Nuclear import of prototype foamy virus transactivator Bel1 is mediated by KPNA1, KPNA6 and KPNA7

JIHUI DUAN, ZHIQIN TANG, HONG MU and GUOJUN ZHANG

Clinical Laboratory, Tianjin First Center Hospital, Tianjin 300192, P.R. China

Received October 9, 2015; Accepted May 31, 2016

DOI: 10.3892/ijmm.2016.2635

Abstract. Bel1, a transactivator of the prototype foamy virus (PFV), plays pivotal roles in the replication of PFV. Previous studies have demonstrated that Bel1 bears a nuclear localization signal (NLS); however, its amino acid sequence remains unclear and the corresponding adaptor importins have not yet been identified. In this study, we inserted various fragments of Bel1 into an EGFP-GST fusion protein and investigated their subcellular localization by fluorescence microscopy. We found that the PRQKRPR fragment, which accords with the consensus sequence K(K/R)X(K/R) of the monopartite NLS, directed the nuclear translocation of Bel1. Point mutation experiments revealed that K218, R219 and R221 were essential for the nuclear localization of Bel1. The results of GST pull-down assay revealed that the Bel1 peptide 215-221, which bears the NLS, interacted with Karyopherin alpha 1 (importin alpha 5) (KPNA1), karyopherin alpha 6 (importin alpha 7) (KPNA6) and karyopherin alpha 7 (importin alpha 8) (KPNA7). Finally, in vitro nuclear import assays demonstrated that KPNA1, KPNA6 or KPNA7, along with other necessary nuclear factors, caused Bel1 to localize to the nucleus. Thus, the findings of our study indicate that KPNA1, KPNA6 and KPNA7 are involved in Bel1 nuclear distribution.

Introduction

Foamy viruses (FVs), which comprise the Spumaretrovirinae in the retrovirus family, are also known as spumaretroviruses. FVs are found in primates, including humans, as well as in non-primates, including cows, cats and horses (1-5).

The prototype foamy virus (PFV) Tas protein, also known as Bel1, is a 300-amino-acid nuclear protein that is essential for virus replication (6), and can highly transactivate the PFV promoters, LTR and IP (7-9). Similar to most typical transcriptional activators, nuclear localization is required for the transactivation activity of Bel1 (10). Bel1 bears a putative nuclear localization signal (NLS) in the central highly basic region (11,12). Earlier studies have indicated that peptide 211-225 and/or 209-226 are necessary and sufficient for Bel1 nuclear localization (13-15). Later studies demonstrated that another two basic amino acids, R199-H200, also regulate Bel1 nuclear localization, which suggests that Bel1 carries a bipartite NLS consisting of residues 199-200 and residues 211-223 (10,16). However, Ma et al further found that residues R221R222, but not R199H200, are essential for the nuclear distribution of Bel1 (17).

Importin is a type of karyopherin (18) that transports protein molecules into the nucleus by binding to nuclear localization sequences. Importin has two subunits, karyopherin alpha (KPNA; also known as importin alpha) and karyopherin beta KPNB (also known as importin beta). Members of the KPNA family can bind and transport cargoes by themselves (19-21), or can form heterodimers with KPNB (22,23). As part of a heterodimer, KPNB mediates the interaction with nuclear pore complex (NPC), while KPNA acts as an adaptor protein to bind KPNB and the NLS on the cargo (24). The NLS-KPNA-KPNB trimer dissociates after binding to RanGTP inside the nucleus (25), with the two importin proteins being recycled to the cytoplasm for further use. Although KPNA and KPNB are used to describe importin as a whole, they actually represent larger families of proteins that share a similar structure and function. A variety of genes have been identified for both KPNA and KPNB, such as KPNA1-KPNA7 and KPNB1 (26). Different KPNA members show preferences for particular types of NLS cargo, although there is no absolute boundary (26,27).

In this study, we aimed to determine which adaptor importins are required for Bel1 nuclear translocation. We found that the PRQKRPR fragment, which accords with the consensus sequence K(K/R)X(K/R) of monopartite NLS, directs the nuclear localization of Bel1. Point mutation experiments revealed that residues K218, R219 and R221 were essential for the nuclear accumulation of Bel1. The results of GST pull-down assay revealed that the Bel1 NLS fragment 215-221 interacted with KPNA1, KPNA6 and KPNA7. Finally, in vitro nuclear import assays demonstrated that KPNA1, KPNA6 and KPNA7 caused Bel1 to localize to the nucleus. Our findings thus indicate that KPNA1, KPNA6 and KPNA7 are involved in Bel1 nuclear translocation.

Key words: nuclear import, Bel1, nuclear localization signal, KPNA, KPNB
Materials and methods

Plasmids. The Bell gene was amplified from the PFV full-length infectious clone, pCHFV, kindly provided by Maxine L. Linial (28). The mammalian cell expression plasmids, pC3-EGFP-X-GST, pC3-EGFP-NLS-GST, pC3-EGFP-BiNLS-GST, pC3-EGFP-Bel-GST, pC3-EGFP-215-221-GST and other truncated Bell plasmids were generated as previously described (17). The Bell mutants K218R, K218A, R219A and R221A were generated using a QuikChange™ site-directed mutagenesis kit (Stratagene, Palo Alto, CA, USA) and R221A was expressed by exposure to X-ray films (Kodak, Xiamen, China).

Cell culture and transfection, antibodies and reagents. HeLa and 293T cells (both from the Cell Bank of the Chinese Academy of Sciences, Shanghai, China) were grown in Dulbecco's modified Eagle's medium (high glucose; HyClone, Logan, UT, USA) supplemented with 10% fetal bovine serum (Gibco-BRL, Grand Island, NY, USA) and Pen Strep Glutamine (PSG) (Gibco). The cells were maintained in a humidified atmosphere containing 5% CO₂ at 37°C and transfected with polyethylamines (PEI) (Polysciences, Inc., Warrington, PA, USA) in accordance with the manufacturer's instructions.

Anti-EGFP (sc-9996), anti-GAPDH (sc-32233), anti-GST (sc-138) and HRP-conjugated goat anti-mouse secondary antibodies (sc-2005) were purchased from Santa Cruz Biotechnology, Inc. (Santa Cruz, CA, USA). Anti-Flag (F3165) antibody and 4,6-diamidino-2-phenylindole (DAPI) were purchased from Sigma-Aldrich. FITC-conjugated affinipure goat anti-mouse secondary antibodies (115-095-003) were purchased from Jackson ImmunoResearch Laboratories, Inc. (West Grove, PA, USA).

Immunoﬂuorescence microscopy assay (IFA). The HeLa cells were seeded on glass coverslips. Following fixation in 4% paraformaldehyde for 10 min on ice, the cells were permeabilized in 0.2% Triton X-100 for 10 min on ice. After blocking in 3% BSA + 5% fat-free milk at 4°C for 2 h, the cells were incubated with anti-EGFP antibodies at 4°C for a further 2 h, and subsequently washed with 0.1% Triton X-100 in PBS 5 times at room temperature. FITC-conjugated secondary antibodies were added at 4°C for 45 min. After the nuclei were stained with 0.2 µg/ml DAPI for 10 min at room temperature, the coverslips were observed under an Olympus IX71 ﬂuorescence microscope (Olympus, Tokyo, Japan).

In vivo GST pull-down assay and western blot analysis. The 293T cells were transfected with the pC3-EGFP-X-GST empty vector or pC3-EGFP-Bel1-GST along with plasmids that encode Flag-KPNAs or Flag-KPNB1; at 48 h post-transfection, the cell lysates were incubated with Glutathione Sepharose 4B beads (20182003-2) (GE Healthcare, Cleveland, OH, USA) for 4 h at 4°C. Samples from both cell lysates and Glutathione Sepharose 4B beads were mixed with loading buffer. After boiling for 20 min at 100°C, the protein samples were resolved by SDS-PAGE and transferred onto PVDF membranes. Prior to incubation first with primary antibodies overnight at 4°C and then with HRP-conjugated secondary antibodies for 1 h at room temperature, the membranes were blocked in 5% fat-free milk for 1.5 h at room temperature. After the membranes were treated with LuminataTM Western HRP chemiluminescence substrates (WBLUC0100; Millipore, Billerica, MA, USA), the specific protein signals were detected by exposure to X-ray films (Kodak, Xiamen, China).

In vitro nuclear import assay. In vitro nuclear transport assays were carried out as previously described with some modifications (29,30). Briefly, the HeLa cells (70-80% confluent), plated on glass coverslips, were washed 3 times with ice-cold transport buffer (TB) and permeabilized with digitonin (40 mg/ml) for 5 min on ice. The cells were then washed twice with...
ice-cold TB and soaked in TB for 10 min on ice. The complete transport solution contained import substrates (~2 μM), an adenosine triphosphate (ATP)-regenerating system (1 mM ATP, 5 mM creatine phosphate and 20 U/ml creatine phosphokinase) as a source of energy and some other soluble import factors. The import reaction was performed for 30 min at 37˚C or on ice in a humidified chamber. After the transport reaction, the cells were washed twice with ice-cold TB followed by fixation with 4% paraformaldehyde for 10 min on ice. The cells were washed 3 times first with TB and then twice with PBS. Following permeabilization with 0.2% Triton X-100 in PBS for 5 min on ice, the cells were blocked with 3% BSA + 5% fat-free milk in PBS and incubated with anti-EGFP antibodies and FITC-conjugated secondary antibodies as mentioned above. After being mounted on slides in PBS containing DAPI for 10 min on ice, the cells were visualized using an Olympus IX71 fluorescence microscope (Olympus).

**Results**

The NLS of Bel1 is \(215\text{PRQKRPR}^{221}\). In order to accurately determine the NLS of Bel1, we inserted into the EGFP-GST fusion protein the truncated fragments of Bel1 that encompass the amino acid positions 211-223, 221-223, 218-223, 217-223, 216-223, 215-223, 214-223, 213-223, 212-223 and 211-220 (Fig. 1) and observed their subcellular distribution by performing indirect IFA. The monopartite NLS of SV40 large T antigen (NLS) and the bipartite NLS of Xenopus laevis nucleoplasmin (BiNLS) were also inserted into EGFP-GST as positive controls for nuclear localization. As illustrated in Fig. 2, similar to the activity of SV40-NLS and the BiNLS, the 211-223 peptide of Bel1 enabled the nuclear localization of the fusion protein. In view of the fact that residues R\(^{221}\)R\(^{222}\)R\(^{223}\) are necessary for Bel1 nuclear distribution (10,13-17), we extended the N-terminal of the peptide segment to observe the effects. As shown in Fig. 2, the 221-223, 218-223, 217-223 and 216-223 fusion proteins still mainly distributed in the cytoplasm with little nuclear distribution, although containing the residues R\(^{221}\)R\(^{222}\)R\(^{223}\). Until the N-terminal extended to residue P215, the fusion protein 215-223 localized to the nucleus (Fig. 2). We then shortened the C-terminal of 215-223 to continue our observation. As shown in Fig. 2, the both (215-222)- and (215-221)-containing EGFP-GST fusion proteins were distributed in the nucleus, whereas 215-220 was distributed in the cytoplasm. Taken together, these data suggest that peptide \(215\text{PRQKRPR}^{221}\) is the NLS of Bel1 and is essential for nuclear distribution.

The NLS of Bel1 is monopartite. Sequence analysis indicated that \(215\text{PRQKRPR}^{221}\) accords with the consensus sequence K(K/R)X(K/R) of monopartite NLS, comprised primarily of lysine (K) and arginine (R) residues (31), wherein the basic amino acids are critical. To confirm this result, we generated four mutations of the Bel1 protein sequence, named K218R that changed K\(^{218}\) to R\(^{218}\), K218A that turned K\(^{218}\) into A\(^{218}\), R219A and R221A: Bel1 mutants. Capitalized letters represent amino acid sequences; the numbers denote the amino acid position in Bel1 protein; asterisks indicate Bel1 amino acids.

Bell interacts with KPNA1, KPNA2, KPNA6 and KPNA7. In the conventional nuclear transport pathway, cargoes are recognized and bound by the transport receptor adaptor, importin alpha, to translocate to the nucleus (32,33). In this study, in order to determine which importins mediate the transportation of Bel1 into the nucleus, we detected the interaction between Bel1 and 7 isoforms of importin alpha (KPNA1, KPNA2, KPNA3, KPNA4, KPNA5, KPNA6 and KPNA7) and the most common importin beta protein, KPNB1, in 293T cells by in vivo GST pull-down assay. The results of western blot analysis revealed that Bel1 interacted with KPNA1, KPNA2, KPNA6 and KPNA7 solidly, as opposed to other isoforms of importin alpha or KPNB1 (Fig. 4). This suggests that Bel1 may use KPNA1, KPNA2, KPNA6 and KPNA7 to enter the nucleus.
The NLS peptide of Bel1 interacts with KPNA1, KPNA6 and KPNA7 separately. Classical NLSs (cNLS) are directly recognized and bound by the adaptor protein importin alpha (34-36). To confirm this, we then determined the interrelation between truncated mutant 215-221, the NLS peptide of Bel1, and KPNA1, KPNA6 and KPNA7 in 293T cells by in vivo GST pull-down assay. As shown in Fig. 5, although KPNA2 bound to WT Bel1 (Fig. 4), truncated 215-221 did not interact with KPNA2. In accordance with the above findings, KPNA1, KPNA6 and KPNA7 bound solidly to the NLS sequence 215-221 of Bel1. These results further confirm that these three nuclear-import receptors are involved in the translocation of Bel1 into the nucleus.

Figure 2. Subcellular distribution of Bel1 truncated mutants (x400 magnification). HeLa cells were transfected with pC3-EGFP-X-GST vector or fusion expression plasmids by polyethylenimines (PEI) and fixed 24 h post-transfection. The subcellular localization of EGFP-GST fusion proteins was visualized by indirect fluorescence microscopy assay. Nuclei were stained with 4,6-diamidino-2-phenylindole (DAPI). Representative images are shown.

**Discussion**

As a key positive regulator of viral gene expression, Bel1 contains a conventional NLS and is located in the nucleus to conduct its transactivational activity (11,37). Previous studies have confirmed the key role of R^{221}R^{222}R^{223} in Bel1 nuclear import with recombinant transport receptors (Fig. 6). As a control for active transport, the SV40-NLS-containing EGFP-NLS-GST fusion protein was included in the same experiment. In contrast to KPNA2, KPNA1, KPNA6 and KPNA7 were sufficient for the nuclear accumulation of Bel1 (Fig. 6). The subcellular distribution of the positive control EGFP-NLS-GST was consistent with that previously reported (26,27): KPNA1, KPNA2 or KPNA6 were able to mediate SV40-NLS alone, while KPNA7 failed to do that. Taken together, these findings indicate that the efficient nuclear import of Bel1 in cells is mediated by KPNA1, KPNA6 and KPNA7 via the importin alpha/beta transport pathway.
Figure 3. Subcellular distribution of Bel1 wild-type (WT) and mutants (x400 magnification). HeLa cells were transfected with pC3-EGFP-Bel1-GST plasmids WT, K218R, K218A, R219A or R221A by polyethylenimines (PEI) and fixed 24 h post-transfection. The subcellular localization of EGFP-GST fusion proteins was visualized by indirect fluorescence microscopy assay. Nuclei were stained with 4',6-diamidino-2-phenylindole (DAPI). Representative images are shown.

Figure 4. Interaction between Bel1 and KPNAs or KPNB1. 293T cells were transfected with pC3-EGFP-X-GST empty vector or pC3-EGFP-Bel1-GST plasmid along with plasmids that encode Flag-KPNAs or Flag-KPNB1; 48 h post-transfection, cell lysates were incubated with Glutathione Sepharose 4B beads for 4 h at 4°C. Samples from both cell lysates and Glutathione Sepharose 4B beads were subjected to western blot analysis and probed with anti-Flag, anti-GST and anti-GAPDH antibodies. Representative results from 3 independent experiments are shown.
localization, yet the accurate nuclear localization sequence is controversial and the adaptor-mediated Bel1 nuclear transport is unclear.

In this study, with the purpose of defining the peptide sequences that are essential for the nuclear distribution of Bel1, we introduced an EGFP-GST fusion expression system that has been widely utilized in studying the subcellular localization of retrovirus transactivators (38,39). With the Bel1 shortened mutant 211-223, we finally confirmed that PRQKRPR is necessary and sufficient for the nuclear localization of Bel1. Furthermore, we found that residues KRPR of Bel1 are indispensable for its nuclear accumulation by the results of mutagenesis experiments. Comprehensive analysis of the consensus sequence K(K/R)X(K/R) of monopartite NLS indicated that KRPR is the core sequence of Bel1 NLS and the NLS of Bel1 is monopartite.

Consistent with the characteristics of the NLS sequence of mammalian cells, the residue K in the consensus sequence was replaced by the positive charge residue R, which had no change in the subcellular distribution. This suggests that the importance of the basic amino acid residues in the nuclear protein is closely related to the positive charge. That is to say, Bel1 may use a similar way to enter the nucleus

Figure 5. Interaction between 215-221 and KPNAs. 293T cells were transfected with pC3-EGFP-X-GST empty vector or the Bel1 truncated mutant 215-221, as for 215-221 in Fig. 1, along with plasmids encoding Flag-KPNAs; 48 h post-transfection, cell lysates were incubated with Glutathione Sepharose 4B beads for 4 h at 4°C. Samples from both cell lysates and Glutathione Sepharose 4B beads were subjected to western blot analysis and probed with anti-Flag, anti-GST and anti-GAPDH antibodies. Representative results from 3 independent experiments are shown.

Figure 6. In vitro nuclear import assay using KPNAs (x400 magnification). Digitonin-permeabilized HeLa cells were washed with transport buffer (TB) and then incubated with ~2 µM either EGFP-Bel1-GST or EGFP-NLS-GST in 50 µl TB containing adenosine triphosphate (ATP) regeneration system, 2 µM His-RanGTP WT, 2 µM GST-KPNA1 or GST-KPNA2 or GST-KPNA6 or GST-KPNA7 used in combination with 2 µM GST-KPNB1. The cells were washed with TB and then fixed and stained with 4',6-diamidino-2-phenylindole (DAPI). The cells were visualized by indirect fluorescence microscopy assay. Representative images are shown.

| EGFP-Bel1-GST | EGFP-NLS-GST |
|---------------|---------------|
| EGFP | DAPI | Merge | EGFP | DAPI | Merge |
| KPNA1 | | | | | |
| KPNA2 | | | | | |
| KPNA6 | | | | | |
| KPNA7 | | | | | |
the nucleus as a host cell transcription factor to complete its transactivational function.

NLSs are categorized into cNLSs and non-classical NLSs (ncNLS) (40). cNLSs are characterized by either monopartite (e.g., PKKRRK from SV40 large T antigen) or bipartite (e.g., KPRATKQKAKKKK from nucleoplasmin) stretches of basic amino acids (41,42). There is no consensus on whether different types of NLS have different biological functions. As RNA virus, the genome fidelity of foamy virus is lower than that of DNA genome. In the course of viral inheritance, monopartite NLS, less conserved nucleic acid sequence, may have some certain evolutionary advantages. In addition, monopartite NLS, shorter stretches of basic amino acids, may be more conducive to efficiently use of limited resources for virus.

Human KPNA isoforms are well conserved, with 26% identity and 42% conservation in their amino acid sequences (43,44). They can be divided into three subfamilies according to phylogenetic analysis: i) the α1 subfamily containing KPNA1, KPNA5 and KPNA6; ii) the α2 subfamily containing KPNA2 and KPNA7; and iii) the α3 subfamily containing KPNA3 and KPNA4. Although the α1 subfamily shares a maximum of 82.1% identity and 82% sequence conservation (26), their affinity for Bel1 differed markedly, which may due to the restriction of KPNA5 expression to the testes (45), in our GST pull-down experiments using 293T cells. The α2 subfamily is the least conserved of the KPNA subfamilies, with 55% identity and 71% conservation (24). In addition, phylogenetic analysis of the ARM repeats, responsible for identifying and combining with the NLS of cargo proteins (36), of the KPNA shows that the KPNA7 ARM repeats is more divergent than that of KPNA2 (26). To a certain extent, this explains the different performance of KPNA2 and KPNA7. Besides, the combination between KPNA2 and Bel1 may be involved in the other amino acids apart from the NLS, and/or the peptide PRQKRPR21 is not sufficient to mediate the binding between the two. It is thus revealed that KPNA2 may participate in other biological functions of Bell except nuclear transport.

The present study provided evidence that KPNA1, KPNA6 and KPNA7 may be ‘hijacked’ by PFV Bell for efficient nuclear import and viral replication. Given this fact, the restricted expression of KPNA isoforms may provide a mechanism for the suppression of PFV replication and disease progression.

Acknowledgements

This study was supported by a grant from the National Key Clinical Specialist Construction Programs of China (grant no. 2013-544). We would like to thank Maxine L. Linial (University of Washington and Fred Hutchinson Cancer Research Center) for the PFV full-length infectious clone pCHFV.

References

1. Broussard SR, Comuzzie AG, Leighton KL, Leland MM, Whitehead EM and Allan JS: Characterization of new simian foamy viruses from African nonhuman primates. Virology 237: 349-357, 1997.
2. Hatama S, Otake K, Omoto S, Murae Y, Ikemoto A, Mochizuki M, Takahashi E, Okuyama H and Fujii Y: Isolation and sequencing of infectious clones of feline foamy virus and a human/feline foamy virus Env chimera. J Gen Virol 82: 2999-3004, 2001.
3. Herchenröder O, Renne R, Loncar D, Cobb EK, Murthy KK, Schneider J, Mergia A and Luciw PA: Isolation, cloning, and sequencing of simian foamy viruses from chimpanzees (SFVcpz): high homology to human foamy virus (HFV). Virology 201: 387-399, 1994.
4. Materniak M, Bicka L and Kuzmak J: Isolation and partial characterization of bovine foamy virus from Polish cattle. Pol J Vet Sci 9: 207-211, 2006.
5. Tobaly-Tapiro J, Bitouin P, Neves M, Guillemin MC, Lecellier CH, Puvion-Dutilleul F, Giorgi B, Zientara S, Giron ML, de Thé H, et al: Isolation and characterization of an equine foamy virus. J Virol 74: 4064-4073, 2000.
6. Löchelt M, Zentgraf H and Flügel RM: Construction of an infectious DNA clone of the full-length human spumaretrovirus genome and mutagenesis of the bel 1 gene. Virology 184: 43-54, 1992.
7. He F, Blair WS, Fukushima J and Cullen BR: The human foamy virus Bel-1 transcription factor is a sequence-specific DNA binding protein. J Virol 70: 3902-3908, 1996.
8. Kang Y, Blair WS and Cullen BR: Identification and functional characterization of the Bel-1-DNA binding site located in the human foamy virus internal promoter. J Virol 72: 504-511, 1998.
9. Löchelt M, Flügel RM and Aboud M: The human foamy virus internal promoter directs the expression of the functional Bel 1 transactivator and Bet protein early after infection. J Virol 68: 6315-6318, 1994.
10. Chang J, Lee KJ, Jang KL, Lee EK, Baek GH and Sung YC: Human foamy virus Bet1 transactivator contains a bipartite nuclear localization determinant which is sensitive to protein context and triple multimerization domains. J Virol 69: 801-808, 1995.
11. Venkatesh LK, Theodorakis PA and Chinnadurai G: Distinct cis-acting regions in U3 regulate trans-activation of the human spumaretrovirus long terminal repeat by the viral bel1 gene product. Nucleic Acids Res 19: 3661-3666, 1991.
12. Flügel RM: Spumaviruses: a group of complex retroviruses. J Acquir Immune Defic Syndr 4: 739-750, 1991.
13. He F, Sun JD, Garrett ED and Cullen BR: Functional organization of the Bel-1 transactivator of human foamy virus. J Virol 67: 1896-1904, 1993.
14. Venkatesh LK and Chinnadurai G: The carboxy-terminal transactivation enhancement region of the human spumaretrovirus transactivator contains discrete determinants of the activator function. J Virol 67: 3868-3876, 1993.
15. Venkatesh LK, Yang C, Theodorakis PA and Chinnadurai G: Functional dissection of the human spumaretrovirus transactivator identifies distinct classes of dominant-negative mutants. J Virol 67: 161-169, 1993.
16. Lee CW, Chang J, Lee KJ and Sung YC: The Bel1 protein of human foamy virus contains one positive and two negative control regions which regulate a distinct activation domain of 30 amino acids. J Virol 68: 2708-2719, 1994.
17. Liu J, Tan J, Cui X, Luo D, Yu M, Liang C and Qiao W: Residues R(199)H(200) of prototype foamy virus transactivator Bell contribute to its binding with LTR and IP promoters but not its nuclear localization. Virology 449: 215-223, 2014.
18. Görlich D, Prehn S, Laskey RA and Hartmann E: Isolation of a protein that is essential for the first step of nuclear protein import. Cell 79: 767-778, 1994.
19. van der Watt PJ, Stowell CL and Leaner JD: The nuclear import receptor Kpnβ1 and its potential as an anticancer therapeutic target. Crit Rev Eukaryot Gene Expr 23: 1-10, 2013.
20. Flores K and Seger R: Stimulated nuclear import by β-like importins. F1000Prime Rep 5: 41, 2013.
21. Zehorai E and Seger R: Beta-like importins mediate the nuclear translocation of mitogen-activated protein kinases. Mol Cell Biol 34: 259-270, 2014.
22. Goldfarb DS, Corbett AH, Mason DA, Harreman MT and Adam SA: Importin alpha: a multipurpose nuclear-transport receptor. Trends Cell Biol 14: 505-514, 2004.
23. Cimica V, Chen HC, Iyer JK and Reich NC: Dynamics of the STAT3 transcription factor: nuclear import dependent on Ran and importin-β1. PLoS One 6: e20188, 2011.
24. Pumroy RA and Cingolani G: Diversification of importin-α isoforms in cellular trafficking and disease states. Biochem J 466: 13-28, 2015.
25. Mattaj IW and Englmeier L: Nuclearcytoplasmic transport: the soluble phase. Annu Rev Biochem 67: 265-306, 1998.
26. Kelley JB, Talley AM, Spencer A, Gioeli D and Paschal BM: Karyopherin alpha7 (KPNA7), a divergent member of the importin alpha family of nuclear import receptors. BMC Cell Biol 11: 63, 2010.

27. Köhler M, Speck C, Christiansen M, Bischoff FR, Prehn S, Haller H, Görlich D and Hartmann E: Evidence for distinct substrate specificities of importin alpha family members in nuclear protein import. Mol Cell Biol 19: 7782-7791, 1999.

28. Life RB, Lee EG, Eastman SW and Linial ML: Mutations in the amino terminus of foamy virus Gag disrupt morphology and infectivity but do not target assembly. J Virol 82: 6109-6119, 2008.

29. Cassany A and Gerace L: Reconstitution of nuclear import in permeabilized cells. Methods Mol Biol 464: 181-205, 2009.

30. Adam SA, Marr RS and Gerace L: Nuclear protein import in permeabilized mammalian cells requires soluble cytoplasmic factors. J Cell Biol 111: 807-816, 1990.

31. McLane LM and Corbett AH: Nuclear localization signals and human disease. IUBMB Life 61: 697-706, 2009.

32. Marfori M, Lonhienne TG, Forwood JK and Kobe B: Structural basis of high-affinity nuclear localization signal interactions with importin-α. Traffic 13: 532-548, 2012.

33. Jans DA, Xiao CY and Lam MH: Nuclear targeting signal recognition: a key control point in nuclear transport? BioEssays 22: 532-544, 2000.

34. Stewart M: Molecular mechanism of the nuclear protein import cycle. Nat Rev Mol Cell Biol 8: 195-208, 2007.

35. Marfori M, Mynott A, Ellis JJ, Mehdi AM, Saunders NF, Curmi PM, Forwood JK, Bodén M and Kobe B: Molecular basis for specificity of nuclear import and prediction of nuclear localization. Biochim Biophys Acta 1813: 1562-1577, 2011.

36. Fontes MR, Teh T and Kobe B: Structural basis of recognition of monopartite and bipartite nuclear localization sequences by mammalian importin-alpha. J Mol Biol 297: 1183-1194, 2000.

37. Keller A, Partin KM, Löhelt M, Bannert H, Flügel RM and Cullen BR: Characterization of the transcriptional trans activator of human foamy retrovirus. J Virol 65: 2589-2594, 1991.

38. Meertens L, Chevalier S, Weil R, Gessain A and Mahieux R: A 10-amino acid domain within human T-cell leukemia virus type 1 and type 2 tax protein sequences is responsible for their divergent subcellular distribution. J Biol Chem 279: 43307-43320, 2004.

39. Gu L, Tsuji T, Jarboui MA, Yeo GP, Sheehy N, Hall WW and Gautier VW: Intermolecular masking of the HIV-1 Rev NLS by the cellular protein HIC: novel insights into the regulation of Rev nuclear import. Retrovirology 8: 17, 2011.

40. Korlimarla A, Bhandary L, Prabhu JS, Shankar H, Sankaranarayanan H, Kumar P, Remacle J, Natarajan D and Sridhar TS: Identification of a non-canonical nuclear localization signal (NLS) in BRCA1 that could mediate nuclear localization of splice variants lacking the classical NLS. Cell Mol Biol Lett 18: 284-296, 2013.

41. Soniat M and Chook YM: Nuclear localization signals for four distinct karyopherin-β nuclear import systems. Biochem J 468: 353-362, 2015.

42. Lange A, Mills RE, Lange CJ, Stewart M, Devine SE and Corbett AH: Classical nuclear localization signals: Definition, function, and interaction with importin alpha. J Biol Chem 282: 5101-5105, 2007.

43. Larkin MA, Blackshields G, Brown NP, Chenna R, McGettigan PA, McWilliam H, Valentin F, Wallace IM, Wilm A, Lopez R, et al: Clustal W and Clustal X version 2.0. Bioinformatics 23: 2947-2948, 2007.

44. Henikoff S and Henikoff JG: Amino acid substitution matrices from protein blocks. Proc Natl Acad Sci USA 89: 10915-10919, 1992.

45. Köhler M, Ansieau S, Prehn S, Leut A, Haller H and Hartmann E: Cloning of two novel human importin-alpha subunits and analysis of the expression pattern of the importin-alpha protein family. FEBS Lett 417: 104-108, 1997.