The Nucleoplasmin Nuclear Location Sequence Is Larger and More Complex than That of SV-40 Large T Antigen

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Abstract. The carboxy-terminal tail of nucleoplasmin, which specifies entry into the cell nucleus, contains four short sequences that are similar to previously identified nuclear location sequences. We show that none of these is able to locate chicken muscle pyruvate kinase to the cell nucleus. Deletion analysis was used to determine the limits of a nuclear location sequence and indicated that a 14-amino acid segment (RPAATKKAGQAKKK) should function as a minimal nuclear location sequence. When tested directly, however, this sequence was unable to locate pyruvate kinase to the cell nucleus. Restoration of three amino acids of nucleoplasmin sequence at either end of this sequence generated sequences that were able to locate pyruvate kinase to the cell nucleus.

The 14-amino acid proposed minimal nuclear location sequence is present in the functional sequences, AVKRPAATKKAGQAKKK, RPAATKKAG-QAKKKKLD, and the sequence AVKRPAATKKAG-QAKKKKLD, which has additional amino acids at both ends. The minimal sequence element is therefore necessary but not sufficient for transport into the cell nucleus. This unusual feature of the nucleoplasmin nuclear location sequence suggests ways in which it could interact with the nuclear transport mechanism.

In addition to its established role in chromatin assembly and histone storage in the Xenopus oocyte and egg (4, 16, 18, 21), nucleoplasmin has emerged as the protein of choice for studies of protein transport into the cell nucleus (reviewed in reference 5). Investigations using this acidic, thermostable, pentameric protein provided the decisive evidence that a mechanism of uptake involving selective entry into the nucleus operates in oocytes (8).

Nucleoplasmin has also been used to demonstrate that entry into the Xenopus oocyte nucleus occurs through the nuclear pore complex (9), to evaluate the transport properties of nuclei reconstituted in vitro, and to show that ATP is required for the transport process (28).

Primary amino acid sequences specifying entry into the nucleus have been identified in a number of proteins (5, 23, 25, 30). The sequence that has been studied in greatest molecular detail is the SV-40 large T antigen nuclear location sequence (3, 11, 14, 15, 19, 20, 33, 37). The earlier studies in this series demonstrated that the amino acid sequence PKK~28KRKV is able to translocate chicken muscle pyruvate kinase to the cell nucleus (15).

The chicken muscle pyruvate kinase expression vector system has also been used to map the polyoma virus large T nuclear location sequences (32) and more recently to investigate the effect of protein context upon the function of the SV-40 large T antigen nuclear location sequence (33). Therefore we chose this system to identify the nuclear location sequence or sequences in the nucleoplasmin “tail” region since a large body of data exists against which we can interpret our results.

We have sequenced nucleoplasmin and shown that the tail region which is both necessary and sufficient for selective entry of nucleoplasmin into the nucleus corresponds to the carboxy-terminal 50 amino acids (6, 7). In addition we have pointed out the presence of four sequences in the tail region which show significant degrees of similarity to other nuclear location sequences (6, 7).

Recently Burglin and De Robertis (1) have shown that one-half of the tail region of nucleoplasmin can translocate Escherichia coli β-galactosidase to the Xenopus oocyte nucleus. This region includes a sequence which bears a striking similarity to the SV-40 large T antigen nuclear location sequence, differing only by two amino acid substitutions. Each of these substitutions alone still allows the SV-40 large T antigen sequence to locate the protein to the cell nucleus (37).

Here we describe the construction of recombinant plasmids in which portions of the nucleoplasmin tail region are linked to the chicken muscle pyruvate kinase cDNA. We confirm that the entire nucleoplasmin tail region can translocate a cytoplasmic protein to the cell nucleus (in agreement...
with reference 1) and that the sequence that is responsible for this activity lies in the amino-terminal half of the tail. We find, however that the sequence in this half of the tail region which most strongly resembles the SV-40 nuclear location sequence is unable to locate chicken muscle pyruvate kinase to the cell nucleus when fused to the carboxy or amino terminus of the protein. In fact none of the four sequences similar to other nuclear location sequences that we have observed and discussed previously (1, 6, 7) is capable of translocating pyruvate kinase to the cell nucleus.

Deletion analysis reveals that a single sequence in the nucleoplasmin tail region is capable of translocating pyruvate kinase to the nucleus. However this sequence is longer and more complex than previously identified nuclear location sequences. The unusual properties of this sequence suggest a model for the interaction of the nuclear location sequence with the putative receptor in the transport mechanism.

**Materials and Methods**

**Microinjection**

Vero (African Green Monkey) cells were grown on glass coverslips and microinjected with glass capillaries as described previously (12, 31). Plasmid DNAs were injected at a concentration of 100 µg/ml into the nucleus delivering ~10,000 molecules/cell. Proteins at a concentration of 1 mg/ml were injected into the nucleus or cytoplasm of the cell delivering on the order of 10⁶ molecules/cell.

**Immunofluorescence Microscopy**

Nucleoplasmin/ pyruvate kinase fusion proteins were detected with rabbit polyclonal antiserum against purified chicken muscle type M1 pyruvate kinase (22) followed by FITC-conjugated goat anti-rabbit IgG (Nordic Immunological Labs., Maidenhed, UK) or with a mixture of monoclonal antibodies to the tail region of nucleoplasmin (7) followed by FITC-conjugated rabbit anti-mouse IgG.

**Proteins**

Nucleoplasmin, the trypsin resistant core of nucleoplasmin and the 12-kD tail fragment of nucleoplasmin were all prepared using previously published procedures (8).

**Plasmid Construction**

**Vector Fragments.** The materials and methods for plasmid construction are identical to those described previously (15, 32, 33).

Nucleoplasmin Fragments. An M13 clone containing a full-length nucleoplasmin cDNA (7) with Eco RI linkers was used as the start material for the production of nucleoplasmin DNA fragments. All procedures used were conventional techniques described in detail elsewhere (24).

pPK-NPI. A fragment encompassing the tail region (amino acids 149-200) was prepared by digestion of 1 µg of the M13 clone (RF) with Hinf I, repaired with DNA polymerase (Klenow) and addition of an Xho I linker. The required Xho-Pk fragment was purified from an acrylamide gel and ligated into suitably cleaved plasmid (pUC 13-NP3) which had the required nucleoplasmin fragment and a universal terminator linker. After digestion with Xho I and Eco RI this fragment was purified from an acrylamide gel and ligated to the appropriate vector fragments to produce pPK-NP3.

pPK-NP4. 1 µg of the plasmid pUC13-NP3 was digested with Bst NI, repaired with DNA polymerase large fragment, and Xho linkers were added. The DNA was digested with Xho I and Eco RI and the DNA fragment encoding amino acids 165-176 of nucleoplasmin was purified from an acrylamide gel and ligated to the appropriate vector fragments to produce pPK-NP4.

pPK-NPS. An Xho I-Eco RI fragment encoding amino acids 165-176 was ligated in a three fragment ligation with a Bam HI-Xho I fragment from pSSR 40 and an Eco RI-Bam HI fragment from RL18 PK12 to link the nucleoplasmin sequence at the amino terminus of pyruvate kinase.

pPK-NP5. 1 µg of the plasmid pUC13-NP3 was digested with Bst NI, repaired with the DNA polymerase large fragment, and a universal translation terminator linker and an Eco RI linker were added to the repaired ends. The resulting Xho I-Eco RI fragment encoding amino acids 149-165 was purified from an acrylamide gel and ligated to the appropriate vector fragment to produce pPK-NP5.

pPK-NP6. 1 µg of the plasmid pUC13-NP1 was digested with Bst NI, repaired with DNA polymerase large fragment, and an Xho I linker was added to the repaired ends. The DNA preparation was then digested with Eco RI and the appropriate Xho I-Eco RI fragment encoding amino acids 165-200 was purified from an acrylamide gel and ligated to the appropriate vector fragment to produce pPK-NP6.

Bal31 Deletion. 10 µg of the plasmid pUC13-NP3 was linearized with Eco RI or Xho I. The linear DNA molecule was digested with Bal31 nuclease (New England Biolabs, Beverly, MA) at 30°C for 5 min. Appropriate linkers were added to the digested DNA and excess linkers were removed by restriction enzyme digestion. The linear plasmid was purified from an agarose gel, ligated to form circular molecules, and used to transform competent E. coli strain 603. DNA was prepared from colonies of transformed cells and screened by restriction enzyme digestion. Fragments showing a reduction in size were sequenced and suitable deletions were ligated to the pyruvate kinase vector fragment to give the pPKΔ series of plasmids.

**Minimal Sequence Constructs**

The Xho I-Eco RI fragments encoding the chosen deletions were subcloned into pUC13 and grown in NM545 (thr leu b plsp1 dam dac [GM 48 Sm])'. The Xho I-Eco RI and Eco RI-Eco RI fragments from the chosen original deletions were ligated to the appropriate pyruvate kinase vector fragment in a three fragment ligation. An Xho I-Eco RI fragment was excised from the recombinants and sequenced to confirm their identity.

**Results**

**Selective Entry Model for Nuclear Accumulation of Proteins Is Applicable to Mammalian Cells**

We wished to determine if the selective entry model for nuclear accumulation established in *Xenopus* oocytes (8) applies to cultured mammalian cells since these cells were used in the analysis of the nuclear location sequence (see below). Therefore, as a prelude to these studies, the transport mechanism for nucleoplasmin was investigated by microinjection of unlabeled protein followed by detection with antibodies.

Nucleoplasmin accumulates rapidly in the cell nucleus after microinjection into the cytoplasm (Fig. 1 a). To determine the initial rate of nucleoplasmin uptake, cells were cooled to 20°C, microinjected with protein, and then fixed (Fig. 1 b). This would correspond to ~2 min after the injection of the first cell in a group of cells. A number of cells show protein accumulation within the nucleus while in others the protein is predominantly cytoplasmic. This indicates that under normal conditions the bulk of the injected nucleoplasmin...
would accumulate in the nucleus in <1 min. In some instances a distinct, sharply defined zone of increased staining is observed at the level of the nuclear envelope (Fig. 1 b). We and others have shown elsewhere, that inhibition of ATP generation reversibly blocks intranuclear accumulation but not perinuclear association, and on this basis have proposed a two step mechanism for the accumulation of proteins in the cell nucleus (27, 35). The trypsin-resistant core of nucleoplasmin does not enter the nucleus (Fig. 1 c) while the purified tail fragment does (Fig. 1 d). In the latter case distinct punctate cytoplasmic fluorescence is observed and contrasts with the absence of cytoplasmic staining in cells injected with intact nucleoplasmin (compare Fig. 1, a and d). The tail fragment may be unstable in these cells as it is in *Xenopus* oocytes and the fluorescent spots may represent regions of protein turnover.

In the *Xenopus* oocyte the decisive evidence for a selective entry mechanism for nuclear accumulation was the retention within the nucleus of the trypsin-resistant core of nucleoplasmin after direct microinjection into the nucleus (8). When the same experiment was performed in Vero cells the nucleoplasmin core molecule remains stably in the nucleus (Fig. 1 e).

These experiments demonstrate that as in the *Xenopus* oocyte nucleoplasmin accumulates extremely rapidly in the cell nucleus by a selective entry mechanism and not by free diffusion and binding within the nucleus.

**Identification of a Region Encoding a Nuclear Location Signal (NLS)**

As mentioned above the sequence of a full-length nucleoplasmin cDNA has been determined and we have shown that the nucleoplasmin polypeptide is 200 amino acids long and the carboxy-terminal 50 amino acids constitute the tail region that specifies entry into the nucleus (7). In this region there are four sequences which are similar to nuclear location signal sequences that have been identified in other nuclear proteins (6, 7). Two regions show close similarity to the SV-40 large T antigen NLS (37) and two are similar to the consensus sequence identified in the MAT α2 protein of yeast (13).

Fig. 2 a shows the location of these putative nuclear location sequences in the tail region and the position of selected restriction enzyme cleavage sites. These sites were used to construct recombinant plasmids (Fig. 2 b) in which these se-
sequences are linked to the COOH terminus or NH2 terminus of chicken muscle pyruvate kinase using the plasmid vector system described previously (15, 33). The intracellular distribution of the fusion protein expressed from each plasmid was examined by indirect immunofluorescence with antiserum against pyruvate kinase and where possible with monoclonal antibodies against the tail region of nucleoplasmin.

When the carboxy-terminal 51 amino acids of nucleoplasmin (residues 149-200) that constitute the entire tail region are linked to the carboxyterminus of pyruvate kinase (plasmid pPK-NP1), the fusion protein becomes located in the cell nucleus (Fig. 3). Both an antiserum against pyruvate kinase and a mixture of three tail-specific monoclonal antibodies specific for the tail region of nucleoplasmin showed strong nuclear fluorescence (Fig. 3, a and a'). This result indicates that the entire nucleoplasmin tail sequence can locate PK to the cell nucleus.

In pPK-NP2 the carboxy-terminal 23 amino acids of nucleoplasmin (residues 178-200) are linked to the carboxyterminus of PK. The fusion protein expressed from this plasmid was unable to locate to the cell nucleus as shown by the presence of cytoplasmic fluorescence with antiserum to pyruvate kinase (Fig. 3 b). The three tail-specific monoclonal antibodies did not give a fluorescent signal with the fusion protein. This lack of reaction and the inability of the fusion protein to locate to the cell nucleus is not due to linkage of the two DNA fragments in the incorrect reading frame as the same result was obtained when the nucleoplasmin fragment was linked to pyruvate kinase in all three reading frames (data not shown) and agrees with the findings of Burglin and De Robertis (1) using β-galactosidase.

Therefore the sequence lying between amino acids 178 and 200 of the nucleoplasmin tail region is unable to locate PK to the cell nucleus even though it contains sequences that resemble previously identified nuclear location sequences (see above). Also the three monoclonal antibodies to nucleoplasmin do not recognise an epitope in the 23 amino acids at the extreme COOH terminus of nucleoplasmin.

In the plasmid pPK-NP3, amino acids 149-176 of nucleoplasmin are linked to the COOH terminus of pyruvate kinase. The fusion protein expressed from this plasmid is located in the cell nucleus as indicated by staining with both the antiserum to pyruvate kinase and the tail-specific anti–nucleoplasmin monoclonal antibodies (Fig. 3, c and c'). This demonstrates that this region of nucleoplasmin contains a sequence which can locate pyruvate kinase to the cell nucleus. This region has also been shown to locate β-galactosidase to the Xenopus oocyte nucleus (1). In this 27-amino acid segment there is a sequence (residues 156-162) that shows a reasonable similarity to the yeast MAT α2 consensus and a sequence (residues 166-172) that is strikingly similar to the SV-40 large T antigen nuclear location sequence. These two sequences encoded within the nucleoplasmin fragment in pPK-NP3 were separately linked to the COOH terminus of pyruvate kinase to give two plasmids, pPK-NP4 and pPK-NP5 (Fig. 2 b). Surprisingly the fusion protein expressed from either of these plasmids is located in the cell cytoplasm (Fig. 3, d and d'). The nucleoplasmin fragment in pPK-NP4 which encodes the sequence with closest similarity to the SV-40 nuclear location sequence is also unable to locate the fusion protein to the cell nucleus when linked to the amino terminus (pNP4-PK) rather than the carboxy terminus (pPK-NP5-PK) (Fig. 2; Fig. 3, e and e'). These fusions are directly comparable with those used to

Figure 2. (a) Diagramatic representation of the tail region of nucleoplasmin. The bar represents amino acids 149–200 of the nucleoplasmin tail (7). Residue 200 is the carboxy terminus of nucleoplasmin. Protein sequences similar to the MAT α2 consensus or the SV-40 nuclear location sequence are shown above. Restriction enzyme cleavage sites in the DNA are indicated by arrows below. (b) Structure and subcellular location of nucleoplasmin/pyruvate kinase fusion proteins. The plasmid vectors listed on the left were made as described in Materials and Methods. The fusion proteins consist of a major part of pyruvate kinase (solid bars) to which are linked sequences from the nucleoplasmin tail region (hatched bars). The subcellular location of the fusion protein listed on the right was determined by immunofluorescence assay after microinjection of plasmid DNA into Vero cells. N, nuclear location; C, cytoplasmic location.
demonstrate the ability of the SV-40 nuclear location sequence to locate pyruvate kinase to the cell nucleus. Therefore we have demonstrated conclusively that the sequence in nucleoplasm that is closely similar to the SV-40 nuclear location sequence is unable to locate PK to the cell nucleus when it is fused to either the amino or carboxy terminus of the protein.

To establish that the sequence encoded in pPK-NP3 is the
only sequence in the tail region of nucleoplasmin that is capable of translocating pyruvate kinase to the cell nucleus, we linked amino acids 165-200 of nucleoplasmin to the carboxy terminus of pyruvate kinase (plasmid pPK-NP6 in Fig. 2 b). The fusion protein expressed from this plasmid is located in the cell cytoplasm.

We therefore undertook a detailed analysis of the region of nucleoplasmin (amino acids 149-176) linked to pyruvate kinase in pPK-NP3 in order to determine precisely the sequence responsible for nuclear location.

**Deletion Analysis of the Nuclear Location Sequence**

The DNA fragment which encodes amino acids 149-176 of nucleoplasmin was treated with Bal 31 nuclease (see Materials and Methods) to generate molecules with deletions from either the 5' (amino) end or the 3' (carboxyl) end of this fragment. The amino acid sequences of the deletions produced by this procedure and the subcellular location of the corresponding fusion protein are shown in Fig. 4 top. The immunofluorescence patterns observed for certain of the deletions are shown in Fig. 4 bottom.

**Deletions from the Amino Terminus.** Analysis of deletions from the amino terminus (amino acid 149) indicate a transition of the location of the fusion protein from nuclear to cytoplasmic when the deletion end point lies in the region of amino acid 157 (proline).

In PK-Δ1 the fusion protein is nuclear (Fig. 4 bottom, a). The end point of this deletion lies at arginine residue 156. However the lysine residue 155 present in the original fragment is replaced in the deletion by an arginine residue encoded by the linker DNA. Therefore the presence of two basic residues at this location is maintained; lys-arg-pro becomes arg-arg-pro.

In pPK-Δ4 the fusion protein is cytoplasmic (Fig. 4 bottom, b). The end point of this deletion lies at alanine residue 158. In this case the nucleoplasmin sequence lys-arg-pro is

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*Figure 4. (Top) Deletion analysis of the minimal nuclear location sequence. The plasmid vectors listed on the left were made as described in Materials and Methods. Nucleoplasmin sequences (boxed) were linked to the COOH terminus of pyruvate kinase. Amino acid sequences encoded by linker DNA are shown outside the boxed sequences. Numbers indicate amino acids in the nucleoplasmin sequence. Subcellular location of the fusion proteins (N, nuclear; C, cytoplasmic) was determined by immunofluorescence assay after microinjection of plasmid DNAs into Vero cells. (Bottom) Subcellular location of pyruvate kinase/nucleoplasmin fusion proteins. Plasmid DNAs encoding the fusion proteins shown in the top were microinjected into the cell nucleus of Vero cells. After overnight incubation the cells were fixed and stained for pyruvate kinase as described in Materials and Methods. (a) pPK-Δ1; (b) pPK-Δ4; (c) pPK-Δ159; (d) pPK-Δ190.*
Figure 5. (Top) Minimal sequence for nuclear location of pyruvate kinase fusion proteins. The plasmid vectors listed on the left were made as described in Materials and Methods. Nucleoplasmin sequences (boxed) were linked to the carboxy terminus of pyruvate kinase. Amino acid sequences encoded by the linker DNA are shown outside the boxed sequences. (Bottom) Subcellular location of pyruvate kinase/nucleoplasmin fusion proteins carrying the proposed minimal nuclear location sequences. Plasmid DNA encoding the fusion protein shown in the top were microinjected into the nucleus of Vero cells. After overnight incubation the cells were fixed and stained for pyruvate kinase as described in Materials and Methods. (a) pPK-A1/A159; (b) pPK-A2/A159; (c) pPK-A1/A138; (d) pPK-A2/A138.

Deletion from the Carboxy Terminus. Deletion of amino acids from the carboxy terminus (amino acid 176) reveals that only part of the lysine tract (residues 167-170) is necessary for location to the cell nucleus. The fusion protein expressed from pPK-A159 bears amino acids 149-169 of nucleoplasmin (Fig. 4 top) and three of the four contiguous lysine residues remain. This fusion protein is located in the cell nucleus (Fig. 4 bottom, c), however, when these three remaining lysine residues are removed (pPK-A190) the fusion protein is located in the cell cytoplasm (Fig. 4 bottom, d).

This result clearly shows that within the context of this fragment, at most three of the four lysine residues are necessary for the location of pyruvate kinase to the cell nucleus, however, as described below the local sequence context of these deletions influences their ability to locate the fusion protein to the nucleus.

Minimal Sequence for Nuclear Location

The deletion analyses described above indicate that a minimal sequence of fourteen amino acids (amino acids 156-169) should be sufficient to locate pyruvate kinase to the cell nucleus. This minimal sequence was tested directly by linking a fragment of DNA from pPK-A1 (nuclear located fusion protein) to a fragment of DNA from pPK-A159 (nuclear located fusion protein) through the common EcoRII site (Fig. 5 top).

Surprisingly we found that the pyruvate kinase fusion protein bearing this proposal minimal sequence (pPK-A1/A159) was located in the cell cytoplasm (Fig. 5 bottom, a).

To determine which additional sequences were required to generate a sequence which could locate pyruvate kinase to the nucleus we restored nucleoplasmin sequence at the amino-terminal end (pPK-A2/A159), the carboxyl-terminal end (pPK-A1/A138), or at both ends (pPK-A2/A138). We found that the fusion proteins from all three plasmid con-
The rapid rate of accumulation of nucleoplasmin that we have observed is greater than that reported for fluorescently labeled nucleoplasmin in hepatoma tissue culture heterokaryons (34). We feel that this difference is probably due to the chemical modification of lysine residues by their fluorochrome leading to reduced efficiency of the nuclear location sequence.

None of the Putative Nuclear Location Sequences in the Tail Region Can Locate Pyruvate Kinase to the Cell Nucleus

As mentioned above we chose the pyruvate kinase vector system to analyze the nuclear location sequence in nucleoplasmin because of the wealth of data that has been accumulated for the SV-40 and polyoma virus large T antigen nuclear location sequences using this system. The advantage of pyruvate kinase is its exclusively cytoplasmic distribution while the structural data available for the closely homologous cat muscle enzyme (26) indicates that the carboxy and amino termini are both exposed in the monomer and are ideal locations for the placement of a nuclear location sequence. E. coli β-galactosidase has been used as a marker protein for several studies of nuclear location sequences (1, 13, 25, 36). However, β-galactosidase enters the nucleus of cells to some extent (15) complicating the assay by producing a background level against which the activity of any putative nuclear location sequence must be assessed.

The investigations described here have led to the surprising finding that none of the four sequences in the nucleoplasmin tail region that resemble other nuclear location sequences are able to locate pyruvate kinase to the cell nucleus. In particular, one of these sequences differs from the SV-40 large T antigen nuclear location sequence at two amino acid positions, both of which have been mutated individually and shown not to influence nuclear accumulation of large T (37). However we find that this similar sequence from nucleoplasmin is unable to locate pyruvate kinase to the cell nucleus whether it is linked to the amino acid or carboxy terminus of the protein. Instead we have demonstrated that a single sequence able to locate pyruvate kinase to the cell nucleus lies between amino acids 149 and 176, in the amino-terminal 28 amino acids of the tail region.

Minimal Sequence of 14 Amino Acids Is Revealed by Deletion Analysis

Deletion analysis of the 28 amino acid segment able to locate pyruvate kinase to the nucleus allowed us to define the limits of a nuclear location sequence.

Deletion of amino acids from the amino terminus of this segment showed that a transition in the location of the fusion protein from nuclear to cytoplasmic is concomitant with simultaneous substitution of lysine155 by alanine, and proline157 by glycine. Both of these substitutions are nonconservative; a positively charged residue (lysine) is replaced by a hydrophobic amino acid (alanine) and proline, which fixes the conformation of the polypeptide backbone, is replaced by glycine which permits maximum flexibility. Analysis of single mutations in the sequence is required to determine the individual contribution of these two amino acid substitutions to the transport defect.

Deletion of amino acids from the carboxy terminus of this segment removes lysine residues which are part of the sequence showing greatest similarity to the SV-40 large T antigen nuclear location sequence. Removal of the most carboxyterminal lysine residue does not abolish transport into the cell nucleus but removal of the next three does. This shows that at most three basic amino acids are needed at this end of the sequence for efficient nuclear location.

However, the minimal sequence defined by these end points (RPAATKKQAKKK) is unable to locate pyruvate kinase to the cell nucleus, indicating that the context within which this sequence is presented influences its activity. Thus, restoration of three amino acids of the original nucleoplasmin sequence at either the amino- or carboxy-terminal end of this minimal sequence (or both ends) produces sequences that can locate pyruvate kinase to the cell nucleus. Therefore the functional nuclear location sequences contain the proposed minimal sequence as a necessary central sequence element. This sequence has periodic structure in which two sequences of four hydrophobic residues separate three basic regions.

It is notable that the functional sequences contain a sequence showing similarity to the yeast MATα2 nuclear location sequence and the sequence having the closest similarity to the SV-40 large T antigen nuclear location sequence.

The unusual finding that restoring sequences at one end or the other is sufficient to generate a functional nuclear location sequence suggests models for the interaction of this sequence with a putative receptor in the nuclear transport machinery. We will briefly consider three.

First, it could be envisaged that the termini of the minimal sequence defined by the deletions described above represent two discrete binding domains that interact with corresponding binding sites in the receptor. Disruption of one binding domain can be tolerated but only as long as the other domain is maintained intact. A sequence in which both domains are disrupted cannot locate the fusion protein to the cell nucleus due to the combinatorial effect of weak binding at both sites.

Secondly, the nucleoplasmin–receptor interaction may not be limited to distinct sites but could occur over an extended region which includes these sites. Weak interaction at either...
end of the region can be tolerated, but weak interaction at both ends reduces the overall binding to a level that renders the sequence inactive. In this case amino acids in the central region of this sequence would be expected to play a significant role in the function of this sequence. Third, a precise conformation is necessary for recognition and transport and only the longer sequences can assume this conformation.

These models can be tested by in vitro mutagenesis.

**Comparison with Other Nuclear Location Sequences**

The properties of the nucleoplasmin nuclear location sequence described here are unusual but there are indications that other nuclear location sequences may have similar features. In the case of N1, another abundant, acidic Xenopus nuclear protein also involved in chromatin assembly (4, 16, 18), there is a sequence which closely resembles the SV-40 nuclear location sequence (17). As in nucleoplasmin, this similar sequence is not sufficient to locate the protein to the nucleus. Further amino acids are required that provide additional positively charged residues in a manner analogous to nucleoplasmin (16a).

In bovine HMG 1 (29) there is a sequence that bears a striking resemblance to the nucleoplasmin nuclear location sequence identified here. This sequence is KKGVVKASKSKKK, compared with (AVK)RPAATKQAGQAKK (KLD) in nucleoplasmin and lies on the amino-terminal side of the polyglutamic acid tract.

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