Legionella Subvert the Functions of Rab1 and Sec22b to Create a Replicative Organelle

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Abstract

*Legionella pneumophila* is a bacterial pathogen that infects eukaryotic host cells and replicates inside a specialized organelle that is morphologically similar to the endoplasmic reticulum (ER). To better understand the molecular mechanisms governing transport of the *Legionella*-containing vacuole (LCV), we have identified host proteins that participate in the conversion of the LCV into a replicative organelle. Our data show that Rab1 is recruited to the LCV within minutes of uptake. Rab1 recruitment to the LCV precedes remodeling of this compartment by ER-derived vesicles. Genetic inhibition studies demonstrate that Rab1 is important for the recruitment of ER-derived vesicles to the LCV and that inhibiting Rab1 function abrogates intracellular growth of *Legionella*. Morphological studies indicate that the Sec22b protein is located on ER-derived vesicles recruited to the LCV and that Sec22b is delivered to the LCV membrane. Sec22b function was found to be important for biogenesis of the specialized organelle that supports *Legionella* replication. These studies demonstrate that *Legionella* has the ability to subvert Rab1 and Sec22b function to facilitate the transport and fusion of ER-derived vesicles with the LCV, resulting in the formation of a specialized organelle that can support bacterial replication.

Key words: brefeldin A • ARF1 • Sar1 • phagosome • endoplasmic reticulum

Introduction

After uptake by a eukaryotic cell, the bacterial pathogen *Legionella pneumophila* has the ability to modulate transport of the vacuole in which it resides. The initial compartment in which virulent *Legionella* resides is derived from the plasma membrane and we refer to it as the *Legionella*-containing vacuole (LCV). Unlike avirulent microorganisms, which typically reside in vacuoles that interact sequentially with endocytic vesicles resulting in their delivery to lysosomes, the LCV avoids endocytic maturation and intercepts secretory vesicles as they exit the ER (1–5). To modulate transport of their vacuole, *Legionella* requires a specialized secretion apparatus called the Dot/Icm system, which injects bacterial proteins into the cytosol of host cells during infection (6–9). The functions of most bacterial proteins injected by the Dot/Icm system and the host factors that participate in LCV transport remain unknown.

The LCV transport pathway has been defined primarily by morphological studies. Fluorescence microscopy indicates that the LCV diverges from the default endocytic pathway within the first 5 min of infection (4). Although endocytic markers are excluded (10, 11), proteins residing in secretory vesicles cycling between the ER and Golgi apparatus are associated with most LCVs within 30 min, and resident ER proteins such as calnexin are found associated with the LCV within 1–2 h (3). Vacuoles containing dot/icm mutants of *Legionella* do not stain positive for secretory proteins or ER markers (3, 12–14). Electron microscopy has been used to confirm and extend observations made by fluorescence microscopy. Electron micrographs show that ER-derived vesicles attach to LCVs within the first 30 min of infection (12). Several hours later, attached ER-derived vesicles are less frequent, and LCVs at this time have ribosomes decorating the cytoplasmic surface of their membrane (12, 15). It is within this ER-derived organelle that *Legionella* begins to replicate (15), which is why this specialized...
compartment is called a replicative organelle. These data demonstrate that the Legionella Dot/Icm system is necessary for intercepting secretory vesicles. However, it is unknown how these ER-derived vesicles are recruited to a LCV and whether these vesicles participate directly in the remodeling of this compartment into a replicative organelle.

Herein, to better understand the cell biology of LCV transport, we examine host proteins that regulate the transport and fusion of ER-derived vesicles to see if they are important for biogenesis of the Legionella replicative organelle. Rab1 is a small guanosine triphosphatase (GTPase) that plays an important role in fusion of ER-derived vesicles with preGolgi intermediate compartments and the Golgi apparatus (16–18). Rab1 recruits factors necessary for the tethering and fusion of ER-derived vesicles with target membranes (16, 17). This membrane fusion process requires the pairing of soluble N-ethylmaleimide–sensitive factor attachment protein receptors (SNAREs) found on the vesicle membrane (v-SNARE) and target membrane (t-SNARE; references 19–21). The Sec22b protein in mammalian cells is found on ER-derived vesicles and functions as a v-SNARE (22–24). The proteins Membrin, Syntaxin 5, and Bet1 comprise the cognate t-SNARE complex to generate transport vesicles (28–32). The COPII subunits of the COPII coat and binds to this prebudding complex to generate transport vesicles (36–38). Rab1 fusion proteins with an amino terminal green fluorescent protein (GFP) tag were created in pEGFP-C1 (CLONTECH Laboratories, Inc.) using plasmids described previously that encode Rab1 (45). Plasmids encoding ARF1T31N-GFP and Sar1H79G were described previously (3). Plasmids encoding GFP-tagged VAMP4 were provided by N. Andrews (Yale University School of Medicine, New Haven, CT). Fluorescein-5 (Roche) was used to transfect CHO and CHO FcRII cells with the indicated plasmids according to the manufacturer’s instructions. BMMs were transfected with plasmids encoding Sec22b and Membrin using the CD34 cell Nucleofector™ kit (Amaxa Biosystems).

**Materials and Methods**

**Bacterial Strains and Tissue Culture.** Growth and macrophage infection by L. pneumophila strains derived from serogroup 1 strain LP01 have been described previously (6, 38, 39). Bone marrow–derived macrophages (BMMs) from the A/J mouse were prepared as described previously (40). Chinese hamster ovary (CHO) cells (41) and CHO FcγRII cells (42) were maintained in minimal Eagle’s media α (GIBCO BRL) supplemented with 10% FBS.

**Plasmids.** The cDNA encoding ARF1T31N (43) was ligated into pCLXSN (44). Plasmids producing myc-tagged versions of the mammalian Sec22b and Membrin were described previously (23, 24). Rab1 fusion proteins with an amino terminal green fluorescent protein (GFP) tag were created in pEGFP-C1 (CLONTECH Laboratories, Inc.) using plasmids described previously that encode Rab1 (45). Plasmids encoding ARF1T31N-GFP and Sar1H79G were described previously (3). Plasmid encoding GFP-tagged VAMP4 was provided by N. Andrews (Yale University School of Medicine, New Haven, CT). Fucose (6–10 µg/ml) was used to transflect CHO and CHO FcγRII cells with the indicated plasmids according to the manufacturer’s instructions. BMMs were transfected with plasmids encoding Sec22b and Membrin using the CD34 cell Nucleofector™ kit (Amaxa Biosystems).

**Inmunofluorescence Microscopy.** To examine LCV localization of the Rab1b, Rab2, and Rab6 proteins in BMMs, cells were infected with Legionella for 30 min as described previously (3). Where indicated, 10 µg/ml brefeldin A (Molecular Probes) was added to host cells 45 min before infection and remained in the medium during the course of the experiment. Cells were fixed in 1% paraformaldehyde in PBS for 20 min, permeabilized with ice-cold methanol, and blocked with 2% goat serum in PBS. Samples were stained with Rab-specific antibodies (Santa Cruz Biotechnology, Inc.) and fluorescein-labeled anti–rabbit IgG secondary antibodies. Bacteria and the host cell DNA were labeled using 0.1 µg/ml 4,6-diamidino-2-phenylindole. The same fixation and DNA staining procedure was used to localize GFP–Rab1a in transfected CHO FcγRII cells. To localize Sec22b and Membrin in CHO FcγRII cells, Legionella were opsonized with rabbit anti–Legionella antibody (1:1,000), and cells were infected for the times indicated. Cells were fixed in 1% paraformaldehyde in PBS for 20 min. Cells were permeabilized in 0.4% saponin for 15 min. Sec22b and Membrin were identified by anti–myc staining using a monoclonal antibody 9E10 (46) and a fluorescein-labeled anti–mouse secondary antibody. Bacteria were identified using a rhodamine-labeled anti–rabbit secondary antibody. Images were collected on an inverted microscope (model TE200; Nikon) equipped with a digital Hamamatsu ORCA ER camera. Images were exported, and color was added using Photoshop 7.0 (Adobe). When measuring the percentage of vacuoles that stained positive for Sec22b in CHO FcγRII cells producing dominant-interfering proteins, it was important to distinguish between intracellular and extracellular bacteria. This was done by differential labeling where the extracellular bacteria were first rhodamine-labeled with an anti–rabbit secondary antibody before permeabilization and 4,6-diamidino-2-phenylindole staining was used to identify all bacteria.

**Intracellular Growth Assays.** CHO cells were plated in 24-well dishes at a density of 3 × 10⁴ cells/well. After an overnight incubation, cells were cotransfected with plasmids encoding the FcγRII receptor (42) and the other indicated proteins. 16 h later, cells were infected with Legionella that were opsonized with rabbit polyclonal anti–Legionella antibodies (1:1,000 dilution). Plates were centrifuged at 150 g for 5 min and placed in a 37°C water bath for 5 min. Cells were washed three times with PBS, and fresh medium containing gentamicin (20 µg/ml⁻¹) was added for...
3 h to kill remaining extracellular bacteria. Cells were washed in PBS and either lysed to determine the efficiency of *Legionella* uptake or refreshed with 1 ml of antibiotic-free medium and lysed after an additional incubation period of 24 h. Bacterial numbers were determined in each lysate by measuring CFUs on charcoal yeast extract agar plates. Intracellular replication was measured as the fold increase in bacteria recovered after 24 h.

**Secretion Assays.** CHO FcγRII cells were plated in 24-well dishes at $3 \times 10^4$ cells/well. After an overnight incubation, the cells were cotransfected with plasmids encoding the proteins indicated and a plasmid encoding a secreted alkaline phosphatase (SEAP) protein. After a 24-h incubation period, cells were washed, and fresh tissue culture medium was added. SEAP activity was measured in triplicate wells 7 h later using the Phospha-light SEAP kit (Applied Biosystems). Data are presented as a secretion index, which is the ratio of SEAP activity detected in the culture medium to the cell-associated SEAP activity.

**Immunoelectron Microscopy.** CHO cells were plated in six-well dishes at a density of $2 \times 10^5$ cells/well. After an overnight incubation, cells were cotransfected with plasmids encoding FcγRII and Sec22b. Cells were infected 16 h after transfection with *Legionella* that were opsonized with rabbit polyclonal anti-*Legionella* antibodies (1:1,000 dilution). Plates were centrifuged at 150 g for 5 min and placed in a 37°C water bath for 10 min. Cells were washed three times with PBS and refreshed with prewarmed media for 1 h at 37°C. Cells were fixed in 4% paraformaldehyde (Electron Microscopy Sciences) in 0.25 M Hepes, pH 7.4, for 1 h at room temperature, followed by 8% paraformaldehyde in the same buffer overnight at 4°C. Cells were scraped, pelleted, and embedded in 10% bovine skin gelatin in PBS. Pieces of the pellet were infiltrated overnight with 2.3 M sucrose in PBS at 4°C, mounted on aluminum studs, and frozen in liquid nitrogen. Blocks were sectioned at $-108°C$ in an Ultracut cryo-ultramicrotome (Leica). 60-nm thick sections were collected using a 1:1 mixture of 2.3 M sucrose and 2% methyl cellulose and transferred onto formvar- and carbon-coated nickel grids. Sections were incubated for 10 min with 0.1 M NH$_4$Cl in PBS and for 20 min with 0.5% fish skin gelatin (FSG; Sigma-

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**Figure 1.** Specific recruitment of Rab1 to the LCV. (A–C) Rab-specific antibodies were used to stain BMMs infected for 30 min with wild-type *Legionella*. LCVs are indicated using arrows. Images show Rab1b (A) staining of the LCV. LCV staining was not observed using antibodies specific for Rab2 (B) or Rab6 (C). (D) BMMs were fixed at the indicated times after infection by wild-type *Legionella* and Rab1b staining of the LCV was determined using fluorescence microscopy. Values represent the mean ± SE of three independent experiments in which 50 vacuoles were scored for each time point. Bars, 5 μm.
Aldrich) in PBS (PBS-FSG) and incubated for 30 min at room temperature with rabbit anti-myc polyclonal antibody (Santa Cruz Biotechnology, Inc.) diluted in PBS-FSG. After four washes in PBS (5 min total), the sections were labeled with 10 nm protein A–gold conjugate (Utrecht University, Netherlands) diluted in PBS-FSG for 30 min. Sections were washed again in PBS and fixed with 1% glutaraldehyde in PBS. The sections were rinsed in distilled water and incubated with 1.8% methyl cellulose (25 centipoises; Sigma-Aldrich) and 0.5% uranyl acetate for 10 min on ice. The sections were dried and examined in a Philips Tecnai 12-electron microscope.

Online Supplemental Material. An unbiased assay to measure the effect of dominant-interfering proteins on \textit{Legionella} intracellular growth is provided in Fig. S1. Online supplemental material is available at http://www.jem.org/cgi/content/full/jem.20031706/DC1.

Results

Specific Recruitment of \textit{Rab}1 to the LCV. Because \textit{Rab} proteins are key regulators of most vesicle transport and fusion processes, it is predicted that a subset of \textit{Rab} proteins should be involved in biogenesis of the \textit{Legionella} replicative organelle. Given that the LCV is remodeled into an ER–derived organelle, we asked whether \textit{Rab} proteins that function in the transport of secretory vesicles between the ER and Golgi apparatus are recruited to the LCV. Immunolocalization studies demonstrated that the LCV acquires \textit{Rab}1 within the first 30 min of infection (Fig. 1, A and D). There was no detectable staining of LCVs using antibodies specific for either \textit{Rab}2 or \textit{Rab}6 (Fig. 1, B and C), which are two other \textit{Rab} proteins that function in vesicular transport between the ER and Golgi apparatus (47–51). \textit{Rab}1 staining was not observed on LCVs harboring a \textit{dotA} mutant of \textit{Legionella} (Fig. 2 A and Fig. 3), indicating that \textit{Rab}1 recruitment to the LCV requires factors delivered into host cells by the Dot/Icm system. \textit{Rab}1 recruitment to LCVs was also observed in \textit{CHO}-FcγRII cells (Figs. 2 D and 3), a stable cell line that produces the FcγRII protein. \textit{Rab}1 staining in \textit{CHO}-FcγRII cells is consistent with previous work showing that the LCV transport pathway in these cells mirrors the pathway defined in BMMs (3).

The majority of LCVs stained positive for \textit{Rab}1 at 30 min after infection (Fig. 1 D). \textit{Rab}1 staining diminished

![Figure 2](image-url)

Figure 2. Recruitment of \textit{Rab}1 to the LCV requires the Dot/Icm system and is independent of vesicular transport from the ER. (A and B) BMMs were fixed 30 min after infection with a \textit{Legionella} \textit{dotA} mutant (A) or \textit{ralF} mutant (B) and stained with a \textit{Rab}1b-specific antibody. Arrows indicate the location of the LCV that is magnified in the inset of each panel. (C) \textit{Rab}1b staining is detected on an LCV containing wild-type \textit{Legionella} in BMMs treated with brefeldin A. Notice that the expected Golgi fragmentation resulting from brefeldin A treatment has eliminated the intense \textit{Rab}1b-specific signal observed in the untreated cells in A and B. (D) GFP-\textit{Rab}1a localizes to an LCV containing wild-type \textit{Legionella} in \textit{CHO} FcγRII cells. Bars, 5 μm.
over time with fewer LCVs staining positive for Rab1 at 4 h after infection (Fig. 1 D). The kinetics of Rab1 staining of the LCV were more rapid than that observed previously for ARF1 (3) and correlated with a stage of vacuole maturation in which LCVs have intercepted ER-derived vesicles, but have not yet taken on the appearance of an ER-like organelle (3, 12, 15). The Legionella protein RaI, which plays an important role in the recruitment of ARF to the LCV by functioning as a guanine nucleotide exchange factor for ARF (6), was not required for Rab1 recruitment to the LCV (Figs. 2 B and 3). Treatment of BMMs with brefeldin A, which blocks delivery of early secretory markers to the LCV by disrupting vesicular transport from ER exit sites, did not affect Rab1 recruitment to the LCV (Figs. 2 C and 3). These data indicate that recruitment of Rab1 to the LCV is a Dot/Icm-mediated process that does not require remodeling of the LCV by early secretory vesicles.

Inhibition of Rab1 Function Interferes with Intracellular Replication of Legionella. To determine whether Rab1 function is important for intracellular replication of Legionella, a plasmid encoding the GDP-locked dominant-interfering protein Rab1S25N was cotransfected into CHO cells along with a plasmid encoding the FcγRII protein. Because CHO cells are normally nonphagocytic, the uptake of Legionella is inefficient. However, cotransfection of a plasmid encoding FcγRII allowed uptake of IgG-opsonized Legionella preferentially into cells that were transiently transfected (Supplemental Results, available at http://www.jem.org/cgi/content/full/jem.20031706/DC1). Using this approach, we consistently found >90% of the internalized bacteria residing in transfected cells; even under intracellular growth of Legionella in untransfected cells over a 24-h period, the total number of bacteria in these untransfected cells did not equal the starting number of bacteria in the transfected cells (Fig. S1, available at http://www.jem.org/cgi/content/full/jem.20031706/DC1). Using this assay, we have confirmed previous studies based on single cell analysis to show that the interfering variants ARF1T31N and Sar1H79G both disrupt intracellular replication of Legionella (Fig. 4 A). Intracellular growth of Legionella in cells producing Rab1S25N was significantly reduced compared with control cells producing GFP alone (Fig. 4 A). These data indicate that Rab1 function is important for conversion of the LCV into an organelle that supports intracellular replication of Legionella.

The degree of Legionella growth inhibition observed using Rab1S25N was not as pronounced as that seen using ARF1T31N or Sar1H79G. To determine how effective these proteins were at disrupting secretory traffic, plasmids encoding SEAP were cotransfected into CHO cells along with plasmids encoding the proteins being used to disrupt the secretory pathway. These data show that ARF1T31N, Sar1H79G, and Rab1S25N all disrupt protein secretion (Fig. 4 B). The degree to which interfering variants affect protein secretion correlates with the severity of defects observed for intracellular replication of Legionella.

Rab1 Function Is Important for the Delivery of the Sec22b Protein to the LCV. The primary role of Rab1 is to facilitate the transport and docking of ER-derived vesicles with the cis-Golgi apparatus. Docking of these vesicles promotes SNARE-mediated vesicle fusion. To determine whether Rab1 is playing a similar role on the LCV, we examined whether SNARE proteins that mediate fusion of ER-derived vesicles with target membranes are delivered to the LCV. Toward this end, we examined the localization of both the v-SNARE protein Sec22b and the t-SNARE protein Membrin. Distinct colocalization of Sec22b was observed on the LCV in CHO-FcγRII cells (Fig. 5 A). Vacuoles containing dotA mutant Legionella did not acquire Sec22b (Fig. 5 B). Interestingly, Membrin colocalization was not detected on LCVs (Fig. 5 C). Similar results were obtained in transfected BMMs (unpublished data). Sec22b colocalization was inhibited by brefeldin A (Fig. 5 D).
Consistent with results obtained with brefeldin A, inhibiting the production of vesicles from ER exit sites using Sar1H79G or ARF1T31N also had a potent effect on delaying the delivery of Sec22b to the LCV (Fig. 5 D). In contrast with what was observed for Rab1, Sec22b recruitment to the LCV requires vesicular transport from the ER. These data indicate that Rab1 binding to the LCV precedes the delivery of Sec22b.

Because Rab1 coordinates the tethering and fusion of ER-derived vesicles with target membranes, next we asked whether inhibiting Rab1 function had any effect on the kinetics of Sec22b localization to the LCV. Ectopic production of the GDP-locked Rab1S25N protein potently inhibited Sec22b localization to LCVs examined 30 min after infection (Fig. 5 D). Within 2 h, most LCVs had acquired Sec22b in cells producing Rab1S25N, which indicates that there is a kinetic defect in Sec22b transport to the LCV but not an absolute block, consistent with Rab1 being involved in vesicle tethering rather than vesicle production. These data suggest that Rab1 recruitment to the LCV promotes the transport and fusion of ER-derived vesicles containing Sec22b.

**The Sec22b Protein Is Delivered from ER-derived Vesicles to the Vacuole Membrane Surrounding Legionella.** To determine whether ER-derived vesicles containing Sec22b fuse with the LCV, immunogold labeling studies were conducted. Electron micrographs clearly show Sec22b-specific gold particles labeling both vesicles attached to the LCV (Fig. 6, A and B, arrowheads) and on the vacuole mem-
brane surrounding Legionella (Fig. 6, A and B, arrows). Sec22b-specific gold particles were not detected on the vacuole membrane surrounding dotA mutant Legionella (Fig. 6, C and D), even though gold particles were in abundance on ER membranes in close proximity to these vacuoles (Fig. 6, C and D, asterisks). These data indicate that Sec22b is transferred from the attached vesicles to the limiting membrane of the vacuole that surrounds Legionella. Because Sec22b is a transmembrane protein, the presence of Sec22b on the LCV membrane indicates that attached ER-derived vesicles have undergone fusion with the LCV.

**Sec22b Function Is Important for Establishment of the Legionella Replicative Organelle.** Overproduction of an individual SNARE protein can have a dominant negative effect on specific vesicular transport processes by affecting the proper distribution and functioning of cognate protein binding partners, including other SNARE proteins (24, 52). Previous studies have shown that overproduction of Membrin, and to a lesser extent overproduction of Sec22b, can interfere with the transport of early secretory vesicles (24). We found that overproduction of Membrin interfered with the intracellular growth of Legionella; however, overproduction of Sec22b did not affect Legionella replication (Fig. 7 A). Overproduction of the trans-Golgi SNARE protein VAMP4 did not affect intracellular growth of Legionella (Fig. 7 A). Defects in secretion of an alkaline phosphatase reporter were observed in cells overproducing either Membrin or VAMP4, but not in cells overproducing Sec22b (Fig. 7 B). These data suggest that overproduction of Membrin might specifically interfere with intracellular growth of Legionella by titration of a host factor that is important for replicative organelle biogenesis.
To test whether this effect might be due to titration of Sec22b, we overexpressed both Membrin and Sec22b in cells by cotransfection and assayed intracellular replication of *Legionella*. These data show that ectopic production of Sec22b can suppress *Legionella* growth restriction mediated by Membrin overproduction (Fig. 7 A). VAMP4 overproduction did not suppress the *Legionella* growth restriction caused by ectopic production of Membrin. Importantly, coproduction of Sec22b did not suppress the defect in secretion of alkaline phosphatase caused by Membrin overproduction. These data strongly suggest that Membrin overproduction restricts the growth of *Legionella* by titration of Sec22b as opposed to the more general effect Membrin overproduction has on protein secretion. Consistent with the hypothesis that Membrin overproduction titrates available Sec22b, we found that increasing the amounts of transfected plasmid DNA encoding Membrin resulted in a decrease in the percent of LCVs that stained positive for Sec22b (Fig. 7 C). These data indicate that Membrin overproduction prevents the delivery of Sec22b to the LCV. Thus, both morphological and functional data indicate that Sec22b plays an important role in the creation of the *Legionella* replicative organelle.

**Discussion**

The goal of this paper was to define transport pathways and host factors involved in the creation of the *Legionella* replicative organelle. Toward this end, first we examined the localization of Rab protein family members that are involved in vesicle transport between the ER and Golgi apparatus. These studies revealed that Rab1 is recruited to vacuoles containing wild-type *Legionella* and is not found on vacuoles containing *dotA* mutants (Fig. 1). The proteins Rab2 and Rab6, which (like Rab1) are involved in vesicular transport between the ER and Golgi apparatus, were not found on LCVs. These data indicate that Rab1 recruitment to the LCV is specific and controlled by the Dot/Icm system.

Rab1 recruitment to the LCV was not inhibited by brefeldin A (Fig. 2). This finding is significant because brefeldin A has been shown to block the recruitment of other early secretory proteins that display LCV colocalization (3). Because brefeldin A inhibits production and transport of ER-derived vesicles, these data demonstrate that Rab1 recruitment to the LCV is not coincident with remodeling of this compartment by ER-derived vesicles, suggesting a role for Rab1 on the LCV before the binding of these vesicles. Interference of Rab1 function by the Rab1S25N protein inhibited intracellular replication of *Legionella* (Fig. 4). These data indicate that Rab1 is playing an important role in biogenesis of the organelle in which *Legionella* replicates.

Because Rab1 is involved in the recruitment of factors required for the tethering and fusion of ER-derived vesicles with target membranes (16, 17), we examined whether the SNARE proteins involved in this fusion reaction were recruited to the LCV. Our data show that the Sec22b protein was recruited to LCVs (Figs. 5 and 6). Sec22b recruitment to the LCV was blocked by inhibiting the production of ER-derived vesicles with either Sar1H79G, ARF1T31N, or brefeldin A, demonstrating that Sec22b is transported to the LCV in secretory vesicles. The finding that Sec22b recruitment to the LCV is delayed in cells producing Rab1S25N indicates that Rab1 function is important for delivering cargo in ER-derived vesicles to the LCV. These data indicate that the principle role for Rab1 on the LCV is to promote transport and fusion of ER-derived vesicles with this organelle.

Immunogold labeling was used to determine whether ER-derived vesicles fuse with the LCV. These studies show that the Sec22b protein is present both on the membrane of vesicles attached to the LCV and also on the membrane surrounding *Legionella* (Fig. 6). Because Sec22b is a transmembrane protein that initially inserts in the ER membrane and exits the ER in early secretory vesicles, detection of the Sec22b protein on the LCV membrane provides evidence that the ER-derived vesicles attached to the LCV are fusing with this organelle. The absence of Sec22b from the membrane of vacuoles containing *dotA* mutants of *Legionella* is consistent with previous data showing that the transport and attachment of ER-derived vesicles to the LCV requires the Dot/Icm system (3, 12–14). Although this is the first direct evidence that ER-derived vesicles fuse with the LCV, these data are consistent with previous electron microscopy data, which had suggested that fusion of attached vesicles could account for a thinning of the LCV membrane that was observed shortly after bacterial uptake (12).

Given that cognate SNAREs form cis–SNARE complexes that colocalize in the cell, it was surprising to find that the Sec22b protein was enriched on the LCV, but that the cognate SNARE protein Membrin was undetectable on these vacuoles (Fig. 5). Recent studies examining COPII-directed recruitment of *Saccharomyces cerevisiae* homologues of these mammalian SNAREs might provide an explanation for the specific recruitment of Sec22b to the LCV (29, 53, 54). Yeast Sec22p can exit the ER in either a monomeric form or in a cis complex with other SNARE proteins. The recruitment of Sec22p is directed by the Sec23/24p subcomplex, which can bind to a motif in Bet1p and recruit cis–SNARE complexes that contain Sec22p into ER-derived vesicles. Additionally, there is a spatially distinct binding site on the Sec23/24p subcomplex that can recruit a monomeric form of Sec22p into ER-derived vesicles. Because the Sec22p binding site in the Sec23/24p complex is likely to be conserved between yeast and mammalian homologues, the packaging of Sec22b into ER-derived vesicles should be regulated similarly. Given the distinct binding site for Sec22b in COPII, it is possible that the LCV preferentially recruits COPII-generated vesicles enriched for the monomeric form of Sec22b, which would explain the absence of cognate SNARE proteins that bind to Sec22b.

It has been shown that dominant negative effects on membrane transport sometimes result from the overpro-
duction of a single SNARE protein (24, 52), presumably by affecting the proper subcellular distribution of cognate proteins to which that SNARE protein binds, including other SNAREs. Overproduction of the SNARE protein Membrin was found to inhibit the intracellular growth of *Legionella* by a process that could be suppressed by overproduction of Sec22b (Fig. 7). Although secretion of alkaline phosphatase from cells was inhibited upon overproduction of the SNARE proteins Membrin or VAMP4, only Membrin overproduction interfered with *Legionella* replication. Additionally, overproduction of Sec22b did not suppress the secretory defect caused by overproduction of Membrin, indicating that inhibition of *Legionella* replication was not simply a consequence of a general perturbation in the secretory pathway caused by overproduction of a SNARE protein. These data are consistent with monomeric Sec22b being an important SNARE protein that is used for biogenesis of the *Legionella* replicative organelle. Accordingly, overproduction of Membrin interferes with replicative organelle biogenesis because it effectively titrates monomeric Sec22b. Overproduction of Sec22b can suppress the inhibition of *Legionella* intracellular growth caused by Membrin overproduction by restoring cellular pools of monomeric Sec22b but does not restore secretion defects caused by the titration of the other proteins to which Membrin binds. Direct evidence in support of this hypothesis comes from data showing that increasing the amount of Membrin produced in host cells interferes with the delivery of Sec22b to the LCV. These data provide the first functional evidence that host ER–Golgi SNARE proteins play an important role in formation of the *Legionella* replicative organelle.

Based on these data, we hypothesize that LCVs preferentially recruit vesicles created by COPII that contain the monomeric form of the Sec22b protein. We demonstrate that Rab1 plays an important role in the recruitment of these Sec22b-containing vesicles to the LCV. Because the LCV is derived from the plasma membrane, the t-SNARE complex composed of Membrin, Bet1, and Syntaxin5 would not be present on this compartment. To initiate fusion with ER-derived vesicles, it is possible that *Legionella* proteins injected into the vacuole membrane by the Dot/Icm system could serve as a functionally equivalent t-SNARE complex for a Sec22b-dependent fusion reaction. Alternatively, a t-SNARE complex present at the plasma membrane might pair with Sec22b and promote the fusion of ER-derived organelles with the LCV. In this second model, the main role for *Legionella* proteins would be to recruit proteins such as Rab1 that can facilitate vesicle tethering and facilitate membrane fusion. Interestingly, previous in vitro studies using purified components have shown that the *S. cerevisiae* Sec22 protein can promote membrane fusion with a plasma membrane t-SNARE complex consisting of the proteins Sso1 and Sec9p (55). Although it has not been shown that the mammalian Sec22b protein can function as a v-SNARE for fusion with a plasma membrane t-SNARE, recent studies that implicate direct fusion between the ER and phagosomes suggest that such interactions are possible (56). Thus, it is likely that determining the mechanism of membrane fusion between ER-derived vesicles and the LCV will reveal new insights into the cell biology of phagosome maturation and vesicular transport in the early secretory pathway.

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References


