Traffic to the inner membrane of the nuclear envelope
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Past research has yielded valuable insight into the mechanisms that regulate the nuclear transport of soluble molecules like transcription factors and mRNA. Much less is known about the mechanisms responsible for the transportation of membrane proteins to the inner membrane of the nuclear envelope. The key question is: does the facilitated transport of integral inner membrane proteins exist in the same way as it does for soluble proteins and, if so, what is it used for? Herein, we provide an overview of the current knowledge on traffic to the inner nuclear membrane, and make a case that: (a) known sorting signals and molecular mechanisms in membrane protein biogenesis, membrane protein traffic and nuclear transport are also relevant with respect to INM traffic; and (b) the interplay of the effects of these signals and molecular mechanisms is what determines the rates of traffic to the INM.

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Introduction
The nuclear envelope (NE) is a specialized area of the endoplasmic reticulum (ER). It is composed of two membranes, the inner and the outer nuclear membrane (INM and ONM), which come together in places where Nuclear Pore Complexes (NPCs) are embedded. In many eukaryotes, a proteinaceous surface, namely the nuclear lamina, underlies the INM. The perinuclear space in between the two membranes is continuous with the ER lumen. The ER, ONM and INM are also continuous, but have distinct functions and sets of transmembrane proteins. Assigning proteins as true INM residents is problematical for multiple reasons, ranging from technical difficulties in microscopically resolving their localization in the INM or ONM, to biological reasons such as their cell type specificity [1]. Bioinformatic predictions are difficult to make, as only a few domains specific to INM proteins have been identified, such as the LEM (for Lap1-emerin-MAN1) and SUN (for Sad-Unc-84 homology) domains for which structures are available [2,3]. A decade ago, the first proteomic studies aimed at identifying putative INM proteins were performed [4,5] but to date, only a relatively small number of these proteins have been both well-characterized and proven to be enriched in the inner membrane compared to the outer membrane and ER.

The importance of the correct trafficking and function of INM proteins is clear from numerous examples of the roles played in the development of nuclear envelopopathies and cancer. Accordingly, the lamina-associated polypeptide 2, Lap2B, is over-expressed in digestive tract cancers [6]. Mutations in the lamin B receptor, LBR, cause both Greenberg dysplasia, a major disease leading to aberrant embryonic development [7], or Pelger-Huet anomaly [8]. Laminopathies are often linked to mutations in lamin A, but recent studies show that the mistargeting of INM proteins could be causative of the disease phenotypes [9,10]. For example, Hutchinson–Gilford Progeria Syndrome (HGPS), a serious accelerated ageing disease, is caused by a dominant \textit{de novo} mutation in lamin A that results in the accumulation of progerin, which is a farnesylated lamin A variant. In HGPS cells, the levels of SUN1 in the INM are increased [9,11], and knocking-down SUN1 alleviates cellular senescence [9]. Similarly, nuclear deformation and cell survival are rescued by SUN1 knock-down in mice cells lacking lamin A or carrying progerin-like mutations [9].

Over the years, multiple mechanisms of INM protein targeting have been proposed, involving a variety of potential sorting signals. Earlier work suggested that the interplay between multiple signals is required for the efficient targeting of INM proteins [12,13]. Clearly, there will be multiple signals encoded on a specific membrane protein to guide its biogenesis and targeting. These signals may encode information for: insertion into the lipid bilayer, cytosolic subcellular sorting to the different membrane compartments, and nuclear transport. For each of these categories short descriptions of the molecular signals and mechanisms (‘molecular toolboxes’) are given (Fig. 1 and Toolbox I, II and III). Table 1 contains an overview of some of the better studied integral membrane proteins that are enriched in the INM in \textit{Saccharomyces cerevisiae} and
Humans. We have also sorted into three molecular toolboxes the plethora of targeting information that has been experimentally validated.

En route to the INM

**En route to the INM: membrane insertion**

Essential steps of the targeting process is the synthesis and insertion of the nascent polypeptide to the membrane environment (Fig. 1, D) [14]. The two conserved insertion machineries, the Sec61 and GET (Guided-Entry of TA proteins) systems, are situated in the ER, including the ONM. An INM localized pool of Sec61 might exist [15] and the GET transmembrane components are small and may also passively reach the INM through the lateral channels of the NPC. Thus, in principle a post-translational mechanism where a chaperoned INM protein is first trafficked to the nucleus, after which it is membrane inserted, could be possible for membrane proteins that are posttranslationally inserted such as very small monotopic membrane proteins and tail anchored proteins. However, this has not been tested directly.

Monotopic membrane proteins could be targeted at either of the two insertion machineries. From the proteins listed in Table 1, only emerin and LAP2B are potentially inserted via the GET pathway. The small splice variant of Heh1(-helix-extension-helix-1)/Src1 known as Src1-small, and Mps3 (monopolar spindle), SUN1 and SUN2 have larger luminal domains and their insertion is probably facilitated by the Sec61 system. The polytopic membrane proteins, Heh1/Src1, Heh2 (helix-extension-helix-1), LEM2 (Lap1-emerin-MAN1-2), MAN1, LBR and numir, are probably inserted by Sec61 co-translationally.

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**Toolbox I  Protein insertion**

Membrane protein integration into the lipid bilayer is a facilitated process. There are two well-characterized insertion systems that are conserved from yeast to man: the Sec61 system and the GET pathway (reviewed in [14,55,56]). The Sec61 system translocates soluble proteins and membrane proteins with single (monotopic) and multiple transmembrane spanning segments (polytopic). The current data supports that polytopic membrane proteins are inserted cotranslationally by the Sec61 system. The GET system evolved for the specialized post-translational insertion of tail anchored proteins, which are proteins with a single transmembrane spanning segment at their C-terminus and a short luminal tail. Small monotopic membrane proteins may also be inserted posttranslationally. Prediction of the topology of membrane proteins is based on hydrophobicity profiles and the characteristic features of the regions flanking the transmembrane segments. For example, if the region N-terminal of the transmembrane domain is long or positively charged (positive-inside rule), then it is likely extraluminal. The translocation of the N-terminus to the lumen is likely when positive charges are lacking, when the preceding region is not well folded and the hydrophobic sequence is long. Besides the Sec and GET pathways, other insertion machineries also exist, for example, in yeast, the Sec61 homolog, Ssh1, and the Sec63 complex.

The question of post-translational or co-translational insertion could be relevant to traffic of INM proteins, but has been little studied. In particular post-translational insertion via the GET pathway could occur at the INM post nuclear import, in which case the transport occurs as a soluble chaperoned protein.

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**Toolbox II  Cytosolic subcellular traffic**

In general, membrane proteins may traffic through different subcellular compartments before they reach their destination, for example to be modified post-translationally. In addition, membrane proteomes are generally dynamic, for example there is a rapid exchange between the plasma membrane, pools of vesicles and the ER network. For the trafficking of membrane proteins to the different cellular membranes, signal sequences exist such as those for Golgi retrieval, ER retention, and peroxisome and mitochondrial targeting. The localization of a protein encoding multiple signals depends on the kinetics of the different trafficking routes.

It may be a mistake to think of INM proteins as stable components of the INM; they may well also have a dynamic localization within the cell that is regulated by the interplay of the above sorting signals. Cases where cytosolic subcellular traffic is relevant for traffic of INM proteins are given in Table 1.

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**Toolbox III  Nuclear import**

The NPCs are anchored where both the INM and ONM come together to form the highly curved pore membrane. Their overall architecture and function is broadly conserved from yeast to humans. There are also distinct differences between yeast and metazoan NPCs: the yeast NPCs are smaller in size and molecular weight and each of them has several unique components [57,58]. NPCs are composed of a scaffold of folded proteins that anchor the 8-fold rotational symmetric structure to the nuclear envelope membrane. A set of intrinsically disordered proteins, the FG-Nups, are anchored to the scaffold of the NPC, and are critical for the selectivity of the pore. For soluble proteins, the mechanisms of nuclear transport are well described [59]. Molecules may diffuse through the NPC passively (efflux/ influx) and equilibrate between the cytosol and nucleus. Transport factors and the gradient of RanGTP–RanGDP across the nuclear envelope are required to specifically ‘pump’ proteins against a concentration gradient and to transport very large macromolecular complexes across the NPC. In these facilitated import and export reactions, soluble transport factors shuttle Nuclear Localization Signal (NLS)-containing proteins or Nuclear Export Signal (NES)-containing proteins across the NPCs. The FG-Nups encode multiple phenylalanine and glycine (FG)- repeats that act as binding sites for the soluble transport factors. Direction to the transport reaction is given by the gradient of RanGTP–RanGDP across the nuclear envelope: in an import reaction, the transport factor dissociates from cargo in the presence of RanGTP, thereby releasing the cargo in the nucleus. In addition, retention mechanisms usually play a role in defining nuclear and cytosolic concentrations of soluble proteins.

Retention mechanisms also play a major role in defining INM localization of membrane proteins. In addition, specifically in yeast, there is good evidence for the facilitated import of Heh1 and Heh2 resulting in accumulation in the INM. Alike for soluble proteins, traffic of these INM proteins depends on FG-Nups, Kaps and the gradient of RanGTP–RanGDP. The sorting signal is composed of an NLS and a long intrinsically disordered linker.
Many INM proteins have relatively large N-terminal extralumenal domains, which often contain regions that have been proved to be relevant for trafficking (Fig. 2). The early recognition of INM proteins, including as early as during translation, was first proposed for viral peptides and later for native INM proteins [16–18]. Here, a shorter isoform of importin-α was shown to both bind a nascent polypeptide chain predisposed for the INM at a stretch of positive charges located 5–8 residues from the transmembrane segment, and direct it to the translocon [16–18]. Whether this is a significant sorting event specifically in Heh2 is unclear, as the absence of this sequence does not affect localization [19]. Accordingly, instead of a sorting sequence, this could be regarded as a manifestation of the positive inside rule guiding membrane insertion. We take that the early recognition of Heh2, being destined for the INM, is more likely to occur through the early binding of the yeast homolog of importin-α, Kap60, to the exceptionally strong NLS of Heh2 [20*].

**En route to the INM: cytosolic subcellular sorting**

The localization of a protein encoding multiple signals depends on the kinetics of the different trafficking routes. Anything that disrupts this balance can cause a change in localization. For example, mitochondria have a separate system for tail anchored protein insertion that could potentially compete with the GET system inserting them into the ER [21]. Knowledge of the cytosolic subcellular sorting (Fig. 1, II) of integral INM proteins is thus far limited, but SUN2 is a clear example of how elements of subcellular sorting between the ER and Golgi are important. This monotopic INM protein possesses an Arg-rich Golgi retrieval signal that is necessary for its INM localization [12]. Similar Arg-rich sequences are found in LBR, Lap2β, emerin and LEM2, but their involvement in targeting has not yet been characterized.

Changes in the concentrations of interaction partners can also disturb proper sorting. For instance, SUN2 was found...
in endosomes when Rab5, a small GTPase responsible for endosomal membrane fusion and complexing SUN2, was over-expressed [22]. Another example is the mislocalization of SUN1 to the Golgi that has been observed in mice lacking functional wild-type lamin A [9*]. Lamin A is probably needed for the retention of SUN1, preventing it from travelling to the Golgi. Indeed, when the N-terminal lamin A-interacting domain of SUN1 is deleted, the SUN1 relocates from the NE to the Golgi [9*]. A balance between subcellular localizations is also required for LBR, which has two separate functions: inside the nucleus it is responsible for regulating the structure of the NE, as illustrated by its role in maintaining the lobulated structure of granulocyte nuclei [8], but it also acts as a sterol reductase for which it has to be ER localized [7]. How the dual localization is controlled is presently unclear.

Intriguing connections with plasma membrane localization also exist. Emerin, for instance, targets to the plasma membrane in the heart tissue of some animals [23]. Also interesting is how the epidermal growth factor receptor (EGFR) travels from the plasma membrane to the nucleus upon EGF binding. The receptor is endocytosed and travels through the Golgi and to the ER via COPII regulated retrograde vesicle trafficking [24,25]. The next steps include translocation to the nucleus and extraction from the membrane, although the order in which this happens is unclear, nor is what triggers membrane extraction.

**En route to the INM: nuclear import**

Current models of the transport of INM proteins disagree significantly on the nature of energy dependence: is there or is there not an active energy dependent import that drives the accumulation of membrane proteins in the INM? When looking at soluble proteins like transcription factors, we mostly see that retention mechanisms, as well as the kinetics of import, export, influx and efflux, define their localization (Fig. 1, III). These kinetics can be adapted by modification or the shielding of import and export signals.

Many membrane proteins in the INM are retained due to interactions with nuclear components, most notably lamins and chromatin and SUN-KASH interactions in the lumen, but there is now good evidence that this ‘selective retention’ is not the sole basis for their nuclear presence. An initial report on the energy (and temperature) dependence of INM protein import [26] suggested that ATP is used for NPC restructuring which creates transient channels through which the proteins could travel. Later reports show that several INM proteins make direct or indirect use of the classical nuclear transport elements, including NLSs, Kaps and FG-Nups. *S. cerevisiae* Heh1 and Heh2 and human SUN2 have confirmed NLS sequences [12,27], while others have predicted sequences [28]. Moreover, Heh1 and Heh2 localization is dependent on the transport factors Kap60 and Kap95 (yeast importin-β), the RanGTP/RanGDP gradient, and a subset of FG-Nups [19,27]. In *S. cerevisiae*, a combination of an NLS and an intrinsically disordered (ID) linker (L) is required and is sufficient for INM targeting. This ‘NLS-L’ motif targets a Heh2 transmembrane domain, a polytopic Sec61 transmembrane domain and a synthetic transmembrane domain composed of leucine alanine repeats to the INM. We propose that the ID linker facilitates binding to the transport factors and interactions with the FG-Nups [19,29]. Alternatively, or additionally, the combination of the strong NLS and the ID linker acts earlier in the membrane protein biogenesis or traffic. Consistent with a facilitated transport mechanism, large extralumenal domains are tolerated [20*]. However, more importantly, using these mobile proteins it was shown that, upon blocking import, the protein leaks out from the INM to the ER. This demonstrates that INM accumulation is the result of fast import and slower efflux, and reflects energy driven accumulation. Facilitated NLS mediated import of proteins with large extralumenal domains has been reproduced with polytopic transmembrane proteins, which should resolve the discussion of whether the transmembrane segments are embedded in the membrane during transport (unpublished). Having reinforced the aspect of the facilitated transport of these yeast INM proteins, we emphasize that retention also plays a role. Full length Heh1 and Heh2 have LEM domains, and their diffusion in the membrane is much slower than that of truncated versions without the LEM domain. This is consistent with them binding to nuclear structures [19]. Overall, as for soluble proteins, the localization of these INM proteins is defined by the kinetics of import, leakage and nuclear retention.

For INM proteins without predicted NLS sequences, other mechanisms for facilitated transport have been proposed, for example via FG repeats encoded on the INM proteins [30], or via a piggyback mechanism in which membrane proteins bind to a soluble NLS-containing protein. The latter mode of transport was proposed for Mps3, which binds histone H2Z.A [31]. Some of the INM proteins that are thought to localize due to retention may in fact make use of the piggyback import. Lamins come to mind as potential piggyback candidates. The current thinking is that lamins contribute to sorting by retaining INM proteins upon arrival at the INM, but a role in piggyback transport of INM proteins cannot be excluded until we measure where they first associate. For example, prelamin A may have such a role in targeting of SUN1 to the INM. In differentiating human myoblasts, farnesylated prelamin A accumulates in and recruits SUN1 to the NE [32]. Additionally, a type of lamin A, possibly the unprocessed or mature forms, prevents SUN1 from travelling to the Golgi [9*]. Farnesylated prelamin A also interplays with SUN2 targeting in differentiating
Here, the enrichment of SUN2 at the nuclear poles depends on farnesylated prelamin A [32]. Moreover, in patients with Mandibuloacral dysplasia with type A lipodystrophy (MADA), which is a rare disease caused by the accumulation of unprocessed prelamin A, SUN2 distribution in the NE is disorganized. This is rescued by drugs that reduce prelamin A farnesylation [10].

In conclusion, particularly in yeast, there is good evidence for the facilitated import of membrane proteins that results in accumulation in the INM. In human cells, there is no definitive answer as to whether facilitated transport alone can result in accumulation in the INM. Nevertheless, it is clear that retention mechanisms play a major role in both yeast and mammalian systems.

**Putative NPC independent traffic**

As discussed [33], NPC independent routes across the NE, such as those used by viruses [34], may also be available to traffic native membrane proteins. For the replication of the Herpes Simplex virus, large nucleocapsids are formed in the nucleus, which have to pass the NE before their...
maturation in the cytosol. An NPC independent export model, namely nuclear egress, is currently accepted as an explanation for this phenomenon (reviewed in [35]). The nuclear localized capsids are enveloped by the INM and cross the perinuclear space as vesicles, which fuse with the ONM and release the capsid to the cytoplasm. The same mechanism is reported in *Drosophila melanogaster* for the export of ribonucleoprotein particles (RNP)s, which are too large to pass the NPC [36]. Perinuclear granules have been observed in other cell types and species, so the nuclear egress might in fact be a conserved export mechanism. Nuclear egress has been hypothesized to be involved in the removal of nuclear protein aggregates [37]. Future studies will have to demonstrate if membrane proteins could exit the nucleus via any such egress pathway.

**Challenges when studying INM import**

**Kinetics matter**

Definitive proof of the existence of the facilitated transport of membrane proteins requires verification that import across the NPC is faster than efflux, as well as a demonstration that import is transport factor and Ran

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**Table 1**

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<td>Yeast Mps3</td>
<td>Cotranslational Sec1 system&lt;sup&gt;a&lt;/sup&gt;</td>
<td></td>
<td>- Indirect dependence on Kap123, Kap95 and RanGTP–RanGDP gradient; piggyback mechanism via binding to histone H2Z.A. - Nuclear retention</td>
<td></td>
<td>[31,60]</td>
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<tr>
<td>Heh1/Src1</td>
<td>- Src1 small: cotranslational Sec61&lt;sup&gt;a&lt;/sup&gt; - Full length Heh1: cotranslational Sec61&lt;sup&gt;a&lt;/sup&gt;</td>
<td></td>
<td>- NLS, RanGTP–RanGDP gradient, Kap60, Kap95, Nup170, Nup2 - Nuclear retention</td>
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<td>[19,27]</td>
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<td>Heh2</td>
<td>- Cotranslational Sec61&lt;sup&gt;a&lt;/sup&gt;; - We interpret ‘INM sorting motif’ is topology indicator</td>
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<td>[17,19,27]</td>
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<td>Human SUN1</td>
<td>Cotranslational Sec61&lt;sup&gt;a&lt;/sup&gt;</td>
<td></td>
<td>- Nuclear retention</td>
<td>Localization depends on farnesylated prelamin A SUN2 mobility requires ATP</td>
<td>[11*,32,61]</td>
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<tr>
<td>SUN2</td>
<td>Cotranslational Sec61&lt;sup&gt;a&lt;/sup&gt;</td>
<td>Golgi retrieval signal</td>
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<td>[12,22,62,63]</td>
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<td>Emerin</td>
<td>Tail anchored protein, possibly posttranslational insertion by GET pathway&lt;sup&gt;a&lt;/sup&gt;</td>
<td>Subpopulation in plasma membrane in heart tissue from human, rat and mouse (sorting signals unknown)</td>
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<td>Emerin mobility requires ATP</td>
<td>[23,63]</td>
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<td>LAP2β</td>
<td>Tail anchored protein, possibly posttranslational insertion by GET pathway&lt;sup&gt;a&lt;/sup&gt;</td>
<td></td>
<td>- Nuclear retention</td>
<td></td>
<td>[26,64–66]</td>
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<tr>
<td>LEM2 MAN1 LBR</td>
<td>Cotranslational Sec61&lt;sup&gt;a&lt;/sup&gt; - N terminal domain probably co-defines topology; - ‘INM sorting motif’</td>
<td>Distinct functions at ER and NE (sorting signals unknown)</td>
<td>- RanGTP dependent interaction with Importinβ (not importin-α dependent)</td>
<td>Mobility of LBR is dependent on RanGTP and Nup35</td>
<td>[7,16,63, 69–72]</td>
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<tr>
<td>Nurim</td>
<td>Cotranslational Sec61&lt;sup&gt;a&lt;/sup&gt; - ‘INM sorting motif’</td>
<td></td>
<td>- Nuclear retention (but not to DNA and not to lamins)</td>
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<td>[73,74]</td>
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<sup>a</sup> Prediction.
dependent. This requires methods that allow the direct assessment of transport kinetics through the NPC, distinct from the kinetics of diffusion in the INM and ONM. Single molecule tracking experiments would be uniquely suitable, but are thus far unexploited.

Alternatively, it is possible to measure rates of bulk efflux or bulk import. Bulk efflux is measured in experiments that start with an accumulation in the INM and then follow the kinetics of equilibration after blocking facilitated import. The steady state accumulation levels together with the efflux kinetics reveal the kinetics of import. These measurements can only be obtained when the proteins of interest are freely diffusing and are not retained in either compartment. The absence of protein turnover over the measured time period is also critical. However, for all known INM proteins, the binding of nuclear localized proteins is an important retention mechanism which makes them unsuitable for bulk efflux measurements. As a consequence, truncated versions that lack retention signals, or even synthetic constructs encoding only the minimally required sorting signals [19,20*], must be used for these studies.

Where do membrane proteins travel through the NPC?

Based on electron tomosgraphs of metazoan NPCs [38-41], the most logical pathway of the extraluminal domains of INM proteins is along the pore membrane through the lateral channels (Fig. 1). These channels are flanked on the cytoplasmic and nuclear sides by the proteins from the outer ring Y-shaped Nup84 or Nup107 subcomplexes in yeast and humans, respectively [42]. An approximately 10 nm space is available between the membrane and this part of the NPC scaffold. More centrally in the NPC, the lateral channels are flanked by its integral membrane proteins, and here the passage seems to be more restricted. Unfortunately, high resolution tomosgraphs are not available of S. cerevisiae.

Membrane proteins have been proposed as passing through (a) the lateral channels, which seems likely judging from the tomosgraphic images of NPCs, or (b) the central channel, which is likely considering the involvement of FG-Nups and Kaps. Three main uncertainties about the structure of the NPC are relevant here. Firstly, yeast and human pores may differ critically, and whereas (a) is largely supported by work regarding metazoans, (b) is mostly from work with respect to baker’s yeast. Secondly, whether the disordered FG-Nups occupy the space in the lateral channels, and whether FG-Nups facilitate karyopherin mediated traffic through the lateral channels, is unknown. Thirdly, NPCs are flexible structures in which the position of the 8-fold rotational symmetric units is variable [39]. At the impressive but still limited resolution available, it is uncertain whether small or temporary openings exist between the centre of the NPC and the lateral channels. Accordingly, to resolve the route(s) through the NPC (even) better knowledge of its dynamic structure is required.

Why INM targeting would be needed

Recent studies have uncovered new exciting functions of integral membrane proteins residing in the INM, and while for some of these activities passive diffusion and selective retention is sufficient, for others a tighter control of protein localization could be expected. Passive diffusion may be enough for LAP2β and MAN1, which have been shown to (redundantly) mediate the assembly of the NE [43]. High enrichment in the INM may possibly be required for proteins that play a role in NPC assembly into an intact NE. NPC assembly in the intact NE in yeast depends on Hef1 and Hef2 [44], while Sun1 and an INM-localized pool of Pom121 play a role in NPC assembly in humans [45-49]. INM proteins can also directly contribute to the INM acting as a ‘transcription factor resting place’ by sequestering transcription factors that illegitimately entered the nucleus and as such prevent transcription of target genes [50].

Functions related to chromatin anchoring to the nuclear periphery might depend on facilitated transport, as they require a higher level of regulation (recent reviews [51,52]). An analysis of cells with inverted chromatin architecture has provided interesting insights into this topic [53**]. The heterochromatin of rod photoreceptor cells of nocturnal mammals is not located on the nuclear periphery, but is shifted to the nuclear interior. This phenotype occurs gradually during differentiation and is caused by the lack of the anchoring proteins LBR and lamin A/C in mature cells, whereas LBR is still present in the cells at early stages of differentiation. This sequential expression of the above-mentioned proteins during differentiation has also been observed in different mouse tissues, and has a potential effect on the expression of cell type specific genes. The deletion of LBR or lamin A in differentiating myotubes have the opposite effects: a lack of LBR increases the expression of muscle specific genes, while the loss of lamin A reduces it. There is no LBR or lamin A regulated effect on the expression level of the same genes in mature muscle cells. These results suggest that INM proteins act as heterochromatin tethers to regulate differentiation. Indeed, several INM proteins are able to reposition specific chromosomes and are restricted to certain tissues [54*].

These observations strongly support the idea that INM proteins localize in the nucleus specifically to shape chromatin and regulate transcription, and do not enter the nucleus by chance and stay there due to an interaction with DNA. So, in addition to the regulation of the expression or turnover of INM proteins, facilitated import may also play an important role in tuning the
proteome of the inner membrane, and with that the proteome of the cell.

Concluding remarks
We asked the question as to whether the facilitated transport of integral inner membrane proteins exists in the same way as it does for soluble proteins, and, if so, what is it used for. We conclude that there is ample evidence that the facilitated import of integral membrane proteins exists in *S. cerevisiae*. Some may argue that facilitated import in yeast is a consequence of its closed mitosis and lack of lamins. However, the biological evidence of INM proteins directing chromosome localization and transcriptional regulation, as well as the presence of NLS sequences, suggests that facilitated transport is also present in humans. A better understanding of the transport of integral membrane proteins to the INM should go hand in hand with research aimed at uncovering new roles of INM proteins in chromatin organization and signal transduction in development, ageing and differentiation.

Acknowledgements
We are grateful to Patrick Lusk for his valuable discussions. This work is financed by the research programmes NWO-vidi and ECHO from the Netherlands Organization for Scientific Research (NWO) and by the Netherlands Proteomics Centre.

References and recommended reading
Papers of particular interest, published within the period of review, have been highlighted as:

- of special interest
- of outstanding interest


The SUN domain is one of the few INM protein specific domains identified, and this paper presents its crystal structure, in complex with KASH. The structure shows a hexameric arrangement of three SUN and three KASH domains.


9. Chen C-Y, Chi Y-H, Mutalif RA, Starost MF, Myers TG.

This study shows the pathology of upregulation and Golgi targeting of SUN1, when Lamin A is mutated or lacking. Knocking down SUN1 in HGPS cells and mutant Lamin A mouse models, rescues the cells from nuclear deformations and senescence.


11. Haque F, Mazzeo D, Patel JT, Smallwood DT, Ellis JA.

Some lamin A mutations, associated with laminopathies as Emery–Dreifuss muscular dystrophy and Hutchinson–Gilford Progeria Syndrome (HGPS), disrupt interactions with SUN1 and SUN2. SUN1 is accumulated at the NE in HGPS cells, which correlates with Prelamin A accumulation.


A quantitative analysis of karyopherin-mediated import and passive efflux of Heh2 derived reporters and their mobility in different membrane compartments is presented. Membrane proteins with large extraluminal domains terminal to the intrinsically disordered linker passively leak out from the INM to the ONM and ER.


remodeling

Current

44

Cell nucleus


This paper provides the first experimental proof for the egress mechanism being an endogenous export pathway. Authors investigate ribonucleoprotein particles (RNPs) formation and transport from the nucleus in Drosophila melanogaster. Using microscopy they show RNPs localize to the perinuclear space and can leave the nucleus. The same molecular remodeling machinery is used as during viral egress.


The authors study the rod photoreceptor cells from nocturnal mammals, which have inverted chromatin architecture (with heterochromatin in the center of the nucleus), and discover the sequential expression of LBR and lamin A during development of many cell types and its effect on chromatin positioning and transcription.


Certain chromosomes are at the nuclear periphery in some tissues, and in the nuclear interior in others, which has obvious effects on expression of encoded proteins. Authors postulate that this observation might be caused by tissue specific expression of novel nuclear envelope proteins that they identified before.


While the cotranslational pathway of membrane proteins insertion into the ER was discovered more than 30 years ago, the first molecular players in the postranslational insertion pathway for tail-anchored proteins were only discovered in Since then great progress in understanding the GET (Guided Entry of Tail Anchored proteins) was made, reviewed here.


