Nuclear Import of the Ran Exchange Factor, RCC1, Is Mediated by at Least Two Distinct Mechanisms

Michael E. Nemergut* and Ian G. Macara‡

*Department of Microbiology and ‡Department of Pharmacology, Markey Center for Cell Signaling, University of Virginia, Charlottesville, Virginia 22908

Abstract. RCC1, the only known guanine-nucleotide exchange factor for the Ran GTPase, is an ∼45-kD nuclear protein that can bind chromatin. An important question concerns how RCC1 traverses the nuclear envelope. We now show that nuclear RCC1 is not exported readily in interphase cells and that the import of RCC1 into the nucleoplasm is extremely rapid. Import can proceed by at least two distinct mechanisms. The first is a classic import pathway mediated by basic residues within the NH₂-terminal domain (NTD) of RCC1. This pathway is dependent upon both a preexisting Ran gradient and energy, and preferentially uses the importin-α3 isoform of importin-α. The second pathway is not mediated by the NTD of RCC1. This novel pathway does not require importin-α or importin-β or the addition of any other soluble factor in vitro; however, this pathway is saturable and sensitive only to a subset of inhibitors of classical import pathways. Furthermore, the nuclear import of RCC1 does not require a preexisting Ran gradient or energy. We speculate that this second import pathway evolved to ensure that RCC1 never accumulates in the cytoplasm.

Key words: nuclear transport • nuclear pore complex • importin • DNA-binding protein • permeabilized cells

Introduction

The exchange of macromolecules between the nucleus and cytoplasm is an essential biological activity in all eukaryotic cells (reviewed in Mattaj and Englmeier, 1998; Melchior and Gerace, 1998; Pemberton et al., 1998; Görlich and Kutay, 1999; Nakielny and Dreyfuss, 1999). Unlike the largely unidirectional transport processes of other organelles, nuclear transport is a bidirectional process. All nuclear transport is believed to occur through the nuclear pore complex, a large (∼125 MD) glycoprotein complex that contains ∼50–100 different proteins in mammals (reviewed in Davis, 1995). The nuclear pore contains an ∼9-nm central channel that is believed to permit the passive diffusion of ions, small molecules, and proteins <∼50 kD (reviewed in Stoffler et al., 1999). Molecules above this diffusion limit require facilitated mechanisms to traverse the nuclear envelope.

Facilitated nuclear import and export are signal-mediated processes. The first such signal to be identified was a nuclear localization signal (NLS), which consists of a short string of basic amino acids, designated as either monopartite or bipartite by the number of basic clusters (K alderon et al., 1984; D ingwall and L askey, 1991). Examples of NLS sequences are widespread in nature. The monopartite NLS from the SV-40 large T antigen and the bipartite NLS from nucleoplasmin are two of the most thoroughly studied. Importin-α (also called karyopherin-α, Srp1p, p56, PTAC 58, and pendulin) was found to bind directly to both types of NLS sequences and mediate their import (Adam and Gerace, 1991; Cortes et al., 1994; Cuomo et al., 1994; G örl ich et al., 1994; K ussel and F rasch, 1995; M oroianu et al., 1995; W eis et al., 1995). In the cytoplasm, importin-α forms a stable complex with the transport receptor importin-β (also called p97, karyopherin-β, and Kap95; C hi et al., 1995; E nenkel et al., 1995; R adu et al., 1995; G örl ich et al., 1996a; W eis et al., 1996). Importin-β is able to associate with the nuclear pore complex and transport NLS/importin-α complexes into the nucleoplasm (G örl ich et al., 1995; M oroianu et al., 1995), where the trimeric import complex is dissociated (M oore and B lobel, 1993; R exach and B lobel, 1995; G örl ich et al., 1996; M or oianu et al., 1996).
Although the genome of the budding yeast, *Saccharomyces cerevisiae*, contains only a single importin-α gene (SRP1) (Kussel and Frasch, 1995), duplication has occurred in metazoans, and mammals possess at least six importin-α isoforms (Cortes et al., 1994; Cuomo et al., 1994; Köhler et al., 1997; Seki et al., 1997; Takeda et al., 1997; Nachury et al., 1998). All of the isoforms bind the SV-40 NLS with similar affinity but show distinct preferences to other basic NLS sequences (Nadler et al., 1997; Seki et al., 1997; Prieye et al., 1998; Köhler et al., 1999; Welch et al., 1999). The structural basis for this preference remains obscure. In contrast to importin-α, the budding yeast genome contains 14 members of the importin-β (karyopherin-β) family, of which nine are known to mediate import and five export; mammals express at least 23 members of this family. (Görlich, 1997; Wozniak et al., 1998; Görlich and Kutay, 1999).

Importin-β family members are also able to transport proteins into the nucleus independently of importin-α (Siomi and Dreyfuss, 1995; Weighardt et al., 1995; Görlich et al., 1997; Pollard et al., 1996; Weis et al., 1996; Jakel and Görlich, 1998; Jakel et al., 1999). In these cases, importin-β binds directly to the import substrate and transports it into the nucleoplasm as a binary importin-β/import substrate complex.

Over the past decade, a wealth of experimental evidence has shown that much of the traffic across the nuclear envelope is governed by the small GTPase Ran (reviewed Mattaj and Englmeier, 1998; Pemberton et al., 1998; Görlich and Kutay, 1999; M acara et al., 2000). Ran is a highly abundant (~10^7 copies/cell), ~25-kD protein that, at steady state, is ~80% nuclear (Bischoff and Ponstingl, 1991b). Like other small GTPases, Ran cycles between GTP- and GDP-bound states. Unlike other GTPases, however, Ran possesses only a single known guanine-nucleotide exchange factor, RCC1 (Bischoff and Ponstingl, 1991a), and a single known GTPase-activating protein, RanGAP (Bischoff et al., 1994). RCC1 and RanGAP possess distinct subcellular localizations. While RCC1 is a nuclear protein (Ohtsubo et al., 1989), RanGAP is found both free in the cytoplasm and attached to the cytoplasmic face of the nuclear pore (Hopper et al., 1990; M atunis et al., 1996; M ahaban et al., 1997; Saitoh et al., 1997). The disparate localizations of these two proteins are believed to create a nuclear compartment with a high concentration of Ran-GTP, while Ran within the cytoplasm is predominantly in the GDP-bound state. This gradient of Ran-GTP across the nuclear envelope is crucial for most forms of nuclear transport (M attaj and Englmeier, 1998; Pemberton et al., 1998; Görlich and Kutay, 1999; M acara et al., 2000).

The prevailing model for the role of Ran in nuclear transport asserts that eukaryotic cells use Ran to control the loading and unloading of cargo in a compartment-specific manner. Biochemically, import complexes can only be formed in the absence of Ran-GTP (R exach and Blo bel, 1995; Chi et al., 1996; Görlich et al., 1996b). Thus, the relative absence of Ran-GTP in the cytoplasm allows for the formation of a stable import complex. These complexes can then translocate into the nucleus where they encounter high levels of Ran-GTP. The binding of Ran-GTP to the import receptor dissociates the receptor-cargo complex, thereby unloading cargo into the nucleoplasm (M oore and Blobel, 1993; Görlich et al., 1996b). This model proposes that the gradient of Ran-GTP across the nuclear envelope is the primary driving force for the directionality of Ran-dependent transport.

Since nuclear transport is an essential biological activity in eukaryotic cells, understanding how the Ran gradient is established and maintained is of obvious interest. One intriguing question is how cells preserve the nuclear localization of RCC1. If nuclear RCC1 is required to establish the Ran gradient, can RCC1 be imported solely by a Ran-dependent mechanism? A recent study has shown that importin-α3 is able to stimulate the import of RCC1 into digitonin-permeabilized cells in vitro (Köhler et al., 1999). It is not known, however, whether there are other, additional transport mechanisms for RCC1.

RCC1 is a nuclear protein of ~45 kD that can bind chromatin (Ohtsubo et al., 1989). The crystal structure for RCC1 has been solved and revealed a seven-bladed propeller (Renaud et al., 1998), reminiscent of the structure for the β subunit of heterotrimeric G-proteins (Wall et al., 1995). Mutational analysis suggests that RCC1 has discrete Ran- and DNA-binding surfaces (Seino et al., 1992; A zuma et al., 1996; R enaund et al., 1998). Mutation of a single aspartic acid residue to alanine (D182A) renders the protein unable to interact stably with Ran and catalyze nucleotide exchange (A zuma et al., 1996, 1999). The first 25 amino acids of RCC1 comprise its NH_2-terminal domain (NTD). The NTD is primarily unstructured, but it is located on the putative DNA-binding surface (Renaud et al., 1998). The NTD harbors two clusters of basic amino acids that each resemble a monopartite NLS (see Fig. 1 A). Mutational analysis of this domain has implicated the NTD in both nuclear transport and DNA binding (Seino et al., 1992); however, a definitive role for this domain remains to be defined.

In this study, we show that the NTD is a bona fide transport sequence for RCC1. Furthermore, we demonstrate that the NTD confers high affinity binding to importin-α3 and access to a Ran-dependent import pathway. Surprisingly, deletion of the NTD from RCC1 abrogated its ability to bind importin-α, but it did not prevent the protein from entering the nucleoplasm. In addition, many treatments that inhibit the transport of Ran-dependent import substrates do not prevent the import of RCC1 constructs that are above the passive-diffusion limit of the nuclear pore. We conclude that RCC1 is imported by at least two different mechanisms. The first is a classic, Ran-dependent import pathway mediated by the NTD of RCC1. The second is a previously uncharacterized pathway that is saturable and temperature-sensitive but requires neither energy, soluble factors, nor a preexisting Ran gradient. We speculate that this second pathway evolved as a mechanism to scavenge RCC1 molecules and prevent any accumulation of Ran-GTP in the cytoplasm.

### Materials and Methods

**DNA Constructs**

The cDNA encoding the full-length RCC1 protein was inserted into pGEX-2T (Pharmacia) as a BamHI-BamHI fragment. Creation of
the RCC1 deletion constructs was accomplished by PCR amplification of the region encoding residues 23–421 and ligation into both pGEX-2T and pGEX-GFP vectors. A BamH1-I fragment from pGEX2T (provided by Dirk Görlich) was amplified by PCR and ligated to a BamH1-BamHI fragment into pGEX-GFP that had been digested BamH1-BglII, thereby removing the GFP and destroying the 3’ restriction site. RCC1 was then ligated into pGEX-zz as a BamH1-BamHI fragment. The CDNA encoding the first 25 amino acids of RCC1 was PCR-amplified and ligated into pQE-70 nucleoexpression core-domain (NPC) (Dirk Görlich) as a Splh-Splh fragment. All constructs were checked by automated sequencing.

**Protein Expression, Purification, and Labeling**

GST-fusion proteins were expressed in E.cherichia coli XL-1-Blue or BL21 essentially as described previously (Seino, 1992; Welch, 1999). The proteins were bound to glutathione-Sepharose beads (Pharmacia), eluted with glutathione, and exchanged into thrombin cleavage buffer (50 mM Tris, pH 7.5, 150 mM NaCl, and 2.5 mM CaCl2) using a PD10 column (Pharmacia). The protein was cleaved with 4u thrombin; the GST and thrombin were then removed by binding to glutathione-Sepharose and p-aminobenzamide beads (Sigma), respectively. The recombinant protein was concentrated using a Centricon 30 (A micon) and analyzed by SDS-PAGE.

RCC1(1-25)-NPC-His6 was expressed in E. coli BL21 and purified by binding to nickel-agarose beads (Q-lagen; Görlich, 1996). The protein was then eluted with 200 mM imidazole, exchanged into thrombin cleavage buffer using a PD10 column (Pharmacia), concentrated using a Centricon 30 (A micon), and analyzed by SDS-PAGE.

A labeling reactions occurred in thrombin-cleavage buffer. Protein samples were labeled with tetramethylrhodamine-5-maleimide (TMRM; Molecular Probes) at a 1:1 molar ratio for 60 min on ice. A fiber quenching unreacted TMRM with 50 mM 2-mercaptoethanol (Sigma), the proteins were separated from unreacted label and exchanged into microinjection buffer (10 mM NaHPO4, pH 7.2, 70 mM KCl, and 1 mM MgCl2) using a Centrisep column (Perinton Separations).

**Cell Culture**

All cells were passaged in Dulbecco’s modified eagle medium (supplemented with 10% fetal calf serum [vol/vol] and penicillin/streptomycin). Baby hamster kidney cells (BHK 21) and HEK 293 cells were cultured in a humidified, 37°C, 5% CO2 incubator. tsBN2 cells were grown at 33.5°C and, where indicated, were temperature-shifted to 39.5°C for 3 h. To energy-deplete BHK 21 cells, cultures were washed twice in PBS, then placed into serum-free, glucose-free Dulbecco’s minimal essential medium containing 10 mM sodium azide (Sigma) and 6 mM 2-deoxy-o-glucose (Sigma) for 3 h before experimentation.

**Microinjection Studies**

Cells were cultured on CELLlocate gridded coverslips (Eppendorf). The samples were placed in Ringer’s solution (25 mM Hapes, pH 7.2, 110 mM NaCl, 5 mM KCl, 2 mM CaCl2, 1 mM MgSO4, 1 mM KH2PO4, 1 mg/ml BSA, and 0.2% glucose) before microinjection. When injecting energy-depleted cells, glucose-free Ringer’s solution was used that was supplemented with metabolic inhibitors. When injecting temperature-shifted tsBN2 cells, the Ringer’s media was supplemented with 10 mM glycol-heximide to prevent the resynthesis of endogenous RCC1. GGNLS, RCC1(1-25)-NPC, and NPC were injected at 2 mlg/ml; Ran(G19V) was injected at 4 mlg/ml. All other samples were injected at 20 mlg/ml. Recombinant protein stocks were injected using the Eppendorf 5242 apparatus, and either a FITC- or TRITC-labeled dextran (Sigma) was used as an injection site marker (1 mg/ml). Samples were incubated for 15–20 min before fixation in 4% paraformaldehyde in pH 7.2 PBS (provided by Dirk Görlich) was quenched in PBS, then washed and stained with the appropriate dye and visualized by fluorescence microscopy using a Nikon microscope with a 60x water immersion lens. Images were captured with a Hamamatsu CCD camera using Openlab software (Improvision).

**Nemergut and Macara Nuclear Import of RCC1**

When the import of proteins was assayed by time-lapse photography, cells were injected in ice-cold Ringer’s solution as described above. The plate was washed quickly with 37°C Ringers before initiating the time-lapse program (~1 min) and images were captured at 1-min intervals. The sample was maintained at the appropriate temperature using a heated stage apparatus. The change in mean nuclear fluorescence intensity was measured using the Openlab software. All time-lapse images were captured as 20x images using the apparatus described above.

To determine whether RCC1 shuttles between the nucleus and cytoplasm, BHK21 cells were cultured on CELLocate coverslips as described above. Single nuclei of multinucleate cells were injected with TMRM-like import receptor binding domain (BIB; Jakel and Görlich, 1998; Welch et al., 1999). The importin-α-like lysate was prepared by resuspension in lysis buffer (50 mM Tris, pH 8.1, 1 mM EDTA, pH 8, 1% Triton X-100, 150 mM NaCl, and 1 mg/ml lysosome), sonication, and clearance by centrifugation. Equal amounts of the GST-fusion proteins were prebound to glutathione-Sepharose beads (Amersham Pharmacia) and the bacterial lysate was added in binding buffer (20 mM MOPS, pH 7.1, 100 mM KOAc, 5 mM MgOAc, 0.1% Tween-20, 5 mM DTT, and 0.5 mM PM SF). Binding reactions were incubated for 2.5 h at 4°C in an Eppendorf thermomixer. The supernatant was sampled to ensure that the beads were incubated with equal concentrations of importin-α. A fiter washing (3×) in binding buffer, the beads were resuspended in Laemmli sample buffer and boiled. The proteins were subjected to SDS-PAGE and transferred to nitrocellulose. Importin-α was then visualized using an anti-His antibody (1:1,000; BabcO), a HRP-coupled secondary antibody (1:20,000; The Jackson Laboratory), and a chemiluminescence reagent (KPL).

**In Vitro Import Assays**

We used a modified protocol from that described originally by A dam et al. (1990). HEK293 cells were cultured on poly-L-lysine-coated coverslips to 50–70% confluence. A fiter washing in ice-cold PBS, the cells were permeabilized with digitonin (0.005%; Calbiochem) in import assay buffer (20 mM Hepes-KOH, pH 7.5, 80 mM KOAc, 4 mM MgOAc, 2 mM DTT, and 250 mM sucrose) for 5 min on ice. Permeabilization was stopped by washing samples in ice-cold buffer. Permeabilized 1% importin-α assay buffer for 5 min at 25°C. A standard import assay consisted of the import substrate (0.6–3 μM), a rabbit reticulocyte lysate (50% vol/vol), and an energy-regenerating system (20 mM phosphocreatine, 1 mM ATP, 1 mM GTP, 20 U/ml creatine phosphokinase) in 1% BSA/importassay buffer. Import reactions were carried out for 25 min for GG-NLS and 5–10 min for the RCC1 constructs. Where indicated, inhibitors were used at the following concentrations: 0.2 mg/ml wheat-germ agglutinin, 100 μM β-mercaptoethanol receptor binding domain (BIB; Jakel and Görlich, 1998; Welch et al., 1999), 6 μM GST-importin-β(45-462), and 36 μM RCC1. Each import reaction was terminated by washing (3×) in import assay buffer. The cells were then fixed in 4% PFA (2% sucrose/PBS, permeabilized in methanol (~20°C), stained with DAPI, and visualized by fluorescence microscopy.

To assess the integrity of the nuclear envelope, import assays were done on digitonin-permeabilized HEK293 cells using RCC1(zz) as an import substrate, as described above. All samples were fixed in 4% PFA (2% sucrose/PBS; however, the methanol permeabilization step was omitted in some samples. All samples were then blocked in 5% BSA/PBS before visualization of the RCC1(zz) by immunofluorescence.

**Chromatin-Binding Assay**

HEK293 cells were cultured on poly-L-lysine coverslips and permeabilized in 0.2% Triton X-100/import assay buffer (vol/vol) for six min at 25°C. RCC1, RCC1-GFP, or RCC1(zz) were added directly to the coverslips for 5 min. The cells were then washed, fixed, stained, and visualized as described previously.
Immunofluorescence

HeLa cells were fixed in 4% PFA/2% sucrose/PBS for 15 min at room temperature and, where indicated, permeabilized in methanol (−20°C) for 2 min. Samples were blocked in 5% BSA/PBS overnight at 4°C. RCC1zz was visualized directly using Texas red-conjugated rabbit IgG (The Jackson Laboratory; 1:1,000) for 60 min at 25°C. Nuclei were stained concomitantly with DAPI. Samples were then washed (5×) and analyzed by fluorescence microscopy.

To visualize endogenous Ran, tsBN2 cells were incubated at 33.5°C or 39.5°C for 3 h and then fixed, permeabilized, and incubated with a monoclonal anti-Ran antibody (Transduction Laboratories; 1:250) for 45 min at 25°C. A secondary, Texas red-conjugated, anti–mouse IgG (The Jackson Laboratory; 1:1,000) was used for 35 min at 25°C. Nuclei were stained concomitantly with DAPI.

Results

The NTD of RCC1 Contains a Functional NLS but Is Not Required for Transport

RCC1, the only known guanine-nucleotide exchange factor for the Ran GTPase (Bischoff and Ponstingl, 1991a), is a nuclear, chromatin-bound protein (Ohtsubo et al., 1989). The NTD of RCC1 is primarily unstructured, but it is located on the putative DNA-binding surface of RCC1 (Renault et al., 1998). It possesses two clusters of basic amino acids that each resemble a monopartite NLS (Fig. 1 A) and has been implicated in both nuclear transport and DNA binding (Seino et al., 1992). To test the functionality of the NTD in nuclear transport, a fusion protein was generated using the NTD of RCC1 and the nucleoplasmin core domain (RCC1(1-25)-NPC); this protein, labeled with TMRM, spontaneously forms pentamers (Görlich et al., 1996a; ~100 kD) that are well above the passive diffusion limit of the nuclear pore (~50 kD; Stoffler et al., 1999). GST-GFP-NLS (GGNLS), a fluorescent protein substrate containing a classic NLS from the SV-40 large T antigen (Kalden et al., 1984; Welch et al., 1999), was used as a positive control; the nucleoplasmin core protein itself was used as a negative control (Fig. 1 B). When injected into the cytoplasm of BHK21 cells, RCC1(1-25)-NPC, but not NPC alone, concentrated rapidly within the nucleoplasm, demonstrating that the NTD of RCC1 contains a functional NLS.

These data showed that the NTD was sufficient to promote the nuclear localization of a fluorescent reporter protein. To test whether the NTD was necessary for nuclear translocation of RCC1, we created RCC1-GFP fusion proteins in which portions of the NTD had been deleted. Since the removal of the first 10 amino acid residues of RCC1 had no detectable effect on the ability of RCC1 to traverse the nuclear envelope (data not shown), we constructed a GFP-fusion protein in which almost the entire NTD was deleted (RCC1(23-421)-GFP). RCC1 and RCC1-GFP were used as controls. When injected into the cytoplasm, full-length RCC1, RCC1-GFP, and RCC1(23-421)-GFP were all imported rapidly into the nucleus (Fig. 1 C). Therefore, the nuclear import of RCC1 does not require its NTD. Since RCC1(23-421)-GFP (~70 kD) is above the diffusion limit of the nuclear pore complex, it seemed unlikely that import of this molecule was due to passive diffusion. Taken together, these data imply that RCC1 possesses at least two independent import pathways.
To explore the transport dynamics of RCC1, we next asked whether RCC1 shuttles continuously between the nucleus and cytoplasm. To investigate this possibility, we injected single nuclei of multinucleate BHK21 cells (Herold et al., 1998). RCC1 did not equilibrate between the nuclei within the cells over a period of 35–40 min (Fig. 1 D), suggesting that RCC1 is not exported readily from the nucleus in interphase cells. Use of heterokaryon fusion assays also demonstrated that RCC1 does not shuttle (Lindsay, M., personal communication). Since RCC1 (~45 kD) is below the theoretical diffusion limit of the nuclear pore (~50 kD), these data suggest that RCC1 is efficiently retained within the nucleoplasm, presumably by binding to chromatin.

The nuclear import of RCC1 in vitro was recently reported to be stimulated by importin-α, and preferentially by the α3 isoform (Köhler et al., 1999). Since the NTD of RCC1 contains a NLS (Fig. 1 B), it seemed likely that this sequence would be necessary to confer binding to importin-α. To determine which RCC1 constructs could bind members of the importin-α family, we immobilized recombinant GST-RCC1 fusion proteins onto glutathione-Sepharose beads. We then tested their ability to bind individual importin-α family members from an E. coli lysate. Bound and unbound fractions were visualized by Coomassie staining (data not shown) and immunoblotting against the His6-tag contained on the COOH terminus of each importin-α family member (Fig. 2). GGNLs, a protein that binds to all known importin-α members tested to date, was used as a positive control (Welch, 1999). While GST-RCC1 was able to bind both importin-α3 and importin-α1, binding to importin-α3 was much more robust (Fig. 2). By Coomassie staining, GST-RCC1 was able to bind nearly equimolar amounts of importin-α3 while binding to importin-α1 was undetectable (data not shown). By immunoblotting, visualization of importin-α1 required an ~12-fold longer exposure relative to that of importin-α3. These data are consistent with the specificity imparted by importin-α3 on the import of RCC1 in vitro (Köhler et al., 1999). The affinity of GST-RCC1(23-421) for these importin-α family members was too low to be detectable (Fig. 2). However, as described above, RCC1(23-421)-GFP is imported into nuclei in intact cells. These data therefore suggest that RCC1 can use both importin-α-dependent and -independent import pathways in vivo.

Transport Kinetics of RCC1

To estimate the rate of entry of RCC1 into the nucleus in vivo, we injected the cytoplasm of tsBN2 cells with the RCC1 constructs and visualized entry into the nucleus by time-lapse photography. GGNLs, a substrate known to be imported in a Ran- and energy-dependent manner, was used as a control (Welch et al., 1999). The rate of import is plotted in Fig. 3, shown below time-lapse stills from individual cells. When GGNLs was injected, we observed a steady increase in nuclear fluorescence that continued for ~30 min. Nuclear import of the RCC1 constructs, however, was significantly different. When RCC1 was injected, almost all of the cells had concentrated the protein within their nuclei before the time-lapse program could be initiated (~1 min, data not shown). RCC1-GFP and RCC1(23-421)-GFP both displayed rapid import rates. In cells injected with RCC1-GFP, ~50% of the cells had concentrated the protein within their nuclei before the time-lapse program was initiated (data not shown); the remaining cells imported RCC1-GFP within ~5 min (Fig. 3). Import of RCC1(23-421)-GFP also occurred rapidly (see Discussion). Furthermore, the early phase of import of all RCC1 constructs tested, the injected protein concentrated at the periphery of the nucleus (Fig. 3, middle panels), suggesting that RCC1 may dock at the nuclear pore more rapidly that it is translocated into the nucleoplasm. A further possibility is that these rings represent the binding of RCC1 to chromatin just under the surface of the nuclear envelope (see Discussion). Shortly after this peripheral accumulation, the protein became almost entirely nuclear within ~2 min. GGNLs did not display this phenotype.

Nuclear Import of RCC1 Does Not Require a Ran Gradient In Vivo

The nuclear import of many proteins has been shown to require a gradient of Ran-GTP across the nuclear envelope. Since RCC1(23-421)-GFP is unable to bind importin-α but is still capable of nuclear import (Figs. 2 and 3), an intriguing question was whether the import of RCC1 was a Ran-dependent process. To address this issue in vivo, we artificially collapsed the Ran gradient by coinjecting the various RCC1 constructs with Ran(G19V). This mutant protein, shown previously to be predominantly in the GTP-bound state, harbors a single amino acid substitution that renders the protein GTPase-deficient and insensitive to both RanGAP and RCC1-catalyzed nucleotide exchange (Lounsbury, 1996). Mammalian cells harbor ~10 million Ran molecules, and ~10–20% are in the cytoplasm at steady state (Bischoff and Ponstingl, 1991). By injecting the cytoplasm with one-tenth the cellular volume (~5 × 10^-13 L) of 40 μM Ran(G19V), ~12 million G19V molecules are injected, an amount that should be sufficient to collapse the Ran gradient in vivo. GGNLs, a protein known to require a Ran gradient for import, was used as a control for Ran gradient collapse (Welch et al., 1999). As...
shown in Fig. 4, the coinjection of Ran(G19V) completely prevented the import of both GGNLS and RCC1(1-25)-NPC. Since RCC1(1-25)-NPC contains a NLS (Fig. 2) that is necessary to confer binding to importin-α (Fig. 3), these data suggest strongly that the NTD of RCC1 is a classic, Ran-dependent NLS imported specifically by the importin-α/β heterodimer. In contrast, the import of RCC1 and RCC1-GFP was not prevented by coinjection of Ran(G19V). Moreover, a mutant form of RCC1, D182A, which cannot interact with Ran stably nor catalyze nucleotide exchange (Azuma et al., 1996, 1999), was imported efficiently in the presence of Ran(G19V). Taken together, these data suggest that RCC1 can be imported by at least two pathways: (a) a classic Ran-dependent pathway and (b) a Ran-independent pathway that is not passive diffusion.

To further test the import of RCC1 in vivo, we conducted microinjection experiments in tsBN2 cells. These cells possess a temperature-sensitive RCC1 allele and, when cultured at 39.5°C, endogenous RCC1 protein is degraded rapidly, resulting in the loss of the Ran gradient (Fig. 5 b; Nishimoto et al., 1978; Uchida et al., 1990; Tsubo et al., 1991; Tachibana et al., 1994; Richards et al., 1997). As a result, it has been shown that the nuclear import of Ran-dependent cargo is severely compromised in temperature-shifted tsBN2 cells (Uchida et al., 1990; Tachibana et al., 1994; Richards et al., 1997). When tsBN2 cells cultured at 33.5°C were injected, GGNLS and all RCC1 constructs tested accumulated within the nucleus (Fig. 5 A). In contrast, when temperature-shifted tsBN2 cells were injected, the import of GGNLS and RCC1(1-25)-NPC was prevented completely. However, RCC1, RCC1(D182A), and RCC1-GFP all entered nuclei rapidly, further suggesting that RCC1 possesses a Ran-independent import pathway.

Since RCC1 is slightly below the diffusion limit of the nuclear pore, one possible explanation for the data described above is that small amounts of RCC1 were able to enter nuclei by passive diffusion, which might reestablish the Ran gradient and prime the import of RCC1 by a Ran-dependent pathway. The fact that RCC1-GFP, a molecule above the diffusion limit of the nuclear pore, accumulated rapidly within nuclei argues against this possibility. In addition, RCC1(D182A), a mutant that is inactive catalytically (Azuma et al., 1996), was able to accumulate within the nuclei of temperature-shifted tsBN2 cells at an efficiency comparable to the wild-type protein. One could speculate, however, that although the ability of the RCC1(D182A) mutant to catalyze nucleotide exchange is severely compromised in vitro, there may be sufficient residual activity to reestablish the Ran gradient in vivo. To test this possibility experimentally, tsBN2 cells were cultured at 39.5°C for 3 h and stained for endogenous Ran. In tsBN2 cells cultured at 33.5°C, Ran is ~80% nuclear while...
in temperature-shifted tsBN2 cells the Ran is distributed diffusely throughout the cell (Fig. 5 B, top). The Ran gradient could be reestablished by a cytoplasmic injection of wild-type RCC1 but not the mutant RCC1(D182A). The fact that RCC1(D182A) did not reestablish the Ran gradient cannot be explained by a lack of import, because it accumulates within the nuclei of temperature-shifted tsBN2 cells (Fig. 5 A). Thus, RCC1(D182A) can enter nuclei in the absence of a Ran gradient. These data demonstrate that (a) a cytoplasmic injection of RCC1 can quickly reestablish the Ran gradient, (b) the ability to reestablish the Ran gradient is dependent upon the catalytic activity of RCC1, and (c) the ability of RCC1 to be imported is independent of both the catalytic activity of RCC1 and a pre-existing Ran gradient.

Import of RCC1 into Digitonin-permeabilized Cells
We have demonstrated that, in intact cells, import of RCC1 is mediated by both Ran-dependent and -independent import pathways. To determine whether RCC1 requires specific, soluble transport factors in vitro, HeLa cells were permeabilized with digitonin (Adam et al., 1990) and the import of TMRM-RCC1 and RCC1-GFP was observed using GGNLS as a control. GGNLS is imported by a classical, temperature-dependent and factor-dependent import pathway (Welch et al., 1999). This process was inhibited by: (a) removal of the soluble factors (reticulocyte lysate); (b) addition of the pore-binding protein WGA; (c) addition of a dominant-negative fragment of importin-β (β(45-462)); (d) addition of an importin-β-binding protein fragment (BIB; Jakel and Görlich, 1998); and (e) carrying out the import reaction at low temperature (Fig. 6 A). In contrast to this classic import substrate, RCC1 and RCC1-GFP were imported rapidly under every condition tested except when the reaction was carried out at 4°C, or when β(45-462) was added. However, since the reaction did not require the addition of importin-β, and the BIB domain did not inhibit the import of RCC1, these data suggest that the importin-β-independent pathway for RCC1 is not mediated by direct binding to importin-β.

The β(45-462) fragment is known to associate almost irreversibly with nucleoporins and to inhibit a variety of different nucleocytoplasmic transport pathways (Kutay et al., 1997). The fact that β(45-462) inhibited the import of the RCC1 constructs confirms that RCC1 uses nuclear pores for transport and that its import does not occur by passive diffusion (see Discussion). To address whether treatment at 4°C was inhibiting nuclear import or retention, HeLa cells were permeabilized with Triton X-100 and the ability of RCC1 to bind chromatin was assayed directly. In Triton X-100-permeabilized cells, the binding of RCC1 and RCC1-GFP to chromatin was a temperature-insensitive event (Fig. 6 B). Thus, low temperature prevented the translocation of RCC1 across the nuclear envelope, and not the nuclear retention by binding to chromatin.

Although WGA inhibited the import of GGNLS completely, the transport of RCC1 and RCC1-GFP appeared much less sensitive. The fact that WGA did not prevent the import of RCC1 was surprising, and raised the possi-
bility that digitonin was perturbing the integrity of the nuclear envelope. Although digitonin is not expected to disrupt the nuclear envelope at the concentrations used in these experiments (0.005%), we wished to test this possibility empirically. If the nuclear envelope was disrupted, then it would allow access of an antibody into the nuclear compartment (Adam et al., 1990). Initially, we observed that the digitonin permeabilization of HeLa cells did not...
permit the access of an antibody targeted to the nuclear antigen C23 (data not shown). As a more direct approach, we designed an assay that would allow recognition of damaged nuclei using identification of the imported RCC1 itself as a marker. A RCC1 fusion protein was constructed with a COOH-terminal zz tag, which binds efficiently to the constant domain from IgG (Görlich et al., 1997). Import assays were performed on digitonin-permeabilized cells using RCC1zz (~60 kD) and GGNLS as import substrates in the presence or absence of WGA. All samples were then fixed in paraformaldehyde; however, only half of the RCC1zz samples were permeabilized with methanol. GGNLS was visualized directly; RCC1zz was stained using a Texas red-conjugated rabbit IgG (TR-IgG). As shown in Fig. 7, when cells were permeabilized with methanol, RCC1zz was detectable in nuclei regardless of whether or not WGA was included in the import reaction. If the cells were not permeabilized with methanol, only background staining was observed. Thus, the ability of TR-IgG to gain access to the nuclear compartment in digitonin-permeabilized cells was entirely dependent on additional permeabilization with methanol. As a positive control, the nuclear envelope was intentionally disrupted with Triton X-100 before incubation with RCC1zz. In these cells, RCC1zz was readily detectable in the absence of methanol permeabilization (Fig. 7, bottom). Since WGA inhibited the import of GGNLS in this assay, these data can not be explained by a defect in the activity of the WGA. Furthermore, the staining of nuclei was dependent upon the addition of RCC1zz; thus, TR-IgG recognized RCC1zz specifically and did not cross-react with endogenous RCC1 (Fig. 7, top). Taken together, these data demonstrate that RCC1 can be imported by a factor-independent pathway that it is resistant to all import inhibitors tested except low temperature and β(45-462).

To determine whether the Ran-independent accumulation of RCC1 in the nucleus was saturable, we performed a competition experiment in vitro using RCC1(23-421)-GFP as an import substrate and unlabeled RCC1 as a competitor. Since RCC1(23-421)-GFP does not possess the NTD, it can only access the Ran-independent import pathway. Furthermore, import reactions were carried out in the absence of a reticulocyte lysate so that only the Ran-independent pathway could be observed. Under these conditions, RCC1(23-421)-GFP rapidly entered the nuclei of digitonin-permeabilized cells in the absence of any competitor (Fig. 8, top). In contrast, the nuclear accumulation of RCC1(23-421)-GFP was prevented by the addition of a 60-fold molar excess of RCC1. Significantly, import of RCC1(23-421)-GFP was also inhibited by low temperature or by the addition of importin-β(45-462) (Fig. 8, bottom). These data confirm that importin-β(45-462) inhibits both the Ran-dependent and -independent import pathways of RCC1 and suggest that, in the absence of soluble factors, RCC1, RCC1-GFP, and RCC1(23-421)-GFP enter nuclei by the same saturable pathway.

**Import of RCC1 in Energy-depleted Cells**

To determine the energy requirements for the import of RCC1 in vivo, BHK cells were treated with 10 mM sodium azide and 6 mM 2′-deoxy-α-glucose to inhibit both mitochondrial and glycolytic mechanisms for ATP production. Under these conditions, it has been shown that ATP pools are depleted rapidly and reversibly (Breeuwer and Goldfarb, 1990; Weldon et al., 1998). Energy-depleted cells were microinjected with the RCC1 constructs and import of the proteins was monitored by fluorescence microscopy. In the absence of metabolic inhibitors, GGNLS, RCC1-GFP, and RCC1(1-25)-NPC were all transported rapidly into nuclei (Fig. 9). In energy-depleted cells, however, RCC1-GFP was still imported rapidly while GGNLS and RCC1(1-25)-NPC remained cytosolic. These data demonstrate that RCC1 is imported by at least two pathways. The first is mediated by the NTD of RCC1 and is a classic Ran (Figs. 4 and 5) and energy-dependent (Fig. 9) pathway. The second pathway is both saturable and sensitive to temperature, but requires neither energy, soluble factors, nor a Ran gradient.

**Discussion**

The protein RCC1 possesses at least two distinct biological functions. The first and best-established function is to generate a Ran-GTP gradient across the nuclear pores (Bischoff and Ponstingl, 1991), which is used to drive the transport of a wide variety of cargo molecules against their concentration gradients (Mattaj and Englmeier, 1998; Melchior and Gerace, 1998; Pemberton et al., 1998; Görlich and Kutay, 1999; Nakielny and Dreyfuss, 1999; Macara, 2000). A second, more recently characterized function is to create a Ran-GTP gradient near the chromatin surface, which is required for mitotic spindle formation (Carazo-Salas, 1999; Ohba et al., 1999; Wilde and Zheng, 1999). In both cases, RCC1 behaves as a sort of identification tag for chromatin and hence, in interphase cells, as a marker for the nuclear compartment. It operates by efficiently catalyzing guanine-nucleotide exchange on the Ran GTPase (Bischoff and Ponstingl, 1991). The amount of RCC1 associated with the chromatin must double with each cell cycle. One can imagine two extreme mechanisms to achieve this result. In the first mechanism, RCC1 is only synthesized in S-phase, and must await breakdown of the nuclear envelope in prophase to associate with the chromatin. This mechanism would be problematic for cells such as those of the budding yeast, Saccharomyces cerevisiae, that use a closed mitotic cycle. In the second mechanism, RCC1 is synthesized throughout the cell cycle and is imported continuously into the nucleus, either by passive diffusion or by a facilitated process.

Although the size of RCC1 (~45 kD) is such that it could, in principle, diffuse passively through the nuclear pores, the rate of diffusion, based on comparison with ovalbumin, which is of similar size (Featherstone et al., 1988), is likely to be too low to be practicable. Also, accumulation of newly synthesized RCC1 within the cytoplasm would likely reduce the efficiency of nucleocytoplasmic transport and disrupt microtubule dynamics formation. Therefore, one might predict that RCC1 is imported by a facilitated mechanism throughout interphase. In support of this prediction, a conserved polybasic sequence that resembles a classic NLS is found near the NH₂ terminus of RCC1 from many species (Fig. 1 A). Attachment of this NTD to β-galactosidase permits accumulation of the fu-
sion protein within nuclei of transiently transfected COS 7 cells (Seino et al., 1992). A iso, a study of multiple importin-α isoforms showed that RCC1 can accumulate within the nuclei of digitonin-permeabilized cells in the presence of energy and soluble transport factors, and that accumulation was enhanced specifically by the importin-α3 isoform (Köhler et al., 1999). These studies suggest that the import of RCC1 can be mediated by the classical importin-α/β pathway through a conventional NLS. However, these studies did not define the NH2-terminal sequence as the

signal recognized by importin-α3. The NTD has also been reported to bind DNA (Seino et al., 1992), thus complicating analysis of the transfection data for the NTD-β-galactosidase construct described above. Because the transfected cells had undergone at least one cell cycle (Seino et al., 1992), one could imagine that the NTD caused binding of the fusion protein to exposed chromatin during mitosis, and hence mediated trapping within nuclei rather than import through the nuclear pores. Moreover, an NH2-terminal deletion mutant of RCC1 was not excluded from the nuclei of transfected tsBN2 cells and was capable of rescuing these cells at the nonpermissive temperature (Seino et al., 1992). Either the deletion mutant could diffuse into the nuclei, where it would be retained by binding to chromatin, or there exists a second transport system that is independent of the putative NH2-terminal NLS.

Given the physiological importance of RCC1 import, we examined the mechanism by which it is translocated into the nucleus using a combination of intact and permeabilized cell assays. We find that the NTD of RCC1 is necessary for binding importin-α and that RCC1 preferentially recognizes the α3 isoform. Attachment of the first 25 amino acids from RCC1 are also sufficient to target a fluorescent reporter protein to the nucleus upon microinjection into the cytoplasm. The NTD of RCC1 is not sufficient, however, to target proteins to chromatin in vitro (data not shown). Furthermore, we show that NTD-mediated transport is dependent upon both a preexisting Ran gradient and energy in vivo. These data confirm that the NTD of RCC1 is a NLS and not a DNA-binding domain.

By time-lapse photography, we found that almost all of the cells concentrated the injected RCC1 protein within their nucleus before we could initiate the time lapse program (<1 min). Assuming that (a) each cell was injected with ~5 × 10^-13 L of substrate, (b) there are ~3,000 nu-
clear pores per nucleus, and (c) ~80% of the injected protein became nuclear within one minute, we can estimate that ~72,000 RCC1 molecules were imported per second (~24 events/pore/s). To our knowledge, this is one of the most rapid nuclear import process recorded to date. In fact, kinetic analysis of RCC1 demonstrated that its import was at least 30-fold greater than the import of a classic NLS-containing import substrate (GGNLS) under similar conditions. Since RCC1 and GGNLS can bind to importin-α3 with similar affinity (Fig. 3), each can access the importin-α/b import pathway; however, GGNLS can both bind and be imported by every importin-α molecule tested to date while RCC1 can not (Köhler et al., 1999; Welch et al., 1999). The fact that RCC1 is imported much more rapidly than GGNLS thus suggests that RCC1 possesses another import pathway that is independent of that mediated by importin-α3.

Removal of the NTD from RCC1 abolished binding to importin-α but did not prevent the RCC1 deletion mutant from entering the nucleoplasm (Fig. 1 C). Kinetic analysis of RCC1(23-421)-GFP in vivo is complicated by the fact that a subset of cells did not import the protein within the time frame of the time-lapse experiment; analysis of those cells that imported, however, showed that the rate was indistinguishable from that of RCC1-GFP. The fact that not all of the cells import RCC1(23-421)-GFP within 25 min suggests that the import might be a regulated process. Efforts to either increase or decrease the import of RCC1(23-421)-GFP have only been successful by treatment at low temperature, the addition of importin-β(45-462), and by competition with excess, unlabeled RCC1 in vitro (Fig. 8, data not shown). The heterogeneity of import may also reflect differences in individual cells to import under time-lapse conditions; however, since cells that did not import RCC1(23-421)-GFP were excluded from kinetic analysis, the overall mean import rate for RCC1-GFP in vivo must be more rapid than for RCC1(23-421)-GFP. This observation, plus the fact that the NTD of RCC1 confers high-affinity binding to importin-α3, suggests that the NTD can facilitate the import of RCC1 in vivo.

Figure 7. Import assays were accomplished on digitonin-permeabilized HeLa cells. Reactions consisted of GGNLS (0.2 mg/ml; top) or RCC1zz (0.7 μM; bottom) as import substrates, an energy regenerating system, and a reticulocyte lysate. Where indicated, 0.2 mg/ml WGA was added. All samples that used GGNLS as an import substrate were permeabilized with methanol and stained with a Texas red-conjugated rabbit IgG (TR-IgG; top, right panels). When RCC1zz was used as the import substrate, half of the samples were permeabilized with methanol (+MeOH) while half were not (−MeOH). RCC1zz was then visualized in all samples using a Texas red-conjugated rabbit IgG. When Triton-permeabilized cells were used (bottom, left panels), the permeabilization with methanol was omitted. All nuclei were visualized with DAPI. Bar, 10 μm.
Interestingly, all of the RCC1 constructs that were tested accumulated at the nuclear periphery before import (Fig. 3). In our hands, the import of molecules by classic transport pathways, such as GGNLS (Fig. 3) and RCC1 (1-25)-NPC (data not shown), do not show this phenotype. A similar accumulation at the nuclear envelope was observed during the import of the RCC1 constructs into the nuclei of temperature-shifted tsBN2 cells (data not shown); therefore, this phenotype is not due to import by a Ran-dependent process. This phenomenon could represent either RCC1 that is bound either directly or indirectly to the nuclear pore, or RCC1 that is binding to chromatin just as it enters the nucleus. The latter hypothesis, however, predicts that accumulation of RCC1 in the nucleus would proceed from the nuclear periphery to the nuclear center, i.e., that the ring of RCC1 would thicken over time, and this process is not observed. If RCC1 is docking at the nuclear pore, then this may be the rate-limiting step in the import of RCC1. The fact that the import of RCC1 was inhibited by importin-β(45-462) suggests that either RCC1 binds to some of the same nucleoporins as importin-β or that access to RCC1-binding sites on nuclear pores is obscured by importin-β. The identity of the putative nucleoporin docking sites for RCC1 remains to be investigated. It is noteworthy that staining of RCC1 at the nuclear periphery was not prominent in digitonin-permeabilized cells.

Figure 8. Import of RCC1 into nuclei is saturable. Digitonin-permeabilized HeLa cells were incubated with RCC1(23-421)-GFP as an import substrate (0.6 μM). All reactions were done in the absence of a reticulocyte lysate (−RL) but in the presence of an energy regenerating system. Import was accomplished in the absence or absence of a 60-fold molar excess of unlabeled RCC1. Nuclei were visualized with DAPI (top left panels). Where indicated, GST-importin-β(45-462) (6 μM) was added; a reaction incubated on ice is also shown. Bar, 10 μm.

Figure 9. Import of RCC1 into nuclei does not require energy in vivo. BHK21 cells were incubated in serum- and glucose-free media with (right panels) or without (left panels) sodium azide and 2′deoxy-D-glucose for 3 h. The indicated protein substrates were coinjected with a fluorescent dextran as an injection site marker at concentrations identical to that described previously. Samples were incubated 15 min before fixation. Bar, 10 μm.
incubated with RCC1 at 4°C. Therefore, either binding is of very low affinity or access to the docking site is reduced at low temperatures. Low-affinity binding to nucleoporins is also a characteristic of NTF2, which has an ~25 μM affinity for nucleoporin-FxFG repeats (Chaillan-Huntington et al., 2000).

A surprising feature of the factor-independent RCC1 import pathway is that, compared with GGNLS, it is much less sensitive to inhibition by WGA. WGA binds with high affinity to O-linked glycoproteins of the nuclear pore (Stöffler et al., 1999), including the p62 complex which is localized near the central gated channel of the pore (Guan et al., 1995). WGA is presumed not to occlude the channel because it does not abrogate the passive diffusion of small dextrans (Finlay et al., 1987). For this reason, WGA has been regarded as a marker to distinguish active from passive transport. However, different transport pathways use different nucleoporins (Shah and Forbes, 1998; Shah et al., 1998; Moy and Silver, 1999; Seedorf et al., 1999), and WGA may inhibit selectively those pathways that require the p62 complex. RCC1 may represent the first member of a class of proteins that are translocated across the nuclear pore complex independently of interactions with O-linked glycoproteins.

In digitonin-permeabilized cells, the import of RCC1 could only be inhibited by importin-β(45-462), low temperature, or competition with excess, unlabeled RCC1. The fact that the import reaction was inhibited by low temperature suggests strongly that import does not proceed by passive diffusion, and that the nuclear envelope was not disrupted during permeabilization of the plasma membrane (Fig. 6 B). The inhibition due to excess RCC1 could be explained by competition for a saturable transport process and/or competition for chromatin-binding sites. Evidence from time-lapse photography, however, suggests that the rate-limiting step for the import of RCC1 in vivo is transit across the nuclear pore.

Interestingly, the process by which RCC1 traverses the nuclear envelope does not require energy in vivo (Fig. 9). These data cannot be explained by the assertion that energy depletion alters the permeability of the nuclear envelope, because neither the injection site marker, GGNLS, nor RCC1(1-25)-NPC entered the nucleoplasm. Furthermore, since RCC1-GFP is beyond the diffusion limit of the nuclear pore, import in energy-depleted cells can not proceed by passive diffusion. Energy-independent translocation of this sort is not without precedent. Evidence from in vitro import assays suggest that single rounds of transport mediated by transportin, Crm1, and importin-β are energy independent (Schwoebel et al., 1998; Engelmeier et al., 1999; Ribbeck et al., 1999). In addition, proteins such as β-catenin (Yokoya et al., 1999) and hnRNPK (Michael et al., 1997) are able to translocate across the nuclear envelope in the absence of soluble factors. It is noteworthy that transport of these proteins is inhibited by the dominant-negative importin-β(45-462), as is the case for RCC1. It seems likely that the translocation of all these proteins occurs by a process of facilitated diffusion that requires specific, low-affinity interactions with nucleoporins that line the central channel of the pores. Facilitated diffusion cannot occur against a concentration gradient. Thus, nuclear pores must import and export RCC1 at the same rate and equilibrate the levels of free RCC1 in the cytoplasm to a concentration identical to that free in the nucleoplasm. The fact that RCC1 is >90% nuclear at steady state and does not shuttle detectably in vivo implies that the binding of RCC1 to chromatin is much more rapid than translocation across the nuclear envelope. In addition, it suggests that binding to chromatin can be sufficient to establish and maintain the nuclear localization of RCC1 and, hence, the Ran gradient. In effect, RCC1 acts as a chromatin marker, an activity key for the evolution of the nuclear compartment and the open mitotic cycle.

Key questions that remain to be explored include the identity of the RCC1-binding proteins on chromatin and at the nuclear pores. It will also be of interest to determine whether other import cargoes use the same pathway for transit across the nuclear envelope as RCC1.

We thank all members of the Macara lab for their continued advice throughout the experimental and preparatory phases of this manuscript. In particular, we thank Clark Peterson, Katie Welch, Amy Brownawell, Mark Lindsay, Alicia Smith, Kendra Pfleffer, and Scott Pfleffer for reagents and helpful suggestions. We also thank Takeharu Nishimoto, Dirk Gorlich, and Matthias Köhler for their generosity in providing some of the expression plasmids used in this study (RCC1, nucleoplasmin, nucleoplasmin core, the BIB zz domains, and importin-α3).

This work was supported by a grant from the National Institutes of Health, DHHS (GM-50526).

Submitted: 2 February 2000
R revised: 6 Apr 2000
A accepted: 13 Apr 2000

References

A dam, S.A., and L. Gerace. 1991. Cytosolic proteins that specifically bind nuclear localization signals are receptors for nuclear import. Cell. 66:837–847.
A dam, S.A., R.S. M arr, and L. Gerace. 1990. Nuclear protein import in permeabilized mammalian cells requires soluble cytoplasmic factors. J. Cell Biol. 111:807–816.
A zuma, Y., H. Seino, T. Seki, S. Uzawa, C. Klebe, T. Obba, A. Wittinghofer, N. Hayashi, and T. Nishimoto. 1996. Conserved histidine residues of RCC1 are essential for nucleotide exchange on Ran. J. Biochem. 120:82–91.
A zuma, Y., L. R euault, J.A. Garcia-Ranea, A. Valencia, T. Nishimoto, and A. Wittinghofer. 1999. Model of the Ran-RCC1 interaction using biochemical and docking experiments. J. Mol. Biol. 289:1119–1130.
B ischoff, F.R., and H. Ponstingl. 1991a. Catalysis of guanine nucleotide exchange on Ran by the mitotic regulator RCC1. Nature. 354:80–82.
B ischoff, F.R., B. Ono, and H. Ponstingl. 1991b. Mitotic regulator protein RCC1 is complexed with a nuclear ras-related polypeptide. Proc. Natl. Acad. Sci. USA. 88:10830–10834.
B ischoff, F.R., C. Klebe, J. Kretschmer, A. Wittinghofer, and H. Ponstingl. 1994. Ran GDP induces GTPase activity of nuclear Ras-related Ran. Proc. Natl. Acad. Sci. USA. 91:2587–2591.
B reeuwer, M., and D.S. Goldfarb. 1990. Facilitated nuclear transport of histone H1 and other small nucleophilic proteins. Cell. 60:999–1008.
C arras-Salas, R.E., G. Guarguaglini, O.J. Gruss, A. Segref, E. KarSENTI, and I.W. Matjaj. 1999. Generation of GTP-bound Ran by RCC1 is required for chromatin-induced mitotic spindle formation. Nature. 400:178–181.
C haiillan-Huntington, C.V., B. Braslavsky, J. K uhrmann, and M. Stewart. 2000. Dissecting the interactions between NTF2, Ran GTP, and the nucleoporins. XFXFG repeats. J. Biol. Chem. 275:5874–5879.
C hi, N.C., E.J. A dam, and S.A. A dam. 1995. Sequence and characterization of cytoplasmic nuclear protein import factor p97. J. Cell Biol. 130:265–274.
C hi, N.C., E.J. A dam, G.D. V isser, and S.A. A dam. 1996. RanGTP stimulates the interaction of ran with p97 in nuclear protein import. J. Cell Biol. 135:559–569.
C ortes, P., Z.S. Ye, and D. B altimore. 1994. RAG-1 interacts with the repeated amino acid motif of the human homologue of the yeast protein SRP1. Proc. Natl. Acad. Sci. USA. 91:7633–7637. function as well as sequence.
C uomo, C.A., S.A. Kirch, J. G yuris, R. Brent, and M.A. O ettinger. 1994. Rch1, a protein that specifically interacts with the RAG-1 recombination-activating protein. Proc. Natl. Acad. Sci. USA. 91:6156–6160.
D avis, L.I. 1995. The nuclear pore complex. A nnu. Rev. Biochem. 64:865–896.
D ingwall, C., and R.A. Laskey. 1991. Nuclear targeting sequences—a consensus? Trends Biochem. Sci. 16:478–481.
Matunis, M.J., E. Coutavas, and G. Blobel. 1996. A novel ubiquitin-like modification targets importin-alpha to mammalian nuclear pore complexes. J. Biol. Chem. 270:16499–16502.

Engelmeier, L., J.C. Olivo, and I.W. Maitaj. 1999. Receptor-mediated substrate translocation through the nuclear pore complex without nucleotide triphosphates. J. Biol. Chem. 274:930–931.

Featherstone, C., M.K. Darby, and L. Gerace. 1988. A monomodal antibody against the nuclear pore complex inhibits nucleocytoplasmic transport of protein and RNA in vivo. J. Cell Biol. 107:1289–1297.

Fleming, D.R., D. Mertz, T.M. Price, and J. Forbes. 1987. Inhibition of in vitro nuclear transport by a lectin that binds to nuclear pores. J. Cell Biol. 104:189–200.

Görlich, D. 1997. Nuclear protein import. Curr. Opin. Cell Biol. 9:412–419.

Görlich, D., and U. Kutay. 1999. Transport between the cell nucleus and the cytoplasm. Annu. Rev. Cell Dev. Biol. 15:675–690.

Görlich, D., S. Prehn, R.A. Laskey, and E. Hartmann. 1994. Isolation of a protein that is essential for the first step of nuclear protein import. J. Cell. Biol. 127:767–776.

Görlich, D., F. Vogel, A.D. Mills, E. Hartmann, and R.A. Laskey. 1995. Distinct functions for the two importin subunits in nuclear protein import. Nature 377:246–248.

Görlich, D., P. Henklein, R.A. Laskey, and E. Hartmann. 1996a. A 41 amino acid motif in importin-alpha confers binding to importin-beta and hence transit into the nucleus. EMBO (Eur. Mol. Biol. Organ.) J. 15:1810–1817.

Görlich, D., N. Pante, U. Kutay, A. Eber, and F.R. Bischoff. 1996b. Identification of different roles for RanGDP and RanGTP in nuclear protein import. EMBO (Eur. Mol. Biol. Organ.) J. 15:5584–5594.

Görlich, D., M. Dabrowski, F.R. Bischoff, U. Kutay, P. Bork, E. Hartmann, S. Prehn, and E. Izaurralde. 1997. A novel class of Ranbpg binding proteins. J. Cell Biol. 137:855–868.

Guan, T., S. Muller, G. Klier, N. Pante, J.M. Blevitt, M. Haner, B. Paschal, U. Aebi, and L. Gerace. 1996. Structural analysis of the Rpn2p complex, anassembly of O-linked glycoproteins that localizes near the central gated channel of the nuclear pore complexes. J. Mol. Biol. 261:949–963.

Herskowitz, A., R. Truant, H. Wiegand, and B.R. Cullen. 1998. Determination of the functional domain organization of the importin alpha nuclear import factor. J. Biol. Chem. 273:14209–14214.

Hoppe, L., K.H. Trajber, and R.G. Wurst. 1990. The yeast RNA1 gene product necessary for RNA processing is located in the cytosol and apparently excluded from the nucleus. J. Cell. Biol. 111:309–321.

Jakel, S., and D. Görlich. 1998. Importin β, transportin, RanBP5 and RanBP7 mediate nuclear import of ribosomal proteins in mammalian cells. EMBO (Eur. Mol. Biol. Organ.) J. 17:4491–4502.

Jakel, S., W. Aibig, U. Kutay, F.R. Bischoff, K. Schwamborn, D. Doenecke, and D. Görlich. 1999. The importin betaimportin 7 heterodimer is a functional nuclear import receptor for histone H1. EMBO (Eur. Mol. Biol. Organ.) J. 18:2411–2423.

Kaldner, D., B.L. Roberts, W.D. Richardson, and A.E. Smith. 1984. A short amino acid sequence to specify nuclear location. Cell 39:499–509.

Köhler, M., S. Ansieau, S. Prehn, A. Leutz, H. Haller, and E. Hartmann. 1997. Identification of a yeast karyopherin alpha family member in nuclear protein import. EMBO (Eur. Mol. Biol. Organ.) J. 16:1153–1163.

Lemonsbury, K.M., S.A. Richards, K.L. Carey, and I.G. M. acara. 1996. Mutations within the Ran/

Ribbeck, K., U. Kutay, E. Paraskova, and D. Görlich. 1999. The transcription of transportin-cargo complexes through nuclear pores is independent of energy. Eur. J. Cell Biol. 78:137–144.

Richards, S.A., K.L. Carey, and I.G. M. acara. 1997. Requirement of guanosine triphosphate-bound Ran for signal-mediated nuclear protein export. Science. 276:1842–1844.

Saha, S., D.B. Sparrow, T. Shiomi, R.T. Pu, T. Nishimoto, T.J. Mohun, and M. Dasso. 1997. Ubc5p and the conjugation of Sumo-1 to RanBP2. Curr. Biol. 7:1214–1218.

Schwoebel, E.D., B. Talcott, I. Cushman, and M.S. Moore. 1997. Identification of a protein complex that is required for nuclear protein import and mediates docking of import substrate to distinct nuclearpores. Proc. Natl. Acad. Sci. USA. 94:1769–1773.

Shikata, T., S. Tada, T. Katada, and T. Enomoto. 1997. Cloning of a cDNA encoding a novel receptor-mediated nuclear protein import pathway. J. Cell Biol. 139:657–666.
Biol. Chem. 269:24542–24545.

Takeda, S., T. Fujiwara, F. Shimizu, A. Kawai, K. Shinomiya, S. Okuno, K. Ozaki, T. Katagiri, Y. Shimada, M. Nagata, et al. 1997. Isolation and mapping of karyopherin alpha 3 (KPNA3), a human gene that is highly homologous to genes encoding Xenopus importin, yeast SRP1 and human RCH1. Cytogenet. Cell Genet. 76:87–93.

Uchida, S., T. Sekiguchi, H. Nishitani, K. Miyauchi, M. Ohtsubo, and T. Nishimoto. 1990. Premature chromosome condensation is induced by a point mutation in the hamster RCC1 gene. Mol. Cell. Biol. 10:577–584.

Wall, M.A., D.E. Coleman, E. Lee, J.A. Iniguez-Lluhi, B.A. Posner, A.G. Gilman, and S.R. Sprang. 1995. The structure of the G protein heterotrimer \(G_{i1}\)βγ. Cell. 83:1047–1058.

Weighardt, F., G. Biamonti, and S. Riva. 1995. Nucleocytoplasmic distribution of human hnRNP proteins: a search for the targeting domains in hnRNP A1. J. Cell Sci. 108:545–555.

Weis, K., I.W. Mattaj, and A.I. Lamond. 1995. Identification of hSRP1 alpha as a functional receptor for nuclear localization sequences. Science. 268:1049–1053.

Welch, K., J. Franke, M. Köhler, and I.G. Macara. 1999. RanBP3 contains an unusual nuclear localization signal that is imported preferentially by importin-alpha3. Mol. Cell. Biol. 19:8400–8411.

Welch, K., J. Franke, M. Köhler, and I.G. Macara. 1995. RanBP1 alpha as a functional receptor for nuclear localization sequences. Science. 268:1049–1053.

Weis, K., U. Ryder, and A.I. Lamond. 1996. The conserved amino-terminus domain of hSRP1 alpha is essential for nuclear protein import. EMBO (Eur. Mol. Biol. Organ.) J. 15:1818–1825.

The text continues...