Modular Organization and Combinatorial Energetics of Proline–Tyrosine Nuclear Localization Signals

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Proline–tyrosine nuclear localization signals (PY-NLSs) are recognized and transported into the nucleus by human Karyopherinβ proteins (Kapβs; Importins/Exportins) that mediate the majority of nucleocytoplasmic protein transport. There are 19 known Kapβs in human and 14 in yeast [1,2]. Kapβs bind substrates through nuclear localization or export signals (NLSs or NESs) and transport them through the nuclear pore complex, and Ran GTPase regulates Kapβ-substrate interactions [3–6]. Ten Kapβs are known to function in nuclear import, each recognizing at least one distinct NLS.

The best-known NLS is the short, basic, classical NLS, which is recognized by Kapβ4/Kapβ1 [4], and this pathway is conserved functionally from human to yeast [7,8]. Classical NLSs can be divided into monopartite and bipartite NLSs. Monopartite NLSs contain a single cluster of basic residues, whereas bipartite sequences contain two clusters of basic residues separated by a 10–12 amino acid linker. Thermodynamic dissection by scanning alanine mutagenesis of monopartite NLSs from the SV40 large T antigen (PKKKRKV) and the c-myc proto-oncogene (PAAKRVKLD) [9–11] confirmed a previously determined consensus sequence of K(K/R)X(K/R) [8,12]. Binding energies of these small signals are dominated by a single lysine residue, in the third position of the SV40 large T antigen and in the fourth position of c-myc, which makes numerous interactions with Kapβ [9]. Thus, in the monopartite classical NLS, it is well-known that a relatively small motif is recognized, and binding energy is concentrated in stereotypical fashion across small sequences. Although numerous structures are available for bipartite NLSs [13–15], thorough thermodynamic analysis of this subclass is not available, and its consensus is less well-defined (one example is KRX_{10–12}KRRK) than that for the monopartite NLS.

Recently, structural and biochemical analyses of human Kapβ2 (Transportin) bound to the hnRNP A1 NLS revealed physical rules that describe Kapβ2’s recognition of a diverse set of 20–30-residue-long NLSs that we termed PY-NLSs [16]. These rules are structural disorder of a 30-residue or larger peptide segment, overall basic character, and weakly conserved sequence motifs composed of a loose N-terminal hydrophobic or basic motif and a C-terminal RX_{2–5}PY motif. The composition of the N-terminal motifs divides PY-NLSs into hydrophobic and basic subclasses (hPY- and bPY-NLSs). The former contains four consecutive predominantly hydro-

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Introduction

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Recently, structural and biochemical analyses of human Kapβ2 (Transportin) bound to the hnRNP A1 NLS revealed physical rules that describe Kapβ2’s recognition of a diverse set of 20–30-residue-long NLSs that we termed PY-NLSs [16]. These rules are structural disorder of a 30-residue or larger peptide segment, overall basic character, and weakly conserved sequence motifs composed of a loose N-terminal hydrophobic or basic motif and a C-terminal RX_{2–5}PY motif. The composition of the N-terminal motifs divides PY-NLSs into hydrophobic and basic subclasses (hPY- and bPY-NLSs). The former contains four consecutive predominantly hydro-
Author Summary

To travel between the cytoplasm and nucleus, proteins rely on a family of transport proteins known as the karyopherinβ family. Karyopherinβ2, the human version of a family member, recognizes cargo proteins containing a class of nuclear localization signal known as the PY-NLS. The yeast homolog of Karyopherinβ2, Kap104p, also recognizes PY-NLSs, indicating that this pathway has evolved to transport proteins between evolutionarily distant species. We mutated residues in the PY-NLSs of two Kap104p cargo proteins and analyzed how tightly these mutants bound Kap104p. These experiments revealed three PY-NLS regions, or epitopes, that are important for binding Kap104p. Each epitope is composed of amino acids that vary between cargoes. The epitopes are energetically independent and bind Kap104p with varying strengths in different PY-NLSs, such that mutating the epitope of one PY-NLS may mistakenly direct cargo to the cytoplasm, while a similar mutation in a different PY-NLS has little effect on cargo localization. This flexible, energetically modular, and combinatorial architecture of PY-NLSs may confer higher tolerance to mutations, but it also allows greater sequence diversity, making prediction of new PY-NLSs difficult. The characteristics of PY-NLSs reported here will assist in the identification of new Kap104p cargoes. And the approach used may be applicable to other biological recognition pathways.

Table 1. Summary of Validated and Potential PY-NLSs

| Description | Proteins | PY-NLSs | References |
|-------------|----------|---------|------------|
| Human, validated in vivo and in vitro<sup>a</sup><sup>e</sup> | PQBP-1<sup>b</sup> | 161 RDRGYDKACSDREKGERRHKREELAPY 187 | [16], [19]<sup>a</sup>, [88]<sup>e</sup> |
| | YBP1<sup>b</sup><sup>c</sup> | 176 SAPEGOAQKRRPRKYRRFPPPYPYRIPY 196 | [16], [19], [69]<sup>a</sup> |
| | PAPB2<sup>d</sup> | 280 FYSGNSPRPRGVRGVRATSWRTSPY 306 | [70]<sup>c</sup> |
| | EWS<sup>hx</sup> | 630 GGGGRRGRRGGPKMRKREDHRRDDPY 656 | [16], [19], [71]<sup>d</sup> |
| | FUS<sup>ld</sup> | 500 GDDGGFPGKPDVQSRGMHRRRREPPPY 526 | [16], [19], [71]<sup>d</sup> |
| | TAF1<sup>ld</sup> | 566 YGGQLGGTGKKRRGGGNKNTDRRNPPY 592 | [71]<sup>c</sup> |
| | SAM68<sup>hx</sup> | 414 DOWNRTPSLKKAPPAPLQVGAAYRIPY 440 | [16], [19], [55]<sup>c</sup>, [72]<sup>d</sup> |
| | ETL2<sup>b</sup> | 317 QQEDWYNQSYKHAPTSARAYRIPY 343 | [71]<sup>c</sup> |
| | hnRNP M<sup>b</sup> | 38 GEGPRAQNKREKRKNGRGKNNTIPY 54 | [16], [19], [73]<sup>e</sup> |
| | hnRNP A1<sup>b</sup> | 263 FNGYNQSSNFPMKGGGGRGGGSPY 289 | [17], [21], [24], [25]<sup>e</sup> |
| | hnRNP A2<sup>b</sup> | 293 NYQNSPSNYPMKGGGSSNNMGPPY 319 | [24]<sup>e</sup> |
| | hnRNP A3<sup>b</sup> | 334 NYSGQSSQNYPMKGGGSGSSPGSPY 360 | [74]<sup>e</sup> |
| | hnRNP A0<sup>b</sup> | 264 YPMKGGGGGSGGGGGSNGSPY 290 | [16], [23], [27]<sup>e</sup> |
| | hnRNP D<sup>b</sup> | 329 YGYSQNGQGVRGRGNGSHGPPPY 355 | [22], [77]<sup>e</sup> |
| | JKBTP-1<sup>b</sup> | 394 ADYSGQOSTYKASGRGGGRQNNIPY 420 | [20]<sup>e</sup> |
| | TAP<sup>b</sup> | 49 SSRLLEDGDOVAMSODAQDPQVRNYPY 75 | [23, 75, 76]<sup>e</sup> |
| Human, potential PY-NLSs validated in vitro<sup>d</sup><sup>e</sup> | HMBA-ind<sup>ld</sup><sup>c</sup><sup>e</sup><sup>g</sup> | 141 WGGQQRQLGKKHRMPSSKKRRH-WKIPY 167 | [16], [19]<sup>c</sup>, [77]<sup>e</sup> |
| | RB158<sup>b</sup> | 258 GLPKPWEARRRSSCSDDGRTWWIPY 284 | [16]<sup>e</sup> |
| | HCC1<sup>cd</sup> | 73 RSKKERRRSSRASSDARKRVRIPY 99 | [16]<sup>e</sup> |
| | SOX14<sup>d</sup> | 72 HPPAYRPBPKPPKKLKKQKVRIPY 98 | [16]<sup>e</sup> |
| | Clik<sup>ld</sup> | 72 RSPFSGEDYYKRSSRRRARLG101 | [16]<sup>e</sup> |
| | hnRNP F<sup>b</sup> | 184 FKSQEEVSYDSPPLKMFVQRIPY 210 | [16], [19], [78]<sup>e</sup> |
| | hnRNP H1<sup>e</sup> | 184 FKSRAEVSYDSPPLKMFVQRIPY 210 | [19]<sup>e</sup> |
| | CPSF6<sup>cd</sup> | 364 PPTNGMPSRDSDGGPPTDPYPRPY 390 | [16], [19]<sup>e</sup> |
| | Cyclin T1<sup>hg</sup> | 661 VNHLHSLLAQQPGVTQTAFEPY 687 | [16], [19]<sup>e</sup> |
| | WBS16C<sup>b</sup> | 75 PSFVQSPSSQPPACARRRRQIPY 301 | [16]<sup>e</sup> |
| Yeast, validated in vivo and in vitro<sup>d</sup><sup>a</sup><sup>e</sup> | Hsp101<sup>b</sup> | 506 RSGGNHRNRRGGRGQRTRNRNYIPPY 532 | [34–37]<sup>b</sup><sup>d</sup><sup>e</sup> |
| | Nab2<sup>h</sup> | 214 AVGKNNRRGGRGRRRGGNHR1NFLP 241 | [34–37]<sup>b</sup><sup>d</sup><sup>e</sup> |

<sup>a</sup>Basic motifs are underlined, italicized, and in bold font. Hydrophobic motifs are underlined and in bold font. R/K/H and PY/L of the R/K/H<sub>2</sub>-PY/L motif are underlined and in bold font.

<sup>b</sup>Proteins were identified experimentally as Kap12 or Kap104p substrates.

<sup>c</sup>Human, validated in vivo and in vitro (includes targeting reporter to nucleus and subcellular mislocalization upon NLS mutation).

<sup>d</sup>W/2 substrates [16–25]. Table 1 summarizes previously reported validated and potential PY-NLSs. Although many of these potential substrates were predicted by bioinformatics [16] and still need experimental testing, more than 20 have been validated for Kap12 binding (Table 1) [16–25]. Comparison of in vivo and in vitro validated PY-NLSs shows large sequence diversity, which is reflected in weak consensus sequences [16]. Structures of five different Kap12-bound PY-NLSs also show substantial variability, with structurally diverse linkers separating the convergent consensus regions [16,26,27]. The PY-NLS is significantly larger than the short monopartite classical NLS. The well-defined consensus and concentrated binding energy of the latter may reflect compactness of the signal. In contrast, the binding energy of the PY-NLS is spread over a much larger sequence. Physical properties of the multipartite PY-NLS may be more similar to those of the less-studied, larger, and sequentially more diverse bipartite classical NLS.

Diverse PY-NLSs are described necessarily by weak consensus motifs. Therefore, instead of the traditional way of describing a linear recognition motif with a strongly restrictive consensus sequence, PY-NLSs were described by a collection of individual

<sup>e</sup>Yeast, validated in vivo and in vitro (includes targeting reporter to nucleus and subcellular mislocalization upon NLS mutation).
usually weak physical rules that together were able to provide substantial limits in sequence space for reasonable predictions of new Kapβ2 substrates [16]. However, the currently predicted substrates are most likely only a fraction of all PY-NLS-containing proteins because narrow sequence patterns were used in the initial search to achieve optimal accuracy. In fact, the sequence patterns used [16] were too narrow to predict PY-NLSs in known substrates HuR, TAP, hnRNP F, and JKTBP-1. The coverage of conventional sequence-based bioinformatics searches is expected to be severely limited due to PY-NLS diversity. Although sequence patterns obviously need to be expanded, we do not yet understand the limits of sequence diversity within motifs or how the different motifs may be combined. Knowledge of how binding energy is parsed in PY-NLSs will shape future efforts to decode these highly degenerate signals. Furthermore, physical understanding of how diverse PY-NLS sequences can achieve common biological function also will provide unique insights into many biological recognition processes that involve linear recognition motifs with weak and obscure consensus sequences, such as vesicular cargo sorting and protein targeting to the mitochondria and the peroxisome [28–33].

The yeast homolog of Kapβ2 is Kap104p (32% sequence identity) [34]. Only two Kap104p substrates, the mRNA processing proteins Nab2p and Hrp1p, are known. Several groups have mapped and validated NLSs of these substrates using both in vivo and in vitro methods to arginine–glycine (RG)-rich regions that were termed rg-NLSs [35–37]. Little sequence homology was detected between NLSs recognized by Kapβ2 and Kap104p. Furthermore, substrate recognition by the two karyopherins appears nonanalogous, as Kap104p does not recognize human substrate hnRNP A1 [35,37]. Given the recent physical understanding of Kapβ2–NLS interactions, we seek to examine the evolutionary conservation and energetic organization of signals in this pathway through studies of Kap104p–NLS interactions.

First, we present biochemical and biophysical analyses showing that RG-rich substrates of yeast Kap104p share similar physical characteristics to those of human PY-NLSs. Kap104p recognizes the basic but not hydrophobic PY-NLS subclass, and structural analyses of Kapβ2–NLS complexes suggested the origin of this specificity, enabling prediction of PY-NLS subclass specificity for all eukaryotic Kapβ2s. Thermodynamic analyses of Kap104p–NLS interactions revealed biophysical properties that govern binding affinity of PY-NLSs. These signals contain at least three energetically significant binding epitopes that are also linear motifs. Each linear epitope accommodates significant sequence diversity, and we have characterized some of the limits of this diversity. The linear epitopes are also energetically quasi-independent, a property that is probably due to intrinsic disorder of the free signals. Finally, in different PY-NLSs, a given epitope can vary significantly in its contribution to total binding energy. When combined with multivalency, this energetic variability can amplify signal diversity through combinatorial mixing of energetically weak and strong motifs.

**Results**

**Yeast rg-NLSs Are Also PY-NLSs**

In vivo validated RG-rich NLSs of Hrp1p and Nab2p (or rg-NLSs) are located at residues 494–534 and 201–250, respectively (Figure 1A) [35–38]. Examination of their sequences revealed physical characteristics similar to those of human PY-NLSs. Hrp1p and Nab2p NLSs are located within structurally disordered segments of 120–190 residues (DisEMBL, structural disorder probabilities of 0.72 and 0.63 for Hrp1p and Nab2p, respectively [39]) in the full-length proteins (Figure 1A). 506 RSQGNAHRNGRRGGR of Hrp1p and 529 KNNRRGCGGGNRRGGR of Nab2p contain many basic residues, like basic N-terminal motifs in human Kapβ2 substrates hnRNP M, PQQBP-1, and YB-1 (Table 1) [16]. Closer to the C termini, the Hrp1p 526 RNNGYPHY of and the Nab2p 253 RFPNL segments either match or are homologous to the C-terminal RX2–3PY consensus.

Immobilized full-length Hrp1p, Nab2p, and their NLSs bound Kap104p in stoichiometric proportions in pull-down binding assays (Figure 1B). Although it was previously reported that Ran could not dissociate substrate from Kap104p [36], we observed efficient dissociation of both full-length substrates and NLSs by RanGTP, possibly due to higher activity and GTP loading of the recombinant Ran. Our results suggest that Kap104p–NLS interactions and regulation by Ran are similar to other characterized Kapβ-mediated nuclear import processes in human [3–6]. Thermodynamic parameters for Kap104p binding to Hrp1p and Nab2p NLSs were obtained by isothermal titration calorimetry (ITC) (Figure S1). Both NLSs bound Kap104p with high affinity (K_D of 32 nM for Hrp1p and 37 nM for Nab2p (Tables 2 and 3)), and extensive mutagenesis of NLSs is discussed below. Thus, on the basis of their sequence characteristics, high affinity for karyopherin, and dissociation by RanGTP, yeast NLSs recognized by Kap104p resemble PY-NLSs.

**Kap104p Recognizes the Basic but Not Hydrophobic Subclass of PY-NLSs**

To investigate the PY-NLS subclass specificity of Kap104p, we examined its interaction with several human hPY- and bPY-NLSs as well as several predicted (see below) yeast hPY- and bPY-NLSs. Splicing factor hnRNP A1 and mRNA transport factor TAP/NXF1 contain hPY-NLSs, and splicing factor hnRNP M and FUS contain bPY-NLSs (Figure 2A). All four human PY-NLSs interacted with Kapβ2 [16], but only bPY-NLSs from hnRNP M and FUS bound yeast Kap104p in GST pull-down assays (Figure 2B). Both yeast Hrp1p and Nab2p NLSs bound equally well to Kap104p and Kapβ2 (Figure S2). Hrp1p and Nab2p are the only two known Kap104p substrates [34–36]. We needed to identify additional yeast substrates to test the preference of Kap104p for bPY-NLS. Because Nab2p has a C-terminal PL instead of PY motif, suggesting that PL motifs also may be present in other functional PY-NLSs, we used the program ScanProsite [40] and sequence patterns Φ1-G/A-S-Φ2-R/K/H-X2–5-PY/L (where Φ1 is a hydrophobic residue and Φ2 or Φ1 are hydrophobic residues or R or K) [16] to search for potential hPY-NLSs within Saccharomyces cerevisiae proteins in the UniProtKB/UniProt protein database [41]. A consensus sequence for the N-terminal motif of bPY-NLSs is not available due to lack of an apparent specific pattern. As a result, we modified a previously used sequence pattern that is consistent with the basic motifs of hnRNP M and PQQBP-1 [16] to accommodate additional validated human bPY-NLSs and NLSs in Nab2p and Hrp1p (Table 1). The resulting sequence pattern K/R-X0–6–4-K/R-X0–6–4-K/R-X2–5–R/K/H-X1–5–PY/L is used to search for
potential yeast bPY-NLSs. The resulting lists were filtered for structural disorder [39] and overall basic character. Six hPY/L-containing fragments were tested, but none bound Kap104p (Figure 2A and 2C). However, 11 of 20 bPY/L-containing fragments tested bound Kap104p and were dissociated by RanGTP (Figure 2A and 2D and Figure S3a and S3b). Two bPY/L-containing full-length substrates, Tfg2p and Rml2p, were tested, and both bound Kap104p and were dissociated by RanGTP.

Table 2. Summary of ITC Data for Kap104p Binding to Hrp1p Mutants

| Hrp1p Mutations | K_d (nM) | ΔG° (kcal/mol) | ΔH (kcal/mol) | TΔS° (kcal/mol/K) | ΔΔG° (kcal/mol) | K_d,wild type/K_d,mutant |
|------------------|----------|----------------|---------------|------------------|----------------|------------------------|
| Wild type        | 32 ± 16  | −10.1 ± 0.3    | −10.0 ± 0.5   | 0.09 ± 0.7       | —              | —                      |
| Mutations in the N-terminal motif |          |                |               |                  |                |                        |
| 506 RSGG/AAA     | 42 ± 35  | −10.0 ± 0.6    | −12.6 ± 1.2   | 0.02 ± 0.6       | 1.3 ± 1.1      |                        |
| 512 RGN/AAA      | 48 ± 26  | −9.8 ± 0.3     | −8.3 ± 0.5    | 0.2 ± 0.3        | 1.5 ± 0.1      |                        |
| 516 RGG/AAA      | 24 ± 18  | −10.3 ± 0.5    | −8.3 ± 2.0    | 2.0 ± 2.4        | −0.3 ± 0.5     | 0.7 ± 0.6              |
| 519 RGGYN/AAAA   | 17 ± 6   | −10.4 ± 0.2    | −14.2 ± 1.9   | −3.8 ± 2.1       | −0.4 ± 0.2     | 0.5 ± 0.2              |
| R512A,R513A,R516A,R519A | 194       | −10.0 ± 0.2    | −12.8 ± 0.0   | −2.8 ± 0.1       | −0.05 ± 0.2    | 1.1 ± 0.4              |
| R512K,R513K,R516K,R519K | 13       | −10.0 ± 0.2    | −12.8 ± 0.0   | −2.8 ± 0.1       | −0.05 ± 0.2    | 1.1 ± 0.4              |
| Mutations in the C-terminal RX2–5PY motif |          |                |               |                  |                |                        |
| R524A            | 37 ± 4   | −10.0 ± 0.05   | −11.7 ± 1.5   | 0.07 ± 0.06      | 1.1 ± 0.1      |                        |
| R525A            | 13 ± 3   | −10.5 ± 0.1    | −12.7 ± 0.6   | −0.5 ± 0.1       | 0.4 ± 0.09     |                        |
| 531 RY/AA        | 62 ± 11  | −9.6 ± 0.1     | −11.4 ± 0.2   | 0.4 ± 0.1        | 1.0 ± 0.3      |                        |
| Y529A            | 143 ± 46 | −9.2 ± 0.2     | −10.9 ± 1.3   | 0.8 ± 0.2        | 4.4 ± 1.4      |                        |
| Y529L            | 37 ± 27  | −10.0 ± 0.4    | −10.9 ± 0.8   | −0.9 ± 0.4       | −0.02 ± 0.4    | 1.1 ± 0.8              |
| Mutations in two binding epitopes |          |                |               |                  |                |                        |
| R512A,524R/AA    | 44 ± 28  | −9.9 ± 0.4     | −9.9 ± 0.4    | 0.06 ± 0.02      | 0.3 ± 0.4      | 1.4 ± 0.9              |
| R512A,Y529A      | 60 ± 6   | −9.7 ± 0.05    | −11.9 ± 0.4   | −2.2 ± 0.5       | 0.4 ± 0.05     | 1.9 ± 0.2              |

a Stoichiometry = 0.9–1.1.
b ΔG = −RT ln K_d.
c TΔS = (ΔH − ΔG).
d ΔΔG = −RT ln(K_d,wild type/K_d,mutant).

n.d., not detectable; All binding experiments were performed 2–4 times (± standard deviation).
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Figure 1. Interactions between Kap104p and Import Substrates Hrp1p and Nab2p
(A) Domain organization of Hrp1p and Nab2p. Domains are indicated by gray boxes, and the RGG regions and the glutamine-rich region are labeled. The sequences of the two NLSs are shown, with the basic motif highlighted in black and the RX2–5PY(L) motif in bold and underlined.

(B) Binding assays of Kap104p (arrow) with immobilized full-length Hrp1p and Nab2p (asterisks) or Hrp1p and Nab2p NLSs, in the presence and the absence of RanGTP. Bound proteins are stained with Coomassie blue.
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Mutations in the C-terminal binding epitopes R222A, 238PL239/AA 411 (H12B–H14B) (Figure 3A). Here, Kap phobic N-terminal motifs (H15B–H17B) and at linker regions the structurally overlapping interfaces with the basic/hydrophobic motif (H8B–H10B) are mostly invariant, while differences occur at residue, and (3) the PY residues, both of the C-terminal binding sites: (1) overlapping portions of the N-terminal hydrophobic motif interfaces in different organisms, and used this information to predict species in which Kap104p would lose binding affinity for additional hydrophobic motifs. Thus, recognition of the basic subclass of PY-NLSs is conserved between human and yeast. However, human Kap104p has evolved to recognize an additional hydrophobic motif in the context of crystal structures of Kap104p bound to PY-NLSs.

Kap104p Subclass Specificity

Kap104p and Kap110p sequences were aligned and examined in the context of crystal structures of Kap104p bound to PY-NLSs of hnRNP A1 (hPY-NLS) and M (bPY-NLS) [16,26]. Kap104p has 20 HEAT repeats, each consisting of two antiparallel B helices A and B. Both PY-NLSs bind the Kap104p interface lined by helices of HEAT repeats 8–18 (abbreviated H8B–H18B), converging structurally at three spatially distinct binding sites: (1) overlapping portions of the N-terminal hydrophobic and the larger basic motifs, (2) the arginine residue, and (3) the PY residues, both of the C-terminal RX2–5PY motifs [26]. Correspondingly, both structures share many common Kap104p interface residues, especially those that contact the conserved C-terminal RX2–5PY motif (Figure 3A).

Approximately half of the Kap104p interface residues are conserved in Kap110p. Interfaces with the RX2–5PY motifs (H8B–H12B) are mostly invariant, while differences occur at the structurally overlapping interfaces with the basic/hydrophobic N-terminal motifs (H15B–H17B) and at linker regions (H12B–H14B) (Figure 3A). Here, Kap104p residues I722, S729, N726, E734, T766, and I773 that contact the hnRNP A1 hydrophobic motif are replaced with T, P, I, L, S, and V, respectively, in yeast (Figure 3A and B) such that many hydrophobic contacts with the FGPM N-terminal motif of hnRNP A1 are expected to be lost in yeast (detailed description in Text S1). In contrast, among Kap104p residues that contact basic side chains of bPY-NLSs, only E653 of Kap104p is different in yeast (Figure 3A and 3C), and several amino acids have been replaced by more electronegative amino acids in Kap104p (Figure 3C), further supporting bPY-NLS recognition in yeast.

Comparison of individual HEAT repeats of Kap104p and Kap110p showed high identity (~50%) at H8–H10, but the similarity dropped to ~20% at H17 (Figure 3D). The B helices that line the interface are generally more conserved than the outer A helices. However, even in the former, sequence identities in H16B–H17B dipped significantly below 40% (Figure 3D). These observations suggest that both helical orientations and interface functional groups are better conserved at recognition sites for the C-terminal PY motif (H8–H10) than at the N-terminal basic/hydrophobic motifs (H16–H17). Consequently, the loss of Kap104p recognition for the N-terminal hydrophobic motif is most likely due to critical interface residue changes in H16B–H17B and to changes in helical orientations in this region. We have aligned sequences of Kap104p homologs, tracked interface residues and potential overall helical similarities at the N-terminal hydrophobic motif interfaces in different organisms, and used this information to predict species in which Kap104p would recognize hPY-NLSs. Results of these studies are discussed in Text S2 and shown in Figure S4A and S4B.

Distribution of Binding Energy along the Hrp1p NLS

We have performed scanning alanine mutagenesis covering residues 506–532 of the Hrp1p NLS (Figure 1A, Table 2, and Table S1). In the N-terminal region of the Hrp1p NLS, none of the four mutants 506RSGG229/AAAA, 512RRNG253/AAAA,
(R512, R513, R516, and R519) were mutated to alanines. This quadruple mutant decreased Kap104p binding by a marginal 5-fold (Figure 4A and Table 2), suggesting that positive charges in the N-terminal basic region are somewhat important for Kapβ-NLS interaction. Quadruple mutant R512, R513, R516, R519/KKKK did not affect Kap104p binding (Table 2), further suggesting that stereospecific interactions with arginine guanido groups are not important for Kap104p binding.

Kap104p binding was not affected significantly when both arginine residues, 524RR, in the C-terminal RX2–5PY motif of the Hrp1p NLS were mutated to alanines (K_{D,mutant}/K_{D,wild type} = 1.7; Figure 4A and Table 2). In contrast, the C-terminal 531PY/AA mutation abolished detectable Kap104p binding (Table 2). The enthalpies of binding for all of the PY-NLSs that we have measured by ITC are similar, and the weakest measurable K_D in this series was 10 μM [26]. Therefore, we assume that the affinity of the Hrp1p 531PY/AA mutant is likely weaker than 10 μM and its K_{D,mutant}/K_{D,wild type} > 200 (Figure 4A). Thus, the Hrp1p NLS contains one strong binding hotspot at its PY motif, similar to the single significant hotspot at the C-terminal PY motif of the human substrate hnRNPM (K_{D,mutant/AA}/K_{D,wild type} = 500 for the hnRNPM NLS) [26]. Interestingly, we also located a modest binding hotspot at residue Y529 (K_{D,mutant}/K_{D,wild type} = 4 for Y529A; Figure 4A and Table 2) in the linker between the arginine and the PY of the RX2–3PY motif. However, the Y529L mutation did not affect Kap104p binding (Table 2), suggesting that a hydrophobic, but not necessarily aromatic, moiety at this position might be important.

Distribution of Binding Energy along the Nab2p NLS

We have performed scanning alanine mutagenesis covering residues 210–239 of the Nab2p NLS (Figure 1A, Table 3, and Table S1). Binding energy along the Nab2p NLS appears quite distributed compared to that of the Hrp1p NLS, with no single binding hotspot that stands out above others (Figure 4B and Table 3). In its basic N-terminal region, 216KNRR219, 222RGGR224, 226RGGRN230 each were mutated to alanines, but only 216KNRR219/AAAA showed a small 3-fold decrease in Kap104p affinity (Table 3). None of the single mutants K216A, R218A, R219A, R222A, R226A, or R229A decreased Kap104p binding (Table S1), and simultaneous mutation of all five arginines to alanines (K_{D,mutant}/K_{D,wild type} = 2.25 μM; Figure 4B and Table 3), suggesting that the collective basic character of this region contributes significantly to the total binding energy of the NLS. Comparison of single arginine to alanine mutants (K_{D,mutant}/K_{D,wild type} = 1.0) to the pentamutant R218, R219, R222, R226, R229/AAAAA (K_{D,mutant}/K_{D,wild type} = 60.8) indicated a binding cooperativity of at least 60-fold within the N-terminal basic motif of Nab2p.

When R235 of the Nab2p C-terminal RX2–5PL motif was mutated to an alanine, Kap104p affinity decreased by 5-fold (Figure 4B and Table 3). Crystal structures of Kapβ2 bound to NLSs of hnRNPs A1 and M showed the equivalent arginine residues making electrostatic interactions with numerous aspartate and glutamate residues, suggesting the importance of a positively charged residue at this position [16,26]. We also mutated R235 to lysine and histidine, but neither mutant

**Figure 2. Kap104 Recognizes Basic PY-NLSs but Not Hydrophobic PY-NLSs**

(A) Sequences of known PY-NLSs in human proteins and predicted hydrophobic-PY(Y) (shaded in light gray) and basic PY motifs (shaded in dark gray) in yeast proteins. The RX2–5PY(L) motif is underlined.

(B) Binding assays of Kap104p and immobilized human bPY-NLSs (hnRNPM and FUS) or human hPY-NLSs (hnRNPA1 and TAP).

(C) Binding assays of Kap104p with six immobilized predicted yeast hPY(L)-NLSs. GST and GST–Nab2p are included as controls. Faint bands at residues 210–239 of the Nab2p NLS (Figure 1A, Table 3, and Table S1) are likely heat shock protein contaminants.

(D) Binding assays of Kap104p and immobilized predicted bPY-NLSs or full-length proteins. GST–Hrp1p is included as a control. Bound proteins are stained with Coomassie blue.

**Table 3.** Distribution of Binding Energy along the Nab2p NLS

| Binding Energy | Nab2p NLS Region | 210–239 | 210–233 | 210–219 | 214–239 | 214–222 | 226–239 | 226–233 | 226–219 | 216–239 | 216–222 | 222–239 | 222–226 | 226–233 | 226–219 | 222–239 | 222–233 | 222–226 | 226–233 |
|---------------|-----------------|---------|---------|---------|---------|---------|---------|---------|---------|---------|---------|---------|---------|---------|---------|---------|---------|---------|---------|
| K_{D} (μM)     |                 |         |         |         |         |         |         |         |         |         |         |         |         |         |         |         |         |         |         |
| 216KNRR219    | 2.25 ± 0.5      | 10.1 ± 1.1 | 20.2 ± 2.3 | 1.4 ± 0.3 | 1.0 ± 0.1 | 1.0 ± 0.1 | 1.0 ± 0.1 | 1.0 ± 0.1 | 1.0 ± 0.1 | 1.0 ± 0.1 | 1.0 ± 0.1 | 1.0 ± 0.1 | 1.0 ± 0.1 | 1.0 ± 0.1 | 1.0 ± 0.1 | 1.0 ± 0.1 | 1.0 ± 0.1 | 1.0 ± 0.1 | 1.0 ± 0.1 |
| 216KNRR219/AAAAA | 4.3 ± 0.7      | 4.3 ± 0.7 | 4.3 ± 0.7 | 4.3 ± 0.7 | 4.3 ± 0.7 | 4.3 ± 0.7 | 4.3 ± 0.7 | 4.3 ± 0.7 | 4.3 ± 0.7 | 4.3 ± 0.7 | 4.3 ± 0.7 | 4.3 ± 0.7 | 4.3 ± 0.7 | 4.3 ± 0.7 | 4.3 ± 0.7 | 4.3 ± 0.7 | 4.3 ± 0.7 | 4.3 ± 0.7 | 4.3 ± 0.7 | 4.3 ± 0.7 | 4.3 ± 0.7 |
| 216KNRR219/AAAA | 2.25 ± 0.5      | 2.25 ± 0.5 | 2.25 ± 0.5 | 2.25 ± 0.5 | 2.25 ± 0.5 | 2.25 ± 0.5 | 2.25 ± 0.5 | 2.25 ± 0.5 | 2.25 ± 0.5 | 2.25 ± 0.5 | 2.25 ± 0.5 | 2.25 ± 0.5 | 2.25 ± 0.5 | 2.25 ± 0.5 | 2.25 ± 0.5 | 2.25 ± 0.5 | 2.25 ± 0.5 | 2.25 ± 0.5 | 2.25 ± 0.5 | 2.25 ± 0.5 | 2.25 ± 0.5 |

514RGGR518/AAAA, and 514RGGRN525AAAAA (Table 2) affected Kap104p binding, suggesting that this N-terminal basic-enriched region may contribute little to total binding energy. However, these mutations may be misleading as glycine to alanine mutations may decrease the entropy of the unbound NLS, thus decreasing the entropic penalty of binding and offsetting affinity loss from arginine mutations. Therefore, we also generated a quadruple mutant where all of the arginines...
affected Kap104p binding significantly (KD,mutant/KD,wild type are 1.0 and 1.7, respectively; Table 3). The C-terminal 238PL239/AA mutation in the Nab2p NLS decreased Kap104p binding by 10-fold (Figure 4B and Table 3). The energetic significance of this mutation suggests its equivalence to the PY motif in human Kapb2 substrates and in Hrp1p. Furthermore, the Nab2p 238PL239/PY mutant bound Kap104p with a slightly higher affinity at a KD value of 13 nM. Mutagenesis of residue L239 to all other amino acids is described below.

The measurable 238PL239/AA mutation in the Nab2p NLS (KD = 376 nM) provided an opportunity to explore cooperativity across binding sites or epitopes. Mutations in the Nab2p triple mutant R222A, 238PL239/AA (KD = 411 nM; KD,mutant/KD,wild type = 11.1; Table 3) show almost perfect additivity when compared to a single R222A mutant (did not affect Kap104p binding; Table S1) and double 238PL239/AA mutant (KD,mutant/KD,wild type = 10.2; Table 3). Strict additivity between the R and the PL sites would give a calculated KD,mutant/KD,wild type value of 56.1 for the triple mutant. Thus, the experimental KD,mutant/KD,wild type value of 14.7 for the triple mutant indicated 3.8-fold cooperativity between the two epitopes. Similarly, Hrp1p triple mutant R512A, 524RR525/AA and double mutant R512A, Y529A showed cooperativity of approximately 1.4- and 2-fold between epitopes, respectively. The couplings between binding epitopes observed here for both Nab2p and Hrp1p are still more than an order of magnitude lower than that observed within the N-terminal basic region of Nab2p (>60-fold cooperativity).

We also located a new binding hotspot at F236 in Nab2p (KD,mutant/KD,wild type = 8 for F236A; Figure 4B and Table 3), which is located in the linker between the R and the PL of the RX2–5PL C-terminal motif. This site is analogous to Y529 of Hrp1p discussed in the previous section, and both residues are located two residues N-terminal of the PY/L motifs. As in

![Figure 3. Kapb2–NLS Structures and Kap104p Specificity](http://www.plosbiology.org/article?id=10.1371/journal.pbio.0060137)
the Hrp1p NLS Y529L mutant, the F236L mutation in Nab2p did not affect Kap104p binding (Table 3). Aromatic or hydrophobic residues occur at this position in many human PY-NLSs, including hnRNPs M, D, and F, JKTBP, TAP, HMBA-inducible protein, PABP2, PQBP-1, RB15B, and WBS-16 [16,22,23,27]. Aromatic side chains at this position overlap in the crystal structures of Kap\textsubscript{2b} bound to the NLSs of hnRNPs M and D and TAP [26,27]. The F61 of the hnRNP M NLS, Y352 of the hnRNP D NLS, and Y72 of the TAP NLS make hydrophobic interactions with Kap\textsubscript{2b} W460A and with the backbones of the PY motifs. A hydrophobic residue here may contribute to binding energy through both favorable enthalpy and a decrease of entropic penalty upon binding by preorganizing the PY motif. Thus, if present, a hydrophobic residue here may be considered as an extension of the PY motif.

Hrp1p contains a single very significant binding hotspot at its PY motif. In contrast, binding energy in Nab2p is more evenly distributed across its N-terminal basic region and the R, F, and PL residues of its C-terminal consensus motif. Thus, distributions of binding energy in the two yeast NLSs are very different. From the N to C terminus, energetic distribution across the three epitopes (N-terminal basic/hydrophobic and the C-terminal PY motifs) of Hrp1p and Nab2p can be described roughly as medium–weak–strong and strong–medium–medium, respectively (Figure 4C–G). In summary, all three PY-NLS epitopes are energetically highly variable, the N-terminal basic/hydrophobic and the C-terminal PY motifs appear to cover the entire energetic continuum from strong to weak, and the arginine of the RX\textsubscript{2–5}PY motif is medium to weakly energetically significant.

**Degeneracy of Tyrosine in the C-Terminal PY Motif**

Of the more than 20 sequences that bind Kap104p and Kap104p (Table 1) [16], two do not contain the PY dipeptide in their C termini. HuR has a PG, and Nab2p has a PL, thus raising the question of degeneracy at this C-terminal position. We mutated Y532 in the PY motif of Hrp1p to the other 19 amino acids (Figure 5A and Table S2). Only Y532F, Y532H, and Y532M showed measurable Kap104p binding by ITC. Y532F best resembles the wild type, with only a 4-fold decrease in Kap104p affinity. Both Y532H and Y532M in Hrp1p bound significantly weaker with \( K_D \) values of 1 and 2 \( \mu \text{M} \), respectively.

We also mutated L239 in the Nab2p PL motif to the other 19 amino acids (Figure 5B and Table S3). Binding energy along the Nab2p NLS is distributed very evenly compared to that of the Hrp1p NLS with the Nab2p 239\textsubscript{PL}239/AA mutation as strong; Figure 4A and 4B). Similarly, in previously characterized PY-NLSs of hnRNPs A1 and D, TAP, and JKTBP [16,27], energetic distributions at the three epitopes also are quite varied, with rough patterns of strong–weak–weak, strong–medium–medium, weak–weak–weak, and weak–medium–strong, respectively (Figure 4C–G). In summary, all three PY-NLS epitopes are energetically highly variable, the N-terminal basic/hydrophobic and the C-terminal PY motifs appear to cover the entire energetic continuum from strong to weak, and the arginine of the RX\textsubscript{2–5}PY motif is medium to weakly energetically significant.
decreasing affinity only 10-fold compared to the >200-fold effect in Hrp1p. Thus, in the energetically distributed Nab2p NLS, changes in the E239 position may be quite permissive. This is indeed the case because only L239D and L239E showed significant affinity decreases of 11- and 7-fold, respectively. L239G, L239I, and L239P showed a modest 3–4-fold affinity decrease. None of the other mutants (to S, T, N, Q, K, R, V, M, F, Y, W, and H) decreased Kap104p binding.

Tyrosine is clearly the most preferred residue in the last position of the Hrp1p NLS. Correspondingly, mutation of the PL motif in Nab2p to PY improves Kap104p binding. These results suggest that, in general, tyrosine may be the most preferred and thus likely the most prevalent amino acid found in the last position of PY-NLSs. It appears that if the transport factor exhibits obvious biologically relevant specificity for nuclear import substrates but at the same time is able to handle a large number of different sequence-diverse substrates. Previous studies have captured the requirement for structural disorder in NLSs and the notion of a few anchoring amino acids such as the N-terminal hydrophobic/basic and RX2–5PY motifs [16,26]. Here, we show that yeast Kap104p is a PY-NLS-recognizing homolog specific for the basic subclass of this signal and that the two different Kap104p substrates have rather different distributions of binding energy for Kap104p. The NLS in Hrp1p largely uses the PY motif, and the NLS in Nab2p uses many positions distributed across three binding regions. Consistent with this, the Y position of the PY motif shows more degeneracy in Nab2p than in Hrp1p. On the basis of all of this and the thermodynamic data from five human PY-NLSs [16,26,27], we propose the following physical properties that govern the affinity of PY-NLS recognition by Kapβ2.

1. PY-NLSs Contain at Least Three Energetically Significant Binding Epitopes

Structures of PY-NLSs from hnRNPs A1, M, and D, TAP, and JKTBP converge spatially at three distinct binding sites or epitopes separated by structurally variable linkers: (1) the N-terminal hydrophobic/basic motif, (2) the arginine residue expressed GFP-tagged full-length Hrp1p and Nab2p wild-type and mutant proteins in yeast. Wild-type Hrp1p–GFP and Nab2p–GFP are localized in the nucleus as has been reported previously (Figure 6A–D) [34–36]. Mutations in the C-terminal PY motif (L238PLL/AA) of Hrp1p, which abolished detectable Kap104p binding, resulted in mislocalization of the GFP fusion protein to the cytoplasm (Figure 6A and 6C). The N-terminal basic motif of Hrp1p is also important for nuclear localization of Hrp1p: the R512,R513,R516,R519/AAAAA mutant, which decreased Kap104p binding by a marginal 5-fold, also is mislocalized (Figure 6A and 6C). Xu and Henry have shown previously that substitutions of R516 and R519 with glutamines mislocalized Hrp1p, but proteins with lysine substitutions are properly localized [38,42]. This further suggests that basic charges rather than stereospecific interactions are necessary for Kap104p interactions.

In the case of Nab2p, mutations in either the N-terminal motif (pentamutant R218,R219,R222,R226,R229/AAAAA; decreases Kap104p binding by 60-fold) or the C-terminal PY motif (L238PLL/AA; decreases Kap104p binding by 10-fold) resulted in increased cytoplasmic localization of the GFP fusion protein (Figure 6B and 6D). Arginine methylation of Nab2p by Hmt1p is required for its export from the nucleus, possibly explaining some nuclear accumulation of the N-terminal mutant despite its low affinity for Kap104p [38,43]. Combined mutations of both the N- and the C-terminal motifs resulted in diffuse localization of the fusion protein, consistent with further affinity reduction for Kap104p (Table 3 and Figure 6B and 6D). We have shown here that mutations in the PY-NLSs of Hrp1p and Nab2p that decrease binding affinity to Kap104 also affect nuclear localization in yeast cells.

Discussion

The problem of deciphering the sequence code for substrate recognition by Kapβ2 is interesting and challenging because the transport factor exhibits obvious biologically relevant specificity for nuclear import substrates but at the same time is able to handle a large number of different sequence-diverse substrates. Previous studies have captured the requirement for structural disorder in NLSs and the notion of a few anchoring amino acids such as the N-terminal hydrophobic/basic and RX2–5PY motifs [16,26]. Here, we show that yeast Kap104p is a PY-NLS-recognizing homolog specific for the basic subclass of this signal and that the two different Kap104p substrates have rather different distributions of binding energy for Kap104p. The NLS in Hrp1p largely uses the PY motif, and the NLS in Nab2p uses many positions distributed across three binding regions. Consistent with this, the Y position of the PY motif shows more degeneracy in Nab2p than in Hrp1p. On the basis of all of this and the thermodynamic data from five human PY-NLSs [16,26,27], we propose the following physical properties that govern the affinity of PY-NLS recognition by Kapβ2.
of the C-terminal RX$_2$–5PY sequence motif, and (3) the PY of the C-terminal RX$_2$–5PY motif [16,26,27]. We have shown here that all three structural epitopes can be energetically significant.

The N-terminal basic-enriched motifs of Hrp1p and Nab2p NLSs constitute epitope 1, where collective basic character and likely charge density drive Kap104p binding. Mutations of all of the arginines in this region to alanines decreased binding energy by 0.9–2.3 kcal/mol for both NLSs. Similarly, the N-terminal hydrophobic motif of the hnRNP A1 NLS and the equivalent region of the hnRNP D NLS that contains both hydrophobic and basic residues are also energetically significant, with mutations decreasing binding energy by $\sim$2 kcal/mol [26].

Epitopes 2 and 3 are contained within the C-terminal RX$_2$–5PY/L sequence motifs. Two linkers of variable lengths, compositions, and structures connect epitope 1 to epitope 2 and epitope 2 to epitope 3 [16,26]. Epitope 2 is located at Hrp1p R$_{521}$R$_{523}$ and Nab2p R$_{235}$ at the first consensus position of the C-terminal RX$_2$–5PY/L sequence motifs. Of the
three PY-NLS epitopes, epitope 2 tends to contribute the least to binding energy, with mutations decreasing binding energy maximally by \(\sim 1\) kcal/mol in Nab2p, hnRNP D, and JKTBP (Figure 4B, 4E, and 4G).

Epitope 3 is located at Hrp1p \(^{531}\)PY \(^{532}\) and Nab2p \(^{238}\)PL \(^{239}\). Mutations at these terminal positions are generally energetically significant, decreasing binding energy by 1.3–4 kcal/mol in Hrp1p, Nab2p, hnRNPs M and D, and JKTBP. However, exceptions are seen in hnRNPs A1 and TAP, where PY mutations decreased binding modestly by only \(\sim 0.7\) kcal/mol.

Because free PY-NLSs are structurally disordered and adopt extended Kap\(\beta\)-bound conformations, epitopes 1–3 are presented as peptides that can be represented by sequence patterns or linear motifs [44–46]. In epitope 1, the N-terminal basic motif may be represented by a collection of sequence patterns covering 5–19 residues, and the N-terminal hydrophobic motif by sequence patterns of approximately 4 residues. Epitopes 2 and 3 are both relatively smaller and simpler and together can be described by a single sequence pattern.

2. Each Linear Epitope Can Accommodate Large Sequence Diversity

Comparison of validated and potential PY-NLSs in Table 1 [16,26] show that sequences within each of the three linear epitopes can be quite variable. The N-terminal basic/hydrophobic motif is the largest and most variable epitope. Mutagenesis of yeast PY-NLSs has provided more information on the diversity and also suggested some limits to the diversity of individual epitopes. In particular, positive charges within the N-terminal basic motifs are important, but arginine and lysine residues are interchangeable, and the exact positions of basic groups may not be important (Tables 2 and 3 and Table S1). Additional biochemical and structural studies will be needed to understand requirements of charge density, segment size, and positively selected amino acids in this epitope. The consensus for this basic region remains elusive. The 55% accuracy for bioinformatics-derived potential yeast bPY-NLSs binding to Kap104p may reflect high sequence variability and undiscovered physical characteristics of this region.

Epitope 2 is usually composed of a single residue. Examination of validated PY-NLSs (Table 1) shows that arginine is most prevalent in this position, although histidines are found in this position in hnRNPs D, JKTBP, and HuR and lysines in potential yeast NLSs of Naf1p, Sbp1p, Arp8p, and Ste20p (Figure S3A). Mutagenesis has shown that arginine, lysine, and histidines are interchangeable in this position. Thus, the appropriate sequence pattern here is R/K/H.

Human Kap\(\beta\)2 substrate HuR (Table 1) has a PG dipeptide, and yeast Nab2p and eight bioinformatics-derived potential yeast NLSs contain PL dipeptides at the C-terminal positions of their NLSs (epitope 3). In some cases, epitope 3 matters energetically more than in others. It is unclear why the dipeptide motif is energetically significant in some peptides and relatively silent in others. We speculate that a hydrophobic amino acid two residues N-terminal of the PY motif may be necessary (though probably not sufficient) and should be included in the sequence pattern for an energetically strong epitope 3. A hydrophobic residue at this position may preorganize the short peptide segment for binding, lowering both strain and entropic penalties. We also note that if epitope 3 is energetically very significant, then the terminal site tends to be phenylalanine, histidine, and methionine. If the dipeptide motif is fairly silent energetically, then many other amino acids are allowed in the terminal position.

3. Energetic Cooperativity Observed within Linear Epitopes but Not between Them

Mutations within a linear epitope such as within the N-terminal basic region of Nab2p show large cooperativity of >60-fold (Table 2 and Table S1). Mutations within the N-terminal basic region of the hnRNPs M NLS also show cooperativity, in a similar regime, of \(\sim 40\)-fold [26]. In contrast, seven examples of simultaneous mutations between different linear epitopes in Hrp1p, Nab2p (Tables 2 and 3), and hnRNPs A1 and M [16,26] show only modest cooperativities of 1.0–3.8-fold. Cooperativity between linear epitopes in PY-NLSs is also very small compared to that typically observed between spatially distinct sites in conformational epitopes. For example, in the interaction of human growth hormone with human growth hormone receptor, mutations at distant sites in the interface showed large cooperativity of \(\sim 60\)-fold [47]. Thus, by comparison, the linear epitopes in PY-NLSs are energetically quasi-independent. In an analogous system, a bipartite interaction in a linear sorting signal in a SNARE and COPII coat also exhibited energetic quasi-independence, showing only a 1.5–2-fold cooperative effect between the two distant sites [48]. In both PY-NLSs and vesicular sorting signals, minimal coupling between linear epitopes, and thus energetic modularity of those epitopes, may be attributed to flexible or structurally variable linkers that connect the epitopes.

4. Energetically Variable Linear Epitopes Can Be Mixed in a Combinatorial Fashion

Finally, the fourth biophysical property that governs PY-NLS affinity stems from the observation that binding energy is distributed very differently amongst the three linear epitopes in all seven thermodynamically characterized PY-NLSs [16,26,27]. In different PY-NLSs, a given linear epitope can vary significantly in its contribution to total binding energy. For example, the N-terminal basic motif in Hrp1p contributes much less to Kap104p binding than the equivalent epitope in Nab2p (compare Figure 4A and 4B). Similarly, PY in hnRNPs A1 contributes only weakly to Kap\(\beta\)2 binding, while PY motifs in hnRNPs M and Hrp1p are the sole binding hotspots in the NLSs (Figure 4A, 4C, and 4D). We had previously taken advantage of the energetic variability of PY-NLS epitopes by harnessing the avidity effect of the NLS hotspot at epitope 1 of hnRNPs A1 fused to the NLS hotspot at epitope 3 of hnRNPs M, which resulted in a chimeric peptide inhibitor that bound Kap\(\beta\)2 200-fold tighter than both substrates and RanGTP [26]. Despite the wide energetic variability of individual linear epitopes, the total binding energies are very similar for various PY-NLS-containing substrates. Therefore, evolution has not combined epitopes randomly but rather tuned them to a range for appreciable Kap\(\beta\)2 binding and efficient Ran dissociation. The extremely tight-binding chimeric peptide inhibitor of Kap\(\beta\)2 [26] is evidence of such evolutionary pressure. Although very high affinity can be achieved easily, nuclear import function is lost as RanGTP can no longer dissociate substrates.

Binding energy in the PY-NLS is distributed over a large
sequence, with three different elements contributing differently in various substrates. It is this feature that makes the PY-NLS fundamentally different from the well-known monopartite classical NLS. A relatively small motif is recognized in a monopartite NLS, and binding energy is concentrated in a stereotypical fashion across small sequences.

**Modular and Combinatorial Design of PY-NLS May Be Highly Evolvable**

In PY-NLSs, the three distinct linear sequence elements are presented on peptides that exhibit intrinsic structural disorder and bind Kapβ2 with extended structurally diverse conformations. This modular and flexible display of multiple sequence motifs is relatively free of spatial constraints that usually relate multiple binding sites within a folded ligand. Furthermore, when binding energy is variably distributed among multiple epitopes in PY-NLSs, single mutations or mutations within single NLS epitopes are likely to have decreased chances of abolishing karyopherin binding. Thus, the modular, flexible, and energetically combinatorial architecture of PY-NLSs may allow significant evolvability to form new interactions while maintaining Kapβ2 recognition. Similar “multifaceted” interactions, where different ligands make energetically significant interactions with different subsets of interface residues, were recently studied in a theoretical context [49] and also suggested to be more tolerant to mutations and are therefore quite evolvable.

Multiple functions have been identified in fact in several PY-NLSs. In Nab2p, the RGG region that overlaps NLS epitope 1 is a putative RNA binding region [50]. The PY-NLSs in Nab2p, Hrp1p, EWS, and FUS interact with and are methylated by arginine methyltransferases [43,51–54]. Phosphorylation sites also have evolved within PY-NLSs to regulate nucleocytoplasmic localization. Serine phosphorylation in the hnRNP A2 NLS and tyrosine phosphorylation in the SAM68 NLS [55] both alter subcellular localization of the proteins. A PY-NLS also may evolve additional NLSs within its sequence. This could generate redundancy in nuclear import pathways and also provide a path to switch substrates from one karyopherin to another and ultimately from one cellular process to another. We have identified a potential classical NLS [56] in the N-terminal basic motifs of eight human bPY-NLSs in Table 1. It is not clear what overlapping NLSs mean in the cellular context, but this question will need to be explored in the future.

**Path to Comprehensive PY-NLS Identification in Genomes**

Identifying correct sequences that will account for most of the very diverse PY-NLS is an extremely challenging task. The core problem is that binding energy is distributed across three epitopes or motifs in many different ways. Thus, simply relaxing sequence constraints in a global search will also increase “noise” and result in many wrong answers.

We predict that if a PY motif (epitope 3) is energetically very significant, then the sequence tolerance for this motif is small, and sequence content of the other two epitopes will likely not matter. Thus, this subset of the PY-NLSs should be identified easily upon identification of PY motifs that can provide large binding energies. Given the relatively small size of this motif, the task of finding strong PY motifs should be experimentally accessible. A similar situation should apply for an energetically strong N-terminal basic/hydrophobic motif (epitope 1). However, as the need for affinity from the PY motif decreases and as more binding energy is provided by the two other motifs, sequence tolerance relaxes. The problem of multiple motifs with varying sequence tolerances seems very complex, but the relatively small size of each motif and energetic independence of the motifs allow the problem to be divided into manageable pieces. Our current inability to identify sequences of individual epitopes that are energetically strong may contribute to the 55% accuracy for bioinformatics-derived potential yeast bPYL-NLS binding to Kap104p. For example, individual epitopes in bioinformatics-derived sequences that did not bind Kap104p may be energetically weak and thus did not provide sufficient binding energy when combined.

First, the range of energies for PY-NLSs that are import-competent in vivo (and to what degree) will need to be determined. The range of suitable binding energies likely will vary depending on cellular concentrations of substrates but should not be unbounded [57]. For example, a designed peptide with a K_D of 100 pM binds Kapβ2 too tightly for in vivo nuclear import [26], thus providing a high-affinity boundary for Kapβ2 import. Second, binding energies of putative PY-NLSs will need to be predicted. Unfortunately, the accuracy of calculating binding affinity for protein–small molecule interaction is still questionable, and predictions of binding energies for protein–protein interactions are even further behind [58]. Our studies here suggest that we can get around this problem by handling each epitope independently and then combining them to assess for functional NLSs. We may use computational alanine-scanning mutagenesis [59] to predict binding energy differences for each of the three PY-NLS linear epitopes and then empirically determine combinations that are functional. Such predictions could be tested against a future experimental thermodynamic database obtained from the initial predicted PY-NLSs [16], and the method was refined iteratively. Binding energy calculation remains problematic. We expect that prevalent sequence- and physical-characteristics-based bioinformatics methods are limited to successful prediction of potential NLSs with at least one energetically strong linear epitope but will miss those composed of multiple weak or intermediate epitopes. A computational method that combines bioinformatics, structural modeling, and prediction of binding energies may be a solution. Many more Kapβ2-NLS structures will be necessary to expand a structural database to facilitate modeling interactions of new sequences by homology modeling and/or physical energy function-based predictions of protein–protein interactions [60–62].

**Conclusions**

PY-NLSs are very diverse in sequence and structure and thus cannot be described sufficiently by their weak consensus motifs. Instead, PY-NLSs are described by a collection of weak physical rules that also include requirements for intrinsic structural disorder and overall positive charge [16]. Here, we examined the energetic organization of PY-NLSs through mutagenic and thermodynamic analyses of these signals in yeast. These studies have revealed physical properties that govern the binding affinity of this variable signal. The PY-NLS is a modular signal composed of three spatially distinct but structurally conserved linear epitopes that can be represented by a series of sequence patterns. Although each
linear epitope can accommodate substantial sequence diversity, we have begun to define limits for each. More importantly, in addition to structural modularity, the three linear epitopes also exhibit energetic modularity. Modular organization of the PY-NLS suggests that the daunting search for these very diverse sequences can be performed in parts. Finally, each linear epitope can contribute very differently to total binding energy in different PY-NLSs, explaining how signal diversity can be achieved through combinatorial mixing of energetically weak and strong motifs while maintaining affinity appropriate for nuclear import function. This collection of physical rules and properties describes how functional determinants of the PY-NLSs are organized and lays a path to decode this diverse and evolvable signal for future genome-wide identification of Kapβ2 import substrates. More generally, many biological recognition processes involve linear recognition motifs with weak and obscure sequence motifs. Physical understanding of how diverse PY-NLS sequences can achieve common biological function may serve as a model for decoding many other weakly conserved and complex signals throughout biology.

Materials and Methods

Plasmds and strains. The Kap104p gene (gift from J. Aitchison) was subcloned into the pGEX-Tev vector [34]. Yeast substrate genes were obtained by PCR from a S. cerevisiae genomic DNA library (Novagen) and subcloned into the BamHI and NotI sites of the pGEX-Tev and/or pMAL-Tev vectors [63,64]. Site-directed mutagenesis of Nab2p 201–251 and Hrp1p 494–534 were performed using the QuikChange method (Stratagene) and confirmed by nucleotide sequencing.

Full-length Nab2p and Hrp1p wild-type and mutant genes were subcloned into the SpeI and Smal sites of a modified pRS415 (CEN6, ARS, LEU2, and AP) shuttle vector containing a C-terminal GFP gene [65].

Cell culture and microscopy. BY4741 (MATa his3Δ1 leu2Δ2 met15Δ0 ura3Δ0) cells harboring pRS415 plasmids were grown at 30 °C in SC-Leu media to mid-logarithmic phase [66]. Cells were transferred to a 1.5% low-melting-point agarose pad made with SC-leu in a coverslip bottom Wilko dish. Cells were observed on an Olympus IX-81 inverse microscope (60× objective) and images were acquired with a Hamamatsu ORCA-ER camera. All images were analyzed in Image-Pro Plus software (Media Cybernetics). To obtain the N/C ratio, mean fluorescence intensity in a 36-pixel box was measured in the nucleus and cytoplasm for at least 50 cells of each mutant.

Protein expression and purification. The GST–Kap104p protein was expressed in Escherichia coli Rosetta(DE3)pLysS cells (Novagen). Cells were lysed using an EmulsiFlex-C5 homogenizer (Avestin). The supernatant was applied to glutathione sepharose (GE Healthcare). The protein was washed with TB buffer and then eluted with Tris buffer containing 20 mM glutathione, pH 8.1. The GST tag was cleaved using 0.5 ml of TEV protease in a total volume of 10 ml and washed extensively with Tris buffer (50 mM Tris, pH 7.5, 100 mM NaCl, 2 mM β-mercaptoethanol, and 10% glycerol). The 90–350 μM MBP-NLS proteins were titrated into a sample cell containing 9–35 μM Kap104p. All ITC experiments were done at 20 °C with 35 rounds of 8-μl injections. Data were plotted and analyzed with a single-site binding model using MicroCal Origin software (version 7.0).

Supporting Information

Figure S1. Isothermal Titration Calorimetry Measurements of Kap104p Binding to (A) MBP–Hrp1p NLS and (B) MBP–Nab2p NLS.

Table S1. Summary of ITC Data for Kap104p Binding to Additional Hrp1p and Nab2p Mutants.

Figure S2. Binding Assays of Kapβ2 with Immobilized Nab2p NLS and Hrp1p NLS in the Presence and the Absence of RanGTP.

Table S2. Summary of ITC Data for Mutations of the PY Motif of Kap104p.

Figure S3. Kap104p Recognizes Basic PL-NLSs.

Table S3. Summary of ITC Data for Mutations of the PL Motif of Nab2p.

Figure S4. Kap104p Specificity.

Text S1. Differences in Interface Residues between Kap104p and Kapβ2.

Text S2. Prediction of Kapβ2 Homologs That Recognize hPy-NLSs.

Accession Numbers

Accession numbers for genes mentioned in this paper from the National Center for Biotechnology Information (http://www.ncbi.nlm.nih.gov) are: Bhp1p (855820), Cig1b (852567), Ino1NPA1 (3175), Hrp1p (853097), Kap104p (852305), Kapβ2 (3842), Nab2p (852755), Nabα9p (856486), Poc5p (855913), Rml2p (856660), Sin3p (854138), Sko1p (855534), Snp1p (854749), TAP (10482), and Tjg2p (852888).
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Author contributions. KES and YMC conceived and designed the experiments. KES and HG performed the experiments. KES and YMC analyzed the data. KES contributed reagents/materials/analysis tools. KES and YMC wrote the paper.

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