 2011), h2NLS accommodates intra-NLS residues as partially folded helices that make minimal contacts with Kap60. Second, both h1 and h2NLSs bind ΔIBB-Kap60 with low nanomolar affinity, comparable with NP-NLS, but their assembly to Kap60 is different from cNLS. The dominant negative mutation in h2NLS that disrupts nuclear localization is at position P2’ of the minor NLS-binding site. This is distinct from a cNLS (Colledge et al., 1986; Kalderon et al., 1984), where mutation at P2’ marginally disrupts nuclear localization (Robbins et al., 1991), reinforcing the idea that h2NLS is not a simple variation of a classical bipartite NLS. Third, h2NLS and to a lesser extent h1NLS compete off the IBB domain in the absence of importin β, which predicts a reduced autoinhibitory role of IBB on membrane protein cargos trafficking from the ER to the INM. This is similar to the influenza polymerase subunit PB2 (Pumroy et al., 2015), which also overcomes IBB autoinhibition by making strong contacts at the minor NLS box. Fourth, nucleoporin Nup2 plays a critical role in displacement of h2NLS from Kap60 by directly competing for binding to the minor NLS-binding site, which provides an anchoring point to both h2NLS and Nup2’s N-terminal NLS-like moiety (Matsuura et
A potential multi-step mechanism describing the recruitment of Heh2 membrane protein NLS by Kap60 can be hypothesized (Figure 8C). Recognition of h2NLS begins at the minor NLS-binding pocket, where the basic box 102-KRKR-105 of h2NLS competes off the equivalent region of IBB (33-RRRR-36) (Table S1). Whereas all four basic residues in h2NLS insert at the Kap60 helical interface between Arm 7–8, only three Args in IBB (at position P1', P2', and P4') make contacts with the minor NLS-binding site, projecting the guanidinium group of R35 (at position P3') at the surface of Arm 7 (Figure 3A). This initial interaction cements h2NLS to the Kap60 minor NLS-binding pocket, increasing its local concentration, and allows zipping to the major NLS site, where the major NLS box 54-KRR-56 of the IBB is readily competed off, overcoming IBB autoinhibition and displacing the IBB in the absence of Kap95. We speculate that early recruitment of importin α could occur while a membrane protein is being synthesized and/or inserted at the ER membrane.

**Physiological Significance of Membrane Protein NLSs**

What is the advantage of bearing a membrane protein NLS instead of a classical bipartite NLS? Although a conclusive answer to this question will require further in-depth analysis of additional membrane protein NLSs, especially from higher eukaryotes (Lusk et al., 2007), and a complete understanding of inner membrane protein full-length 3D-structure (in addition to minimal NLS fragments), a few hypotheses can be formulated on the basis of the data presented in this study. The karyophilic potency of an h2NLS-like import signal is likely to aid in all steps of membrane protein translocation to the INM, thereby providing a selective biological advantage over cNLSs. At first in the cytoplasm, during import complex assembly, we propose membrane protein NLSs facilitate recruitment of karyopherins and formation of a productive membrane-bound import complex. Unlike soluble NLS cargos moving fast by 3D diffusion, membrane proteins move much slower in the 2D plane of the membrane (Meinema et al., 2013). They thus could have a reduced probability to encounter karyopherins, which are soluble factors. However, as shown for h1/h2NLSs in this study, the ability of recruiting importin α in the absence of importin β possibly compensates for the restricted 2D diffusion of membrane-embedded cargos providing a kinetic advantage over classical cargos that assemble into productive import complexes
only when importin α and β are simultaneously present (Pumroy et al., 2015). During translocation through the NPC, although the actual mechanisms of passage are controversial and it is unclear if an import complex undergoes cycles of dissociation and re-association while moving inside the NPC (Bednenko et al., 2003), the advantage of a membrane protein NLS would be its ability to remain bound to importin α even when importin β has been displaced, possibly expediting re-formation of an import complex. Finally, membrane protein NLSs may provide a selective advantage to release cargos at the INM. After importin β- and Ran-dependent passage through the NPC, competition with Nup2 for binding to the importin α minor NLS-binding pocket is likely to promote release of membrane-embedded cargos at the INM, where NETs can be retained by binding interactions with other NE components.

In summary, the present work expands the definition of NLS and provides a framework to identify the molecular mechanisms by which ER-synthesized membrane proteins translocate to INM to play a critical role in nuclear signaling.

Methods

Biochemical Techniques

ΔIBB-Kap60 was co-expressed with GST-h1/h2NLS in E. coli strain BL21-CodonPlus (DE3)-RIL (Stratagene) for 6 hr at 30°C. ΔIBB-Kap60 bound to GST-h1NLS or GST-h2NLS was purified on glutathione-resin (GenScript) and after cleaving off the GST with PreScission Protease, the complex was purified over a Superdex 200 column (GE Healthcare) equilibrated in gel filtration buffer (20 mM Tris [pH 8.0], 150 mM NaCl, 5 mM β-mercaptoethanol, and 0.2 mM PMSF). All GST-tagged constructs used in this study were purified as described above. GST-Nup2 was expressed as described for human Nup50 (Pumroy et al., 2012). All His-MBP-tagged constructs were purified over His-resin (GenScript) followed by gel filtration chromatography. For pull-down assays and ITC analysis, see Supplemental Experimental Procedures.

Crystallographic Studies

Crystals of ΔIBB-Kap60 bound to h1NLS or h2NLS were obtained by mixing equal volume of gel filtration-purified complex at 12.5 mg/ml with 100 mM ammonium acetate, 20% PEG 8000, 100 mM BisTris (pH 6.0) and equilibrating the droplet against 600 ml of the same precipitant. 25% glycerol was added as cryoprotectant before flash-
freezing at -170°C. Crystals were diffracted at beamlines X6A and X29 at the National Synchrotron Light Source (NSLS) on a Quantum Q270 and a Quantum-315r charge-coupled device (CCD) detector, respectively. Data were processed using HKL2000 (Otwinowski and Minor, 1997) and initial phases calculated using Phaser (McCoy et al., 2007). Atomic models were built using Coot (Emsley and Cowtan, 2004) and refined with phenix.refine (Adams et al., 2002). Data collection and refinement statistics are summarized in Table 1 and additional methods are in Supplemental Experimental Procedures.

**Yeast Cultivation and Microscopy**

Yeast strains used in this study are listed in Table S2 and are isogenic to S288C except the Kap95-AA strain (Haruki et al., 2008), which is W303 based (Figure 5). Cells were grown at 30°C and kept at mid-log growth phase for 24 hr before imaging. Reporters were induced at mid-log phase with 0.1% galactose for 1.5 hr (GFP reporters, Figures 5 and 8A) or 5 hr (mCherry reporters, Figure 7). Imaging for Figures 5C–5E was performed on a commercial LSM 710 confocal microscope (Carl Zeiss MicroImaging), using an objective C-Apochromat 403/1.2NA, a solid-state laser (488 nm) for excitation, and a pixel dwell times of 101–177 ms. Imaging for Figures 5B, 6, 7, 8A was on a wide-field deconvolution microscope (DeltaVision; Applied Precision/GE Healthcare), taking 60 x 0.2 mm sections, equipped with a 1003, 1.40 NA objective lens and solid-state illumination; deconvolution was performed using Softworx, ten iterations, and medium noise filtering. The images were acquired using a CCD camera (CoolSNAP HQ2; Photometrics). Data analysis is described in Meinema et al. (2013) and in Supplemental Experimental Procedures.

**Accession numbers**

The atomic coordinates and structure factors for Kap60 bound to h2NLS and h1NLS have been deposited in the PDB with accession codes PDB: 4PVZ and 4XZR.

**Acknowledgments**

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the Zernike Institute for Advanced Materials (to R.A.H.). R.A.P. is supported by NIH grant T32 GM100836. Research in this publication includes work carried out at the Sidney Kimmel Cancer Center X-ray Crystallography and Molecular Interaction Facility, which is supported in part by NCI grant P30 CA56036.

Supplementary documents are also available online (doi: 10.1016/j.str.2015.04.017)


Figure S1. Representative Fo–Fc electron density difference map of h2NLS bound to DIBB-Kap60. A sigmaA weighted Fo – Fc electron density map is displayed as a cyan mesh (contoured at 2.0σ above background) around residues 97-131 of h2NLS. The electron density was calculated using all reflections between 30-2.5 Å resolution, phases calculated from the refined DIBB-Kap60 model (lacking the h2NLS) and by applying a sharpening factor of -50A2, as implemented in Phenix.maps (Adams et al., 2002). The density is overlaid to h2NLS final model (in red) while DIBB-Kap60 is shown in light gray. The figure was generated using PyMol (DeLano, 2002)
Figure S2. Calorimetric analysis of the interaction of ΔIBB-Kap60 (in cell) with MBP-NP-NLS (in syringe). Raw data are in the top panel, and the integrated enthalpy plotted as a function of the NLS:ΔIBB-Kap60 molar ratio is shown in the bottom panel. The equilibrium dissociation constant is $K_d = 46.0 \pm 14$ nM.
Figure S3. Control pull-down assays. No interaction was observed between free MBP and GST-FL-Kap60 immobilized on glutathione beads (lanes 1,2). Similarly, no interaction was observed between free MBP-h2NLS (lanes 3-10) and MBP-h1NLSs (lanes 11-12) and free GST immobilized on glutathione beads. ‘B’ and ‘U’ indicate fractions bound to beads an unbound.

Figure S4. Western-blot analysis confirms GFP-Heh2 mutants are expressed to levels comparable to wild type of Heh2. Different cell lysates (from strains used for Figure 6) were prepared from equal amounts of cells and anti-GFP was used for immuno-detection.
Figure S5. Expression of mCherry-reporters visualized by Western blot analysis. Expression of mCherry-reporters was induced with 0.1% galactose for 5 hrs in a strain expressing chromosomal fusion of Kap60-GFP. Cell lysates are prepared from equal amounts of cells and anti-mCherry (ABcam cat #1C51) was used for immuno-detection. Pgk1 is used as loading control.

* denotes yeast importin α (Kap60).
** denotes rice importin α. In all other cases, mammalian importin α was co-crystallized with NLSs.

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Table S1. Structural alignment of h1NLS and h2NLS with other NLSs visualized crystallographically in complex with Kap60/importin α.
### I. Strains list

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### II. Plasmids list

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Table S2. List of *Saccharomyces cerevisiae* strains and plasmids used in this study.
Supplementary Experimental Procedures

Molecular biology techniques
FL-Kap60, ΔIBB-Kap60 (res. 88-530), h1NLS (res. 171-221), h2NLS (res. 97-137), NP-NLS (res. 151-172) and Nup2 (res. 1-51) were cloned between the BamHI and XhoI restriction sites of pGEX-6P-1 and of an engineered pET28a(+) vector containing the MBP gene downstream of a 6-his tag. ΔIBB-Kap60 (res. 88-530) was cloned between the NdeI and XhoI restriction sites of pET30b(+). Membrane reporters containing different NLSs (Nab2, Cdc6 and nucleoplasmin) were constructed using double fusion PCR to combine the NLSs with the fragments encoding the disordered linker region and first transmembrane domain of Heh2 (res. 137-378 encoding 'L-TM'), and then followed by the LIC method (Amberg et al., 1995; Geertsma and Poolman, 2007). The Nab2 NLS was amplified from pBT032 Nab2NLS-GFP (Timney et al., 2006), the Cdc6 NLS was amplified from genomic DNA of BY4741 and the coding sequence of NP-NLS was obtained from synthetic oligonucleotides. Soluble reporters of NP and h2NLS were generated using the LIC method with PCR fragments from pRAH15, pRAH16, pRAH17, and pRAH21 and pACM021 GFP. The pRAH26 were generated by ligating the SpeI fragments from pRAH16, with pACM061. The strain with a chromosomal tagged Kap60-GFP was obtained from Invitrogen and validated by PCR. All point-mutations were introduced using Stratagene's QuickChange site-directed mutagenesis.

Pulldown assay and quantification
Pull-down assay to study IBB-displacement assay was carried out and quantified as previously described (Pumroy et al., 2015). Briefly, 15 µL of GST-FL-Kap60 or GST-ΔIBB-Kap60 beads were incubated with 2-fold molar excess of MBP-tagged WT or mutant h1NLS, or h2NLSs or control NP-NLS. After 20 min incubation at room temperature, the unbound fraction was collected. After washing, all samples were boiled in 15 µL of 2X SDS-loading buffer and analyzed on SDS-PAGE. The experiment was repeated in triplicate. For quantification, two control gels were run to determine the linear range of Coomassie dye staining density and to confirm that GST-importin α and MBP-tagged cargos absorbed dye comparably. The upper bound of the linear range was determined to be 4 µg, so approximately 2 µg of the GST-importin αs were loaded on gel. The density of each band could be converted to a molar quantity and the molar ratio between the GST-importin α and MBP-cargo in each lane determined. Image Lab software (BioRad) was used to quantify all bands. The error bars represent the standard deviation of three independent bands of Kap60 containing the same quantity of protein and analyzed by SDS-PAGE under identical conditions.

For Nup2 competition assay, various purified recombinant GST-tagged wt or mutant NP- or h2NLSs were immobilized separately on glutathione resin. 15 µL of GST-NLS beads were incubated with 200 µg of ΔIBB-Kap60 for 20 min at room temperature. Excess ΔIBB-Kap60 was extensively washed in G.F. buffer. For ΔIBB-Kap60 displacement, 1.25, 2.5, 5 and 10 times molar excess of Nup2 was added to the beads. The beads were incubated for 20 min at room temperature followed by extensive wash. After washing, all samples were boiled
in 15 µL of 2X SDS-loading buffer and analyzed on SDS-PAGE and quantified as described above.

**Isothermal Titration Calorimetry**

ITC experiments were carried out at 30°C using a nano-ITC calorimeter (TA Instruments). MBP-tagged h1NLS, h2NLS, NP-NLS and various mutants were dissolved in G.F. buffer between 45-100 µM and injected in 2 µL increments into the calorimetric cell containing 300 µL of ΔIBB-Kap60 at 3-5 µM. The spacing between injections was 300 seconds. Titration data were analyzed using the NanoAnalyze data analysis software (TA Instruments). Heats of dilution were determined from control experiments with the ITC buffer and subtracted prior to curve fitting using a single set of binding sites model.

**Structural analysis**

All structural figures in this paper were made using PyMol (DeLano, 2002). Intra- and intermolecular contacts between atoms were determined using Monster (http://monster.northwestern.edu/help/monster2.html). Structural superimpositions were carried out using the program Coot (Emsley and Cowtan, 2004).

**Microscopy data analysis**

NE/ER- and N/C-ratios in Figure 5CDE, 8A was quantified as an average of the pixel intensity at the NE or nucleus over the average pixel intensity at the peripheral endoplasmic reticulum or cytosol as described in (Meinema et al., 2013). For assessment of co-enrichment of Kap60-GFP with mCherry-tagged membrane reporters (Figure 7), deconvolved images were double blind analyzed. In 45% of the cells with detectable mCherry-h2NLS-L(37)-TM expression, Kap60-GFP was detected at the peripheral ER at higher levels than in the cytosol (n=86). When looking only at those cells with higher expression levels of mCherry-h2NLS-L(37)-TM up to 74% of cells with showed enrichment of Kap60-GFP at the ER compared to the cytosol, consistent with (Meinema et al., 2013). Kap60 was never enriched at the peripheral ER as compared to the cytosol when co-expressed with mCherry-L-TM or mCherry-h2NLSP2’-L-TM (n=54, 47).

**Synthetic lethality screening**

DNA constructs (Table S2) encoding GFP tagged full length Heh2, GFP-tagged Heh2(ΔNLS) and GFP-tagged Heh2(P2’NLS) were fused to the NAT marker and the flanking regions as encoded immediately upstream and downstream the HEH2 ORF. The linear DNA constructs were subsequently integrated in BY4742 heh2::KAN by homologous recombination, thereby replacing the KAN marker for GFP-HEH2-NAT. Transformants were selected and analyzed for expression of the GFP tagged proteins by fluorescence microscopy. For assessing the synthetic sick and lethal phenotypes of double mutants, BY4741 nup84::KAN (Mata) was mated with the GFP-Heh2 variant expressing strains AS1 (GFP-HEH2-NAT, Matα), AS2 (GFP-HEH2(ΔNLS)-NAT, Matα), AS3 (GFP-HEH2(P2’H2NLS)-
NAT, Matα) and AS4 (GFP-NAT, Matα). Diploids were selected on YPD containing G418 and Nat. After sporulation, tetrads were dissected and the genotype of each spore was determined by replica plating on YPD containing Nat and YPD containing G418. All constructs generated in this study were fully sequenced to ensure the correctness of the DNA sequence.
SUPPLEMENTARY REFERENCES