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Role of host reticulon proteins in rearranging membranes for positive-strand RNA virus replication
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Positive-strand RNA [(+)RNA] viruses are responsible for numerous human, animal, and plant diseases. Because of the limiting coding capacity of (+)RNA viruses, their replication requires a complex orchestration of interactions between the viral genome, viral proteins and exploited host factors. To replicate their genomic RNAs, (+)RNA viruses induce membrane rearrangements that create membrane-linked RNA replication compartments. Along with substantial advances on the ultrastructure of the membrane-bound RNA replication compartments, recent results have shed light into the role that host factors play in rearranging these membranes. This review focuses on recent insights that have driven a new understanding of the role that the membrane-shaping host reticulon homology domain proteins (RHPs) play in facilitating the replication of various (+)RNA viruses.

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Introduction
(+)-RNA viruses reorganize intracellular membranes to assemble their RNA replication compartments, which are intricate factories or mini-organelles featuring the close association of both viral and host components [1]. Cellular lipid synthesis as well as appropriate lipid composition is required for viral replication, implying that the membrane is an essential, functional component of the RNA replication complex [2]. Pioneering ultrastructural studies using three-dimensional imaging by electron microscope tomography (EMT) have substantially enhanced the understanding of the RNA replication compartments induced by (+)RNA viruses [3–6]. As noted below, various (+)RNA viruses induce membrane rearrangements with distinct morphologies. Nevertheless, in general, the membrane is thought to provide a surface to help localize and increase the local concentration of components required for viral replication [1], protect dsRNA replication intermediates and viral proteins from sensors of the innate immune system [7], and in some cases provide host factors that help target viral proteins or genomic RNAs to the appropriate membrane [8].

Host factors have been shown to be involved in every step of viral life cycles [9,10–12]. In keeping with this, recent results show that host factors also are key determinants in forming viral RNA replication compartments, including by generating appropriate lipid composition of the membranes [2,13,14,15,16,17] and by structurally inducing/supporting relevant membrane curvature [17,18,19]. Since genomic RNA replication is a highly conserved step in the viral life cycle, understanding the role that such host factors play in the formation and function of the viral-induced RNA replication compartments could provide a target to develop broad-spectrum antivirals.

Reticulon homology domain proteins
The reticulons are a group of morphogenic, ER membrane-shaping proteins that partition to and stabilize highly curved ER membrane tubules [20], and as such are candidates for host factors that may affect the generation and/or maintenance of virus-induced membrane rearrangements. The diversity of the reticulon gene family varies among hosts, with four reticulon genes in mammals (RTN1-4), two genes in yeast (RTN1 and RTN2), and 21 and 17 genes, respectively, in the plants Arabidopsis thaliana and Oryza sativa [21,22,23,24]. A related protein family that is also necessary for ER tubule formation consists of six mammalian DP1/REEP proteins and yeast Yop1 [20,25]. The reticulon and DP1 families share little overall homology but do exhibit an important common structural feature—two elongated hydrophobic segments that appear to form partially membrane-spanning hairpin domains [20,26,27]. For simplicity, for the rest of this review we will refer to the reticulon and DP1 families as the reticulon homology domain proteins (RHPs). The bulk of the hydrophobic portions of the RHPs is predicted to reside in the outer leaflet of the phospholipid bilayer, thus inducing curvature by hydrophobic wedging [20]. The oligomerization of the RHPs
is important both for their ability to localize to and to form tubular ER domains, suggesting that such scaffolding might also have a role in curvature induction and stabilization [28]. Consistent with their role in tubule formation, RHPs primarily partition to and stabilize peripheral ER membrane tubules while avoiding the low-curvature ER domains of the nuclear envelope and peripheral ER sheets [20*,29]. Such ER tubules are said to have positive membrane curvature — that is, curved to protrude into the cytoplasm (Figure 1a).

**Role of RHPs in brome mosaic virus RNA replication complex formation**

The replication cycle of brome mosaic virus (BMV), a small alphavirus-like virus, has been well characterized, making BMV an advantageous model to study some of the general features of (+)RNA virus RNA replication [30]. EM shows that for BMV, RNA synthesis localizes to 50–80 nm vesicular compartments, also known as spherules, in which the outer perinuclear ER membrane is invaginated away from the cytoplasm and into the ER lumen [31]. The interior of these vesicles is connected to the cytoplasm through a neck-like opening, providing a channel for ribonucleotide import and product RNA export. The only BMV protein necessary to induce spherule formation is replication factor 1a [31]. The self-interacting 1a protein is membrane associated, localizes to spherule interiors, and is present at high enough levels to imply that it must form a shell covering much of the spherule interior [31–33]. Such a rigid protein shell...
could contribute substantially to forming and maintaining the high-energy membrane deformation state of the spherules (Figure 1b). However, in addition to 1a, RHPs also were recently shown to play key roles in BMV RNA replication compartment formation and function [19**].

Using a loss-of-function experimental approach, RHPs were shown to be required for efficient BMV RNA replication in yeast [19**]. In keeping with the altered RNA replication phenotype, BMV 1a interacts with and incorporates RHPs into the spherule interior, and deleting all three yeast RHPs abolishes 1a-induced spherule formation [19**]. Topologically, the spherule body has negative curvature in that the membrane of the vesicle sunshine is curved away from the cytoplasm, although the neck has positive curvature and can be thought of as a tubule cut in half (Figure 1c). RHPs polymerize into short arcs to form perfect cylindrical tubes [34] and are involved in forming nuclear pores [35], which are topologically equivalent to spherule necks. Thus, RHPs are likely necessary for both spherule formation and maintenance of an open channel to the cytoplasm to facilitate import of RNA templates and nucleotides and export of progeny RNA (Figure 1d). In addition to stabilizing spherule necks, a subset of RHPs must also be incorporated within the vesicular portion of spherules as partial RHP depletion results in spherules with reduced diameter [19**]. In this instance, the positively curving RHPs might partially cancel the negative membrane curvature induced by 1a, allowing expansion of the spherule body (Figure 1d). Although RHPs are involved in shaping smooth ER [36], the site of many lipid synthesis steps [37], RHP deletion did not alter fatty acid levels or composition, suggesting that their main contribution to RNA synthesis is in RNA replication compartment formation. Further supporting this notion, RHP-GFP fusions retain 1a interaction but shift 1a-induced membrane rearrangements from spherules to double-membrane layers of ER surrounding the nucleus [19**].

**Role of RHPs in Picornaviridae RNA replication**

The importance of RHPs in (+)RNA virus replication was first demonstrated when human reticulon 3 (RTN3) was found to interact with the 2C proteins of poliovirus, coxsackievirus A16 (CA16) and enterovirus 71 (EV71) [38]. Moreover, immunoprecipitation showed that EV71 2C protein interacts with the reticulin homology domain of RTN1C, RTN2, and RTN4. siRNA knockdown of RTN3 resulted in inhibited synthesis of viral double-stranded RNA (dsRNA) and proteins in EV71-infected cells [38]. Ectopic expression of a form of RTN3 that was not degraded by siRNA in siRTN3 knockdowns rescued EV71 infectivity, suggesting that RTN3 plays a key role in EV71 replication. Although ultrastructural studies were not performed, immunofluorescence showed that endogenous RTN3 colocalized not only with EV71 2C but also with dsRNA, implying that RTN3 is likely associated with the viral replication compartment [38]. The 2C protein is one of the most highly conserved viral proteins among all picornaviruses and poliovirus 2C has been proposed to be involved in the formation of the vesicle-associated replication compartments [39].

Because of the limitations of conventional, two-dimensional electron microscopy, the RNA replication structures induced by poliovirus and other enteroviruses had been previously described as either single-membrane [40] or double-membrane vesicles (DMVs) [41]. However, EM tomograms of enterovirus-infected cells showed that, during the exponential phase of viral RNA synthesis, closed smooth single-membrane tubules constitute the predominant virus-induced membrane structure, whereas DMVs become increasingly abundant at the expense of these tubules as infection progresses [42*]. Similar observations have been made in poliovirus-infected cells. Early in infection, poliovirus induces the formation of single-membrane connecting and branching tubular compartments [43*]. As infection progresses, the tubular compartments gradually transform into double-membrane structures [43*]. Unlike the negatively curved spherular compartments induced by BMV, the enterovirus71-induced and polio-virus-induced tubule-like and vesicle-like structures are both positively curved (Figure 1c and d). Thus, RHPs may play a role in forming the single-membrane tubules, analogous to their normal role in shaping smooth ER [36], whereas later on they would stabilize the positively curved DMVs. The transformation from single-membrane tubules to DMVs would require both close apposition of the inner tubule membrane with a source for the outer membrane and the induction of curvature to wrap this outer membrane around the tubules, which is another role that RHPs might be involved in.

**Potential RHP role in forming replication compartments of other (+)RNA viruses**

A yeast-two hybrid system used to screen a liver cDNA library showed that RTN3 interacts with the NS4B protein of hepatitis C virus (HCV) [44]. When expressed individually or in the context of the entire HCV polyprotein, NS4B localizes to the ER [45]. Moreover, NS4B is a key organizer of the HCV RNA replication compartment by inducing the formation of ER-derived membranous vesicles, which accumulate in large cytosolic clusters, referred to as the membranous web [46]. Although the role that RTN3 plays in HCV RNA replication was not investigated further in these studies, given the particular functions of NS4B in inducing HCV RNA replication compartments, it is tempting to speculate that RTN3 might be involved in modulating the formation of the membranous web.
Dengue virus (DENV) infection induces several distinct ER-derived membrane structures, including vesicle packets (VPs) and convoluted membranes (CMs) [5,47]. EM tomography shows that the inner vesicles of VPs are invaginations of the outer ER membrane [5], thus, the inner vesicle interiors are connected to the cytoplasm by neck-like pores similar to those in the spherules induced by BMV and many other (+)RNA viruses. Likewise, SARS coronavirus-infected cells contain a mixture of VP and CM rearrangements [4]. However, tomographic reconstructions failed to reveal any membrane openings between the SARS virus-induced DMV interiors and the cytoplasm, as was observed for BMV and DENV, suggesting that the DMVs might be more related to the DMVs formed by poliovirus. On the basis of such ultrastructural data, alpha-like and some flavivirus-like viruses appear to induce structures with negative membrane curvature along the dome of the vesicle [5], with areas of positive curvature along the neck, while picorna-like and coronavirus-like viruses induce structures with positive membrane curvature [4,42,43]. Despite the topological differences, one common theme might be the role of RHPs in forming the various RNA replication compartments, perhaps not necessarily limited to those viruses that use membranes derived from the ER.

Concluding remarks

With RHPs now identified as playing crucial roles in the replication of multiple (+)RNA viruses, the stage is set to identify not only additional participating host factors, but also the detailed molecular mechanisms of how different viruses orchestrate the varied and often complex membrane rearrangements associated with their replication processes. Moreover, the role of RHPs in (+)RNA replication aside from genome amplification remains to be explored, as do the potential interactions of RHPs with other cellular pathways that contribute to virus replication, such as factors that regulate membrane synthesis/composition [2], trafficking, and membrane remodeling. Although mainly found in the ER, RHPs also localize to the Golgi and plasma membrane, implying a potential role in membrane trafficking [48]. Along these lines, poliovirus vesicle formation has been linked to COPI vesicle trafficking [49], tomsbusvirus (TBSV) recruitment of another set of membrane-shaping proteins ESCRT (endosomal sorting complexes required for transport) proteins apparently facilitates the assembly of TBSV-induced spherules [18] and enteroviruses and flaviviruses exploit host phospholipid-modifying enzyme Pl4KIIIβ and replicate their respective viral RNA on phosphatidylinositol-4-phosphate lipid-enriched membranes [50*]. Cryo-EM tomography and super resolution imaging should help to resolve the three-dimensional organization at molecular resolution of viral and host proteins that are part of replication compartments and to enhance our understanding of the role that RHPs and other host factors play in forming the membrane-associated RNA replication compartments.

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