Review

Uptake and Fate of Extracellular Membrane Vesicles: Nucleoplasmic Reticulum-Associated Late Endosomes as a New Gate to Intercellular Communication

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Abstract: Extracellular membrane vesicles (EVs) are emerging as new vehicles in intercellular communication, but how the biological information contained in EVs is shared between cells remains elusive. Several mechanisms have been described to explain their release from donor cells and the initial step of their uptake by recipient cells, which triggers a cellular response. Yet, the intracellular routes and subcellular fate of EV content upon internalization remain poorly characterized. This is particularly true for EV-associated proteins and nucleic acids that shuttle to the nucleus of host cells. In this review, we will describe and discuss the release of EVs from donor cells, their uptake by recipient cells, and the fate of their cargoes, focusing on a novel intracellular route wherein small GTPase Rab7 late endosomes containing endocytosed EVs enter into nuclear envelope invaginations and deliver their cargo components to the nucleoplasm of recipient cells. A tripartite protein complex composed of (VAMP)-associated protein A (VAP-A), oxysterol-binding protein (OSBP)-related protein-3 (ORP3), and Rab7 is essential for the transfer of EV-derived components to the nuclear compartment by orchestrating the particular localization of late endosomes in the nucleoplasmic reticulum.

Keywords: exosome; extracellular vesicle; intercellular communication; late endosome; oxysterol-binding-related protein; nucleoplasmic reticulum; Rab7; VAMP-associated protein A

1. Introduction

Intercellular communication is a fundamental feature for the development and maintenance of multicellular organisms. Diverse molecular mechanisms for the exchange of biological information between cells have been documented. The secretion of soluble proteins and their interaction with membrane receptors located on the target cells or contact-dependent signaling are good examples. To better understand the mechanism that triggers molecular crosstalk between cells, we came to the fascinating and poorly explored world of extracellular membrane vesicles (EVs) [1–4]. We will review here the mechanisms underlying the release of different types of EVs from donor cells into the...
extracellular medium, their uptake by recipient cells, and the fate of their cargoes with a focus on a new intracellular pathway that led to their transfer into the nuclear compartment [5,6].

2. Extracellular Membrane Vesicles and Intercellular Communication at a Glance

EVs are nanobiological membrane structures usually referred to as exosomes or microvesicles depending on their biogenesis and size [7,8]. Exosomes (~40–100 nm in diameter) are produced by inward budding inside an endosome, leading to the formation of a late endosomal multivesicular body (MVB) that could fuse afterward with the plasma membrane and discharge its internal small vesicles in the extracellular milieu, whereas microvesicles, also named ectosomes or shed vesicles (~50–1000 nm), bud directly from the plasma membrane. To complete our non-exhaustive view of the EV catalog, in addition to apoptotic microvesicles (<1000 nm) and apoptotic bodies (1–5 µm) derived from dying cells [9], bulky EVs (1–10 µm), often termed large oncosomes [10], are formed upon cleavage of plasma membrane extensions of tumor cells harboring an amoeboid-like phenotype [11,12]. Nodal vesicular parcels involved in left–right asymmetry during the organism development can be viewed as atypical particles or EVs (~300–500 nm) containing lipophilic granules [13]. Lipoprotein particles can also contribute to exchange materials and promote signaling between cells [14]. Hereafter, we will focus on exosomes and microvesicles.

EVs carry a restricted set of membrane-protected proteins, lipids, and nucleic acids (e.g., messenger (m) RNA, micro (mi) RNA, and long non-coding (nc) RNA) that could act as pivotal mediators in the regulation of neighboring and distant recipient cells [15–20]. The number of bioactive molecules carried by a given type of EVs, especially small-sized ones such as exosomes, can be extremely little; therefore, an efficient mechanism must operate to trigger a cellular response. This is particularly true when the target cells are distant from the donor cells and the amount of EVs is limited [21,22]. The content of EVs depends on the cell type of origin and their physiological conditions. Thus, EVs represent a heterologous population in a given biofluid. Specific isolation of their subpopulations is now an emerging challenge in the field, particularly when the purpose is to use EVs (or their cargo) as potential clinical biofluid markers [23–29]. We could not exclude that distinct EVs interact with each other, resulting in their co-purification [30]. All these pitfalls should be considered. Diverse biological functions are ascribed to EVs in cell-to-cell communication, such as favoring proliferation versus differentiation of stem cells, inducing epithelial-mesenchymal transition, and modulating immune responses among others [31–33]. EV-mediated intercellular communication within an organ is also beginning to be recognized. For instance, neural cells exchange biochemical information upon the release of exosomes or microvesicles [34]. In addition to inter-neuronal communication, EVs have been suggested to play a role in crosstalk between neurons and glial cells, notably oligodendrocytes, and hence support the neuronal physiology [35,36]. In pathological conditions, EVs might participate in intercellular transfer of prions or the progression of various diseases including Alzheimer’s disease [37–39]. In cancer, the components carried by transformed cell-derived EVs play a role in the establishment of a pre-metastatic niche [40–43]. They can also contribute to horizontal propagation of oncogenes among subsets of cancer cells or surrounding healthy cells [44,45]. Lastly, the EV-mediated crosstalk between cancer and non-cancerous cells in the bone marrow microenvironment can modulate the biochemistry and function of stromal cells to stimulate the growth and spreading of cancer cells [46–48].

2.1. Release of Extracellular Membrane Vesicles

In line with a variety of EV subtypes, numerous modes of EV release were described [49]. Our labs gained insight into mechanisms underlying EV secretion by following the cellular trafficking of the stem cell marker CD133 (prominin-1), which is associated with plasma membrane protrusions (e.g., microvilli and cilia) and regulates their organization and function [50–52]. Although the observations with CD133 are not exclusive to this glycoprotein, they summarized the knowledge about the EV biogenesis. CD133+ EVs are released into various internal and external body fluids using different mechanisms that are dependent in part on the cellular type [53]. In non-epithelial cells, CD133+ EVs are discharged
as exosomes via exocytosis of MVB (Figure 1A, point 1) [54]. The interaction of CD133 with syntenin-1 and its ubiquitination might be two essential steps involved in the internalization and sorting to MVB en route to exosomes [55,56]. Together with ALIX and syndecan, syntenin-1 regulates the biogenesis of exosomes [57]. The ubiquitinated CD133 interacts with tumor susceptibility gene 101 protein (TSG 101), a component of endosomal sorting complex required for transport (ESCRT) machinery involved in MVB formation [56]. We invite the readers to consult the following excellent reviews [3,58–60] for mechanistic details related to (i) the sorting of proteins to exosomes; (ii) the MVB biogenesis including ESCRT machinery; and (iii) their fusion with plasma membrane (Figure 1). In epithelial cells, CD133+ EVs bud as microvesicles/ectosomes from primary cilium and/or microvilli present at the apical plasma membrane (Figure 1A, points 2 and 3) [61,62]. Nader and colleagues described that actin polymerization is required for the release of microvesicles from ciliary tips by promoting membrane scission [63]. CD133 could itself play a role in this process [51,64]. ESCRT complex proteins might also participate in the release of microvesicles from plasma membrane [65]. The midbody that bridges nascent daughter cells at the end of cell division is also a source of microvesicles, and the midbody itself can be released depending on cellular status [61,66].

Lipid-based membrane microdomains (known as lipid rafts) and/or certain lipids (e.g., cholesterol, and ceramides), lipid-metabolizing enzymes, and lipid-interacting proteins may play a role in initiating budding processes either at the level of endosomes or plasma membrane (Figure 1A, red segment) [51,54,67,68]. The enrichment of lipid raft-associated cholesterol and sphingomyelin in EVs is consistent with such hypothesis [24,69]. Tetraspanin proteins such as CD9, CD81, and CD82 could be implicated in the sorting of various cargoes to EVs by forming a dynamic platform within the lipid bilayer membrane with other membrane and cytosolic proteins that would favor the formation of a specific dynamic network (also known as the tetraspanin web), and hence stimulate the budding process [70–72]. To conclude, it is important to note that the release of EVs by a given cell type is not restricted to a particular pathway; however, one mechanism can predominate under specific physiological or pathological conditions.

2.2. Uptake of Extracellular Membrane Vesicles

Once released in biofluids or the extracellular milieu between cells embedded in a tissue, circulating EVs can spread the selective information that they carry. All cells that are in direct contact with a given biofluid containing EVs are potential targets. Although poorly characterized, the diffusion of EVs within a tissue might be regulated by a source-sink mechanism, as suggested for morphogens [73,74]. Several barriers can impede the proper delivery of EVs such as their sequestration in the extracellular matrix, their degradation by proteolytic enzymes, and/or unspecific binding to non-targeted cells. The interaction of EVs with cell surface-associated extracellular matrix in a given organ/tissue may allow the specific targeting of EVs, a process mediated by EV-associated specific integrins, and their uptake by resident cells at the predicted metastatic destination [75,76]. The integrins (e.g., β1) carried by EVs can also promote anchorage-independent growth of tumor cells [77,78].

To achieve their vehicle-like function, EVs need to interact with recipient cells, which can be done in certain cases in a specific manner [79]. EVs could bind directly to the cell surface of recipient cells. In kidney, cellular protrusions such as cilia are the preferential sites for EV binding [80]. Exosomes can also “surf” on filopodia to enter cells at specific endocytic hot spots [81]. Molecular bases for the selective cellular targeting of EVs are poorly described [82,83]. The initial cellular binding of EVs can be mediated by adhesion proteins, and several classes of mediators are known to promote EV-cell interaction such as tetraspanins, lectins, heparan sulfate proteoglycans, and certain lipids [60,84–86]. A well-established example is the targeting of EVs (exosomes) to dendritic cells, which is mediated by tetraspanins (CD9, CD81), milk fat globule-E8/lactadherin, CD11a, and CD54/intercellular adhesion molecule 1 present on exosomes and αv/β3 integrin, CD11a, CD54 present on dendritic cells [87]. As reported for apoptotic bodies or certain viruses, phosphatidylserine present at the outer leaflet of EV membrane can play a role in their cellular entry [88–90]. We showed that monovalent antibody against CD9 can impede the
entry of EVs into melanoma cells [91]. Further efforts are needed to dissect all mechanisms regulating the specificity of EV-recipient cell interactions, which is an important step to design new strategies based on EVs as a bio-vehicle for therapeutic drug delivery [92–94].

Several mechanisms that are not mutually exclusive were proposed to explain the transfer of bioactive molecules between EVs and recipient cells (Figure 1B) [95,96]. The machinery responsible for the transmission of information and/or cellular entry of EVs might be determined by composition of EVs and the plasma membrane of recipient cells, as well as the size of EVs or their aggregation [97]. Receptor-mediated binding of EVs, or of EV-derived soluble ligands, to recipient cells could promote a downstream signaling cascade and elicit a pleiotropic response [95,98–101]. Fusion of EVs with the plasma membrane and hence the direct release of their content into cytoplasm might occur [102]. An acidic milieu as observed in cancerous tissue microenvironment might favor EV-cell fusion [103]. The acidity-mediated direct fusion of EVs with tumor cells may be similar to the fusion of endocytosed EVs with limiting late endosomal membranes observed in the low pH endocytic compartment and/or the entry of certain enveloped viruses [104–106]. Under these extreme conditions, alterations in membrane fluidity and/or unmasking of fusogenic proteins may account for the fusion ability [103]. Tetraspanin-rich microdomains and/or lipid rafts present within the EV membrane and/or the plasma membrane of recipient cells may facilitate their fusion [107,108].

In addition to these two processes that trigger a cellular response, recipient cells can internalize EVs en route to intracellular targets [109]. The literature about EV uptake by endocytic mechanisms is growing, but sometimes conflicting [110]. Various mechanisms of endocytosis were described, including clathrin-dependent, caveolae, and/or cholesterol-rich lipid rafts mechanisms (Figure 1B) [95,111–118]. Clathrin-dependent endocytosis involves the engulfment of receptors associated with their ligands in clathrin-coated pits on the plasma membrane that invaginate and form clathrin-coated vesicles. Treatments affecting the formation and/or dynamics of clathrin-coated pits can impede the EV entry [114,119]. Caveolar endocytosis is based on the properties of caveolin proteins to oligomerize, which leads to the formation of caveolin-rich microdomains within plasma membrane ending to the endocytic pathway, notably late endosomes [120]. In early days, this specialized caveolin-rich endosomal compartment was named “caveosome”, which turned out to be “artifactual” endosomes somehow produced by caveolin overexpression [120–123]. Caveolins are cholesterol-binding integral membrane proteins with unusual membrane topology where their uniquely long hydrophobic segment does not span the membrane bilayer. Other players involved in caveola architecture are members of Cavin family, i.e., peripheral membrane proteins that coat the caveolar surface [124,125]. In addition to caveolins and cavins, caveolae contain dynamin, a GTPase also involved in the biogenesis of clathrin-coated vesicles which plays a role in pinching off the caveolar vesicle [123,126]. Similar to lipid raft domains, caveolae are rich in cholesterol and sphingolipids, and membrane cholesterol is essential for their formation [127]. Cholesterol-rich lipid rafts can also promote endocytosis independent of the presence of caveolae or clathrin coats [128]. Lipid raft-associated flotillins and annexins can play a role in such endocytic pathways [129,130]. Ruffling of the plasma membrane of recipient cells can also lead to the internalization of large volumes of extracellular fluid containing EVs, a phenomenon referred to as macropinocytosis [131,132]. Cell type-specific phagocytosis is an alternative way for EV uptake. In these processes, an active remodeling of the actin cytoskeleton and the participation of phosphatidylinositol 3-kinase are essential (Figure 1B) [114,133].
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Figure 1. Mechanisms of the release of EVs by donor cell and their uptake by recipient cell. (A) The release of EVs from a healthy donor cell can occur by two major pathways that are not mutually exclusive. CD133+ EVs can be discharged into extracellular milieu upon fusion of multivesicular body (MVB) with plasma membrane. They will be referred to as exosomes. The internalization of CD133 and its transport to the early and late endosomal compartments can be mediated by its ubiquitination and interaction with syntenin-1. The biogenesis to small intraluminal vesicles within late endosome/MVB involves an inward invagination and budding of the limiting membrane of the endosome toward its own lumen (1). MVB could also fuse to lysosomes, leading to proteolytic degradation of CD133 and other proteins. The endosome movement requires a microtubule network. Alternatively, CD133+ EVs can bud from plasma membrane, notably protrusions such as primary cilium (2) or microvilli (3). They will be referred to as microvesicles or ectosomes. Cholesterol-dependent membrane microdomain (red membrane segment) can play a role in these budding processes. (B) The transfer of EV cargoes to the neighboring or distant recipient cell can occur by several mechanisms. The receptor-mediated binding of EVs to recipient cell could promote a signaling cascade, and hence elicit a pleiotropic response in recipient cells. Fusion of EVs with plasma membrane might occur, and consequently release randomly their content into cytoplasm of host cells. In addition, recipient cells can internalize EVs en route to intracellular molecular targets. Various mechanisms of endocytosis were reported. Clathrin-dependent mechanism required adaptins, which connect membrane cargo to clathrin forming a polyhedral lattice that surrounds the vesicle. Caveolin-1 (Cav1) plays a role in caveola-mediated uptake. Caveolae have a flash-shaped structure and are enriched in cholesterol and sphingolipids. Other lipid raft-mediated EV internalization processes independent of clathrin and caveolin are also proposed models of endocytosis. Phagocytosis and macropinocytosis are alternative mechanisms implicated in the internalization of EVs (see main text for details). After endocytosis, the EV content can be transferred to the cytoplasm of the recipient cells upon fusion of EVs with endosomal membrane and/or transported to nucleus (Figures 2 and 3). The nucleoplasmic reticulum can be involved in the latter process. Both types (I and II) of nuclear envelope invaginations (NEI) are depicted. In type I NEI, only the inner nuclear membrane (INM) penetrates into nucleoplasm, whereas in type II both the INM and outer nuclear membrane (ONM). The endoplasmic reticulum (ER) is an extension of the ONM, and consequently, both membranes share certain constituents.
2.3. Fate of Extracellular Membrane Vesicles upon Internalization

Despite current knowledge on the biogenesis and uptake of EVs and their relevance in various medical areas as biomarkers, therapeutic targets, or biological vehicles for drug delivery, information on the fate of endocytosed EV content remains limited [21,22]. The heterogeneity of EVs and the mechanism of cellular internalization can determine the fate of EV content. Various pathways can operate simultaneously, leading to various effects. Upon endocytosis, internalized EVs traffic to the early endocytic pathway. Little is known about the role or fate there. If a fusion occurs with the early endosomal membrane, their soluble cargo would reach the cytoplasm, while EV-associated membrane proteins could potentially travel to the trans-Golgi network, Golgi complex, and endoplasmic reticulum (ER) via retrograde transport. They can also reach the plasma membrane through recycling endosomes. To our knowledge, however, such EV-early endosome fusion was never reported. Movement of endocytosed EVs from early endosomes to the late endosomes/MVB, and again their discharge into the extracellular milieu upon MVB-plasma membrane fusion, is a potential avenue [134]. Although this recycling mechanism appears futile in regard to the fate of cargo carried by EVs, it could nonetheless allow transfer of EV-associated components through a cell en route to the proper recipient. The transcytosis through a cellular barrier such as an epithelium and/or a blood vessel might allow EVs and/or their specific content to reach distinct cells. Alternatively, EV-derived components can be sent to lysosomes for degradation [135]. Such endosomal-lysosomal degradative pathway leading to EV clearance would need further thought to understand how it could elicit a cellular response. Nonetheless, the EV loading of such degradation pathway might indirectly influence the fate of other intracellular routes (e.g., autophagy), and hence the final destination of their components [136,137].

As communication vehicles, the endocytosed EVs can fuse with late endosomes and release their soluble content directly into the cytoplasm of the host cells [138,139], while their membranous components would mix with those of endosomal membranes. The acidic pH in the late endosome microenvironment would favor such fusion, as discussed above. As recently underlined by Mathieu and colleagues, endocytosed EVs and/or degradation products therefrom can potentially escape the endocytic pathway upon rupture of endosomal/lysosomal membrane [97]. Further investigations are required to decipher the spatiotemporal breakdown of late endosomes, and how this potential mechanism can be coordinated with EV function as a messenger of information. In any case, the subcellular localization of late endosomes (e.g., peripheral areas versus perinuclear) can indirectly regulate the interaction of EV cargo molecules with cellular targets [140]. In this regard, we recently described a new intracellular pathway that led to the transfer of the EV content, notably transmembrane proteins and nucleic acids, to the nuclear compartment of host cells. Nuclear transfer of EV cargo was monitored using engineered EVs expressing CD9-green fluorescent protein (GFP) fusion protein [5]. In a separate study, Read and colleagues reported nuclear transportation of exogenous epidermal growth factor receptor (EGFR) and androgen receptor via EVs [6]. Hereafter, we will present this novel intracellular path, the key organelles involved, i.e., nucleoplasmic reticulum and late endosomes, and the molecular players involved. The perspectives that this alternative intracellular route could bring to the medical field will be highlighted.

3. Nucleoplasmic Reticulum-Associated Late Endosomes: A New Gate to Intercellular Communication

3.1. Nuclear Envelope Invaginations

To grasp the novel intracellular pathway involved in the nuclear transfer of endocytosed EV-related cargoes, we need to introduce the nucleoplasmic reticulum, which is a complex branched network of tubular membrane-bound invaginations of the nuclear envelope that allows deep nuclear structures to be in the proximity of cytoplasmic components, thus increasing the interface between cytoplasmic and nucleoplasmic compartments. Although not all aspects of its biological function are fully understood, nucleoplasmic reticulum plays a role in calcium signaling in the nuclear compartment [141,142], and consequently in all features linked to it such as gene expression [143]. Two phospholipid bilayers, the inner nuclear
membrane (INM) and outer nuclear membrane (ONM), form the nuclear envelope [144]. The narrow space between them is called perinuclear space. ONM is continuous with ER membrane and may share certain constituents (Figure 1B). At the nucleoplasmic side, a lamin-rich proteinaceous meshwork underlies the INM. Nuclear envelope invaginations (NEI) of type I are formed solely by INM penetrating in the nucleoplasm, while type II are formed by both INM and ONM (Figure 1B) [145]. The latter can form channels throughout the nucleus and contain nuclear pores [146,147]. Often, they are in close contact with nucleoli. Type II NEI are enfolded around intermediate filaments, microtubules, and possibly actin cables that are connected to the ONM, and hence link the nuclear compartment to the cellular microenvironment [145,146,148]. The presence of cytoskeleton elements within NEI might be regulated by the cellular status, just like NEI themselves [149]. In certain type II NEI, such as those detected in stressed or polyploid cells, ribosomes and mRNA translation machinery were observed therein [150], which is consistent with the continuity between ONM and ER.

The biogenesis and dynamics of NEI are observed in interphase nuclei [151]. Recently, Vaux and colleagues demonstrated that the formation of the nucleoplasmic reticulum requires the de novo assembly of new components, i.e., nascent membrane phospholipids and lamins, rather than a simple reorganization of the pre-existing nuclear envelope [152]. The protein players involved in the membrane shaping of tubular invaginations during their biogenesis, and possibly their remodeling, remain to be identified. Enzymes involved in membrane lipid metabolism and components of nuclear pores deserve particular attention in this respect [153–155]. Other potential mechanisms explaining the formation of tubular invaginations of the nuclear envelope cannot be ignored [156]. Thus, they can be created by dynamic chromatin movements resulting from the interaction of chromatin-INM-associated proteins by pulling the nuclear envelope into the nucleoplasm [157]. Alternatively, the pressure created by cytoplasmic-associated cytoskeleton elements on nuclear membrane can create type II, but not type I NEI (see below). The abundance of NEI seems linked to the status of cellular differentiation; thus, Johnson and colleagues suggested that NEI indicate the degree of cellular de-differentiation [149]. In cancers, the number of NEI is significantly elevated and can be used for their classification [158,159]. Given that NEI are widely distributed among cells under normal and pathological conditions, they deserve further attention.

### 3.2. Nucleoplasmic Reticulum-Associated Late Endosomes

In search for the mechanism regulating the transfer of biomaterials from EVs to their molecular targets in recipient cells, notably in the nuclear compartment, we observed that a fraction of EV-associated proteins, upon EV endocytosis, ends in type II NEI, suggesting that the endocytic system can be involved in nuclear transfer [5,160]. The literature has documented the presence of membrane-bound vesicles or organelles in NEI in addition to cytoskeleton elements [145,146]. From our side, we have recently demonstrated by immunogold electron microscopy and confocal microscopy [5,160] that a subpopulation of late endosomes labeled with small GTPase Rab7 was associated with type II NEI (Figure 2A–C). As observed on longitudinal cross-sections of NEI, the frequently pseudo-elongated morphology of late endosomes therein resembles a sword in its scabbard [5], which led to propose the name of “spathasome” (from “spathi/spatha” (Greek/Latin) for sword) for this dual membrane-bound structure. The configuration of the double-membrane invaginations can be observed by immunofluorescence using the ER-associated vesicle associated membrane protein (VAMP)-associated protein A (VAP-A; see below), which also labeled ONM (Figure 2C, green), and INM-associated SUN domain-containing protein 2 (SUN2, Figure 2C magenta). Of note, the appearance of these nuclear membranes along the Rab7+ late endosomal membrane, one inside the other, has some similarity to stacking dolls (known as Matryoshka dolls; Figure 2C). Such resemblance becomes striking considering that the endocytosed EVs are present in late endosomes (Figure 2D) [5,160]. Since solely a minute fraction of EV-loaded late endosomes ended up in NEI in a steady-state condition, it remains to be determined whether they are derived from STARD-related lipid transfer domain containing 3 (STARD3)* “early” or OSBP-related protein 1L (ORP1L)* “late” late endosomes, as defined by the fluid-phase cargo transport [162]. STARD3 (also known as metastatic
lymph node 64 (MLN64)) and ORP1L are two cholesterol-binding proteins. The former is a molecular tether transporting membrane cholesterol within ER–endosome contact and is involved in the egress of cholesterol from late endosomes/lysosomes to mitochondria [163–165], while the latter is part of a protein complex that includes Rab7 and regulates the minus-end transport of late endosomes [166]. Notably, neither STARD3 nor ORP1L proteins were present in NEI of melanoma cells, suggesting that late endosomes located there might represent a third subpopulation [5,160]. Additional markers are needed for their complete characterization. Given that the movement of late endosomes in nucleoplasmic reticulum is a dynamic process as observed by time-lapse video, we cannot exclude that a larger proportion of endosomes are moving there [5,160]. All the factors that govern this transfer are waiting to be discovered, as well as the limiting step it involves (see below).

**Figure 2.** Entry of late endosomes into type II nuclear envelope invagination. (A, B) The presence of membrane-bound organelles (red arrowheads) in the nucleoplasmic reticulum (arrows) and cytoplasm of melanoma FEMX-I cells is observed by Rab7 immunogold labeling (black arrowheads) as described [160]. [Anti-Rab7 rabbit monoclonal antibody D95F2, goat-anti-rabbit Nanogold (1 nm), and silver enhancement were applied]. (C) HeLa cells expressing Rab7-red fluorescent protein were double immunolabeled for VAP-A (outer nuclear membrane (ONM) marker, green) and SUN2 (inner nuclear membrane (INM) marker, magenta) [160]. A cross-section of nuclear envelope invagination (NEI) containing Rab7+ late endosomes is displayed. Note the presence of various membranous structures, e.g., the double-layered membrane of the nucleus and the membrane of Rab7+ late endosomes—the latter might contain intact endocytosed EVs. Their arrangement one inside another (M, merge) has some similarity to Matryoshka dolls. (D) Representation of type II NEI that contain late endosomes. Some late endosomes are depicted with endocytosed EVs, which themselves contain cargo molecules. An intact microtubule network is essential for the translocation of late endosomes in a NEI that often lies close to nucleolus. A lamin-rich proteinaceous meshwork underlies the INM. Cross-sections (Cs) and longitudinal (Ls) sections of NEI. Nu, nucleoplasm. [Fluorescence images were originally published in the Journal of Biological Chemistry: Santos MF, Rappa G, Karbanová J, et al. VAMP-associated protein-A and oxysterol-binding protein-related protein 3 promote the entry of late endosomes into the nucleoplasmic reticulum. 2018; 293:13834–13848 [160]. ©The American Society for Biochemistry and Molecular Biology].
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Figure 3. VOR complex is involved in the translocation of late endosomes into the nucleoplasmic reticulum and nuclear transfer of EV-derived components. (A) ORP3, a member of the VOR complex, is associated with nuclear envelope invagination (NEI). Melanoma FEMX-I cells expressing Rab7-red fluorescent protein were double immunolabeled for ORP3 and SUN2 as described [160]. 3D reconstruction of three optical sections (green slice) of a given cell is shown. The protein of interest within NEI is indicated (arrowhead). Nu, nucleoplasm. (B) Schema summarizing the players of VOR complex and their interactions within NEI. In this model, a late endosome containing endocytosed EVs (EV) is transported to NEI and tethered to outer nuclear membrane (ONM) (i). The latter process is mediated by the type II transmembrane, endoplasmic reticulum (ER)/ONM-associated VAP-A protein (yellow), cholesterol-sensor ORP3 protein (green) that binds to VAP-A via its FFAT motif, and late endosome-associated Rab7 (gray). The pleckstrin homology (PH) domain of ORP3 might mediate its binding to late endosomal membrane. The inhibition of transfer of EV-derived components (soluble and membranous) into the nucleoplasm of host cells after importazole treatment suggests that importin α1/β1 (may be those associated with EV; black spot) and nuclear pores play a role in these processes (ii). The potential interaction of nuclear pore components with VOR complex should be addressed in a near future (green question mark). Other urgent questions remain open, notably the mechanism(s) allowing the extraction of EV-derived membrane proteins (red) from endosomal membrane and their transfer into nucleoplasm through the nuclear pores, which are size restricted (iii, black question mark). The low pH milieu in late endosome might favor the fusion of endocytosed EVs with its membrane (i–iii). [Fluorescence images were originally published in the Journal of Biological Chemistry: Santos MF, Rappa G, Karbanová J, et al. VAMP-associated protein-A and oxysterol-binding protein-related protein 3 promote the entry of late endosomes into the nucleoplasmic reticulum. 2018; 293:13834–13848 [160]. ©The American Society for Biochemistry and Molecular Biology].

In contrast to late endosomes, early Rab5+ endosomes were barely detectable in the NEI, as was the Golgi apparatus [5]. Mitochondria remained in most cases at the entrance of the NEI [5,145,146,167].
Under particular situations, mitochondria might shuttle in NEI and influence Ca\(^{2+}\) signaling \([141,168]\). It will be important to further dissect the types of membrane-bound structures that navigate in NEI, particularly in those making tunnels throughout the nucleus. Similarly, it would be of general interest to decipher whether subtypes of type II NEI with specific function(s) exist. The presence of inositol 1,4,5-trisphosphate (IP\(_3\)) receptors in certain NEI suggests it \([141,161]\).

### 3.3. Transport of EV-Loaded Late Endosomes in Nucleoplasmic Reticulum

The mechanism regulating the translocation of late endosomes in type II NEI is an interesting topic that requires further exploration. The movement of late endosomes within the cytoplasmic compartment and their interactions with the ER membrane might be instructive. For instance, the integral ER-anchored protein protrudin contacts late endosomes via Rab7 binding and mediates their plus end-directed translocation along microtubules toward the cell periphery by the intermediate of the motor protein kinesin-1 and motor adaptor protein called FYCO1 (FYVE and coiled-coil domain containing 1) located on late endosomes \([169,170]\). The repeated contacts between late endosomes and ER membrane promote their translocation in a microtubule-dependent manner. Does a similar interplay mechanism occur in NEI? The presence of tubulin in type II NEI and the negative impact of nocodazole treatment on late endosome translocation to NEI support this hypothesis (Figure 2D) \([160]\). It might be more than a coincidence that protrudin binds via its FFAT motif (FFAT being an acronym for two phenylalanines (FF) in an Acidic Tract) to type II membrane protein VAP-A, and the knockdown of the latter impeded the subcellular localization of Rab7\(^{+}\) late endosomes in NEI of melanoma cells \([160]\).

Although protrudin is not detected in all cell types including melanoma cells \([160]\), the interaction of Rab7, Rab7 effector FYCO1 protein, and motor protein kinesin-1 could nevertheless mediate the nuclear translocation of late endosomes towards microtubules, as reported for the transport of autophagic vesicles \([171]\). As described for the positioning of late endosomes, an alternative mechanism involving the interactions of Rab7, ORP1L, Rab-interacting lysosomal protein (RILP), and dynactin subunit p150\(^{Glued}\), which binds to dynein heavy chain, could also explain the movement of late endosomes into NEI \([140,172]\). Although the cholesterol-sensor protein ORP1L, which interacts with VAP-A, is absent from NEI as mentioned above \([160]\), other OSBP-related proteins of this large family could be involved \([173]\). Indeed, the OSBP-related protein-3 (ORP3), in contrast to STARD3 and ORP1L, was found in NEI and co-localized with Rab7\(^{+}\) late endosomes (Figure 3A) \([160]\). ORP3 knockdown, just like VAP-A, abolished the presence of Rab7\(^{+}\) late endosomes in NEI, indicating that this cholesterol-sensor protein could play a certain role in the late endosomes transport process and/or their tether to nuclear membrane within the NEI \([160]\). The next step will be to determine whether RILP and p150\(^{Glued}\) protein are associated with NEI and participate to the selective microtubule minus-end dynein-dynactin-dependent movement of late endosomes. A competition between FYCO1-kinesin and RILP-dynein-dynactin complexes for Rab7 binding and recruitment to late endosomes is conceivable.

The observation that, upon exposure to EVs, the proportion of cells harboring nucleoplasmic reticulum-associated late endosomes increased, and the proportion of cells without NEI decreased \([5]\), suggests that the EV loading of the endosomal compartment regulates NEI biogenesis and consequently represents a new “pushing in” mechanism of their formation \([5]\). Thus, the translocation of late endosomes into NEI and the biogenesis of nuclear membrane folds can be two linked events involving microtubules. In addition to the presence of microtubules in NEI, microtubules running parallel to the nuclear envelope and contacting the cytoplasmic filaments of a nuclear pore complex have been observed by electron microscopy \([174]\). Nucleoporin proteins are indeed present in NEI containing EV-loaded Rab7\(^{+}\) late endosomes \([5]\). Nuclear pore proteins such as nucleoporin 358 (Nup358/RANBP2) and/or nuclear proteins such as SUN1/2 together with member(s) of the ONM-associated Nesprin protein family could anchor microtubules and somehow regulate NEI dynamics \([175,176]\). It is documented that SUN1/2 and Nesprin proteins play an important role in the nuclear envelope organization and nuclear positioning relative to the cell body \([177–180]\). By interacting with motor proteins (e.g., kinesin proteins such as KIF5B and KIF5C), they could anchor the microtubules within
NEI, leading to a highway for the late endosome translocation [181]. These exciting issues await further investigation.

4. The VOR Complex: Interaction of VAP-A, ORP3, and Rab7

4.1. Inter-Organelle Contacts

The regulated interaction between membrane-bound organelles is an exciting and emerging field of cell biology [182,183]. The organelle crosstalk in eukaryotic cells plays an underestimated role in cell signaling and/or membrane dynamics. Exchange of biocomponents such as ions and lipids between connected organelles were described [184,185]. As a general controller of inter-compartmental activities, the ER, the largest organelle that forms a tubular network, appears to be determinant in numerous processes [186,187]. Contact sites between ER and Golgi apparatus, mitochondria, plasma membrane, peroxisomes, lysosomes, lipid droplets, and endosomes were documented [188–190]. Usually, the ER makes multiple contacts with a given organelle, bridging it in close proximity (i.e., less than 30 nm distance). As demonstrated by electron tomography, the distance between ER and late endosomes at the contact site is approximately 8 nm [191]. This short distance between connected organelles facilitates the non-vesicular membrane lipid transport, which requires specific proteins that can extract them from donor membrane, shield in a hydrophobic pocket, and transfer to opposite acceptor membrane. For mechanistic details about lipid transfer between opposite membranes and the involved lipid-transfer proteins, see the following reviews [173,187,192]. The repeated contact between organelles can also regulate the movement of one of them as discussed above, where late endosomes are translocated toward the cell periphery using ER and microtubules as support.

A certain number of molecular players involved in the contact zones between ER and various organelles were identified [186,193]. For instance, integral ER membrane-localized VAP-A and VAP-B (also known as ALS8) have been implicated in tethering and maintenance of structural and functional properties of the Golgi complex by regulating the proper levels of key lipids (e.g., phosphatidylinositol-4-phosphate, sphingomyelin, and diacylglycerol) in the Golgi membrane [194]. The knockdown of VAPs exerts various lipid-mediated effects on Golgi-mediated transport events. These actions are regulated by the interaction of VAPs with lipid-transfer/binding proteins, including PYK2 N-terminal domain-interacting receptor 2 (Nir2), OSBP, and ceramide-transfer protein (CERT). Their FFAT motif mediates the VAP interaction [195–197], while pleckstrin homology (PH) domain of VAP-interacting proteins promotes their binding to Golgi membrane [198,199]. FFAT-binding site in VAPs is associated with the major sperm protein (MSP) homology domain located at the cytoplasmic N-terminal part [200]. In addition to OSBP, the interaction of full-length ORP9 with VAPs also occurs and regulates trans-Golgi/trans-Golgi network structure and function by modifying its membrane environment by means of cholesterol delivery [201]. The contact between VAP-B and outer mitochondrial membrane protein, such as protein tyrosine phosphatase-interacting protein 51 (PTP1P51), regulates the ER-mitochondria association [202,203]. Direct contact between ER and plasma membrane was also described and implicated in various cellular processes such as regulation of intracellular calcium, lipid traffic, and signaling [204,205]. Among the scaffolding proteins involved, VAP-A and ORP3 were described [206]. Interestingly, ORP3 forms a physical complex with the small GTPase R-Ras [207]. The plasma membrane-targeting PH domain as well as VAP-A-interacting FFAT motif found in ORP3 were essential for the activation of R-Ras and its downstream signaling pathway, indicating that ER-plasma membrane contact creates a R-Ras activation platform [208]. Moreover, ORP3 interacts with IQSec1 (IQ motif and Sec7 domain-containing protein 1) protein, and the ORP3/VAP-A/IQSec1 complex plays a role in focal adhesion turnover [209]. Like other OSBP homologs, ORP3 was shown to bind to membrane cholesterol, as demonstrated using live cell photo-crosslinking with \(^{3}H\) photo-cholesterol [210]. The endosomes are other organelles physically involved in the contact with ER. Indeed, ER participates actively in endosome maturation, transport, and fission [211–213]. The literature has documented various molecules implicated in the ER-endosome
tethering, including paired proteins: VAP-A and STARD3 or STARDNL (STARD3 N-terminal like); VAP-A and ORP1L; protrudin/VAP-A and Rab7; and protein tyrosine phosphatase 1B (PTP1B) and endocytosed phosphorylated EGFR [193,214,215]. In the latter case, such inter-organelle contact (i.e., ER–endosomes) can regulate spatially and temporally EGFR signaling [216]. The interaction between VAP-A and endosomal proteins STARD3 and STARDNL could regulate endosome positioning and dynamics. In contrast to STARD3, STARDNL does not contain a START protein domain that allows cholesterol binding and can potentially transfer cholesterol between connected and adjacent membranes [217,218].

4.2. Late Endosome–Nuclear Membrane Contact: Role of VAP-A–ORP3–Rab7 Interactions

Applying a triple fluorescence analysis, fluorescence resonance energy transfer, and para-magnetic immune-isolation techniques, we demonstrated that VAP-A, ORP3, and Rab7 co-localize in discrete areas within NEI and interact with each other [160]. This tripartite complex, named VOR complex (an acronym for VAP-A, ORP3 and Rab7), is essential for the localization and docking of late endosomes in NEI (Figure 3B) [160]. Moreover, ORP3 needs VAP-A to be present in NEI, suggesting that VAP-A is orchestrating these interactions [160]. Surprisingly, the VAP-A homolog, VAP-B, is not involved in these events although it is also localized in nucleoplasmic reticulum [160]. Thus, VAP-A silencing cannot be rescued by VAP-B [160]. These findings highlight the most hidden part of molecular mechanisms controlling these sequential interactions (VAP-A/ORP3/Rab7) and their regulation. The hyperphosphorylation of ORP3 might promote the interaction with VAP-A via the FFAT motif as described for its role in ER-plasma membrane contact zone, while its PH domain could promote its interaction with late endosomal lipids [208,209,219]. In contrast to ORP1L, ORP3 does not contain an ankyrin repeat region, which could mediate Rab7 interaction [220]. It remains to be evaluated whether the R-Ras binding site in ORP3 can be involved [208]. Although the presence of Rab7 late endosomes in NEI is fully abrogated after ORP3 silencing in melanoma cells [160], it will be of interest to evaluate whether other ORPs are engaged in the nucleus—late endosome contact in particular types of cells and/or under peculiar physiological and pathological conditions. A potential candidate is the ER-associated transmembrane protein ORP5, which has been suggested to interact with the late endosome cholesterol transporter Niemann-Pick C1 protein [221] (reviewed in [215]). The co-existence of multiple bridging complexes driven by distinct protein pairs cannot be excluded [204], and it remains to be identified whether other proteins participate in the tether of late endosomes with ONM within nuclear folds.

4.3. Implication of Nuclear Pores in EV-Derived Cargo Nuclear Shuttling

How can EV-associated components within late endosomes reach the nucleoplasm? After the docking of late endosomes to ONM, the fusion of EVs with the endosomal membrane might expose EV content in the proximity of nuclear pores, allowing their nuclear transfer. The presence of the nuclear transport receptor karyopherin (importin α and β1 subunits) in EVs [5,222–225] suggests that it may somehow participate in the nuclear translocation of soluble and perhaps membranous EV components (Figure 3B). The nuclear localization signal (NLS) present on a protein cargo is recognized by importin α1 subunit, which in turn interacts with β1. The resulting protein complex passes through the nuclear pore by binding key nucleoporins [226]. Once translocated into nucleoplasm, small regulatory nuclear GTPase Ran-GTP binds to importin β1 and dissociates the imported protein complex. Consistent with the role of nuclear pores in these processes, nucleoporins were detected in NEI containing late endosomes [5], and the incubation of cells with cell-permeable 2,4-diaminoquinazoline (importazole), a small molecule that inhibits the function of importin β1 by altering its interaction with Ran-GTP [227], impaired the nuclear translocation of EV-derived protein cargo [5]. The intriguing interaction of another ER-located OSBP-related protein, i.e., ORP8, with nucleoporin Nup62 might be relevant in this context [228]. A direct interaction between ORP3, together with VAP-A/Rab7, and nuclear pore constituents should be investigated (Figure 3B, question mark). Further, it remains to be determined...
how a protein complex notably those containing EV-derived membrane glycoproteins (e.g., full-length CD9 and CD133) are extracted from EV-endosomal membrane after the fusion of the former with the endocytic organelle [5]. Are tetraspanin-rich microdomains or lipid rafts present in the membrane bilayer of EVs involved in forming a phase separation with surrounding, more fluid membrane lipids (Figure 3B, double question marks)? Thus, tetraspanin/lipid raft-associated proteins might create a membranous platform, and NLS-containing transmembrane cargo/importin proteins could be extracted and drag the entire complex through the nuclear pore. Considering that nuclear pores are size restricted [229], we cannot expect that an entire endocytosed EV (>40 nm in diameter) reaches the nucleoplasm, although it was recently suggested that very small (nano)vesicles (<30 nm) other than high-density lipoprotein particles may exist [230]. The high degree of membrane curvature in NEI, particularly at their tip may alter the properties of nuclear pores and hence allow larger complexes to pass through. It remains to be investigated whether other pathways independent of nuclear pores are implicated in the nuclear translocation of EV-derived cargoes [231,232].

5. Biological Functions

The localization of late endosomes in the nucleoplasmic reticulum could have a significant impact on the mechanism of intercellular communication mediated by EVs. This novel endosomal pathway adds an alternative route to the soluble and membranous components derived from EVs (e.g., nucleic acids and proteins) to reach and interact with their molecular targets, including those associated with nucleoli [5,6]. This biological issue is particularly important given the limited amount of bioactive molecules carried by EVs. In addition to EV cargo, this pathway might play a role in the nuclear translocation of plasma membrane receptors upon ligand interaction. Similar to integral membrane proteins CD9 and CD133 [5,52,233–235], numerous plasma membrane receptors including EGFR, platelet-derived growth factor (PDGF), insulin growth factor receptor 1 (IGF-1R), and several G-protein coupled receptors that are potential drug targets were surprisingly found in the nuclear compartment [236–239]. Therein, they could be involved in transcriptional regulation and cellular proliferation, and confer chemo- and radio-resistance among other biochemical processes. In this context, it might be more than a coincidence that the nuclear translocation of the cell surface IGF-1R is dependent on the dynactin subunit p150\textsuperscript{Glued} and importin β1 as discussed above for the late endosome movement and nuclear transfer of EV cargo, respectively [238]. To date, the retrograde transport through Golgi and ER was considered the main intracellular route for full-length membrane-anchored proteins to the nucleus [240,241] (Table 1). Our observation with nucleoplasmic reticulum-associated late endosomes is in line with a recent pathway described by Chaumet and colleagues where the bacterial protein Pseudomonas exotoxin A upon binding to the cell surface LDL receptor related protein 1 (LRP1) can reach the nuclear compartment through docking and membrane fusion of early endosomes with the nuclear envelope [232] (reviewed in [242]). In the latter case, SUN1 and SUN2 proteins and Sec61 translocon complex were involved in this novel pathway (Table 1). Sec61 complex mediated the egress of proteins from INM to the nucleoplasm [232,242]. Although the mechanisms regulating these novel intracellular trafficking routes appear distinct, they nonetheless address the question of how cell surface and extracellular proteins reach the nucleoplasm. Potential mechanisms underlying the nuclear transports of membrane proteins present at the plasma membrane and/or EVs are summarized in Table 1.

In addition to the nuclear transfer of EV cargo, nucleoplasmic reticulum-associated late endosomes may create a privileged intracytoplasmic compartment that would favor protein–protein, protein–nucleic acid, and nucleic acid–nucleic acid interactions. The cytoplasmic release of soluble EV constituents upon a tardive fusion of intact endocytosed EVs with late endosomal membrane might facilitate their binding, given a local high concentration, to the molecular targets exported out of the nucleus such as RNA transcripts, and hence exert a function independent of their nuclear import [150]. This issue requires further attention.
Table 1. Potential pathways of nuclear transport of cell surface proteins present at plasma membrane and/or extracellular membrane vesicles.

| Retrograde Transport | Nuclear Envelope-Associated Early Endosomes | Nuclear Envelope Invagination-Associated Late Endosomes |
|---------------------|--------------------------------------------|-----------------------------------------------|
| **PLASMA MEMBRANE** |                                            |                                               |
| **ENDOSOMAL COMPARTMENTS** | Early endosomes | Early endosomes | Early endosomes |
| Golgi apparatus | Late endosomes |                                        |
| Endoplasmic reticulum (ER) | a) cytoplasmic translocation | b) diffusion from ER to nuclear membrane | a) docking to nuclear pore | b) fusion with nuclear membrane | entry in nucleoplasmic reticulum |

Implication of Nuclear Pores

- YES
- Implication of Importin β1
- YES (a)/unclear (b)
- Fusion of Endosome with the Outer Nuclear Membrane
- NO (a)/YES (b)
- Not determined

Implication of Sec61 Translocon Complex

- YES
- unclear (a)/YES (b)
- Not determined

[240,241,243–246] [232,236,238,242,246] [5,91,160]

6. Perspectives and Future Directions

Interference with the new intracellular route used by endocytosed EVs can find applications in the cancer field, for instance by preventing the delivery of cancer cell-derived EV cargo to host cell nucleoplasm. The identification of the VOR complex and its inhibition may be clinically relevant. Thus, disruption of the intercellular communication between cancer cells and stromal components within the metastatic niche may be a novel approach to intercept the mechanisms underlying tumor growth and metastasis formation. A drug screening should be undertaken and the knowledge about the interaction of VAP-A with FFAT motif-containing protein partners, including sterol-binding proteins, would be useful to design such chemical compounds. In addition to cancers, this new pathway may be involved in the delivery of biomaterials (e.g., cell surface receptors) to the nuclear compartment as explained above, and hence other applications can be envisaged.

Although EVs do not have the ability to promote their own replication in host cells unlike enveloped viruses, numerous facets are nonetheless shared between these small phospholipid membrane-enclosed entities, including certain aspects of their biogenesis and release into the extracellular milieu (reviewed...
in Refs [247–249]). The similarity between them is further supported by discoveries revealing that virus-infected cells release modified EVs (i.e., exosomes) containing viral proteins and nucleic acids such as non-coding regulatory miRNAs [250], which could enhance or interfere with infection and/or provoke immune dysregulation [251]. These observations among others have highlighted that the exosomal and/or microvesicle pathways can be hijacked by viruses, notably retroviruses [252]. Moreover, EVs can provide an “envelope” to non-enveloped viruses such as hepatitis, which promotes their spreading without lysis of infected cells [253,254]. The latter observation might also question a “dogmatic” issue that categorizes viruses into enveloped and non-enveloped ones. The internalization of EVs and enveloped viruses might also share some similarity [255]. Direct fusion with the host cell plasma membrane or receptor-mediated endocytosis can explain the entry mechanisms of both particle types (reviewed in [256]). Once internalized into the endocytic pathway, viral components can reach the nuclear compartment, which raises the possibility that they might use the nucleoplasmic reticulum-associated late endosomes to transfer their nucleic acids to the nucleoplasm of host cells. It might be more than a coincidence that the infection of certain viruses (e.g., herpesvirus) could promote morphological alterations of the nuclear membrane of host cells and a redistribution of nuclear pore proteins, notably in NEI-like structures [257]. The interaction of viral proteins with nucleoporins might be significant in these processes [258]. Such nuclear membrane re-organization of infected cells is in line with the impact of EVs on the number of NEI [5], hence the spatial configuration of the nucleus of EV-exposed cells. Therefore, it will be of interest to revisit the cellular entry of enveloped viruses and the delivery of their viral genome into nucleus in light of the shuttling of EV cargo described here.

Lastly, the discovery of nucleoplasmic reticulum-associated late endosomes might open new avenues to deliver and target specific drugs to nuclei of cancer cells or other pathologic cells. Altogether, deciphering at a molecular level all spatiotemporal multi-steps of this new intracellular pathway would benefit the field of nanobiological technology towards new medical therapeutic approaches [91].

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