Proteomic analysis of the Simkania-containing vacuole: the central role of retrograde transport

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Summary

Simkania negevensis is an obligate intracellular bacterial pathogen that grows in amoeba or human cells within a membrane-bound vacuole forming endoplasmic reticulum (ER) contact sites. The membrane of this Simkania-containing vacuole (SnCV) is a critical host–pathogen interface whose origin and molecular interactions with cellular organelles remain poorly defined. We performed proteomic analysis of purified ER-SnCV-membranes using label free LC-MS² to define the pathogen-containing organelle composition. Of the 1,178 proteins of human and 302 proteins of Simkania origin identified by this strategy, 51 host cell proteins were enriched or depleted by infection and 57 proteins were associated with host endosomal transport pathways. Chemical inhibitors that selectively interfere with trafficking at the early endosome-to-trans-Golgi network (TGN) interface (retrograde transport) affected SnCV formation, morphology and lipid transport. Our data demonstrate that Simkania exploits early endosome-to-TGN transport for nutrient acquisition and growth.

Introduction

The obligate intracellular, Gram-negative bacterium Simkania negevensis (Sn) belongs to the family Simkaniaceae in the order Chlamydiales (Everett et al., 1999; Kahane et al., 2007). The genome of Sn is approximately 2.5 Mbp in size and thus two to three times larger than the genome of Chlamydia (Collinge et al., 2011). Chlamydiae are found as symbionts and pathogens in a wide range of eukaryotes, including protists, invertebrates and vertebrates (Subtil et al., 2014). Sn is able to replicate in amoebae, human and simian epithelial cells and macrophages (Kahane et al., 1993; 2007) and has been associated with infections of the upper respiratory tract in infants and adults (Lieberman et al., 1997; Kahane et al., 1999; Horn, 2008; Lamothe and Greub, 2010). Infections with the two closely related human pathogenic bacteria Chlamydia pneumoniae and Chlamydia psittaci can cause community acquired or animal transmitted pneumonia, chronic bronchitis and chronic asthma (Hughes et al., 1997; Hahn and McDonald, 1998; Harkinezhad et al., 2009).

Similar to Chlamydia, Sn undergoes a unique developmental cycle during intracellular replication. Infection is initiated by entry of an electron dense elementary body (EB) into the host cell. Inside of a membrane-bound compartment, termed inclusion, EBs differentiate into reticulate bodies (RBs) within a short period of time. RBs are the metabolically active and replicative form of Chlamydia and divide by binary fission. At the end of every multiplication phase, RBs re-differentiate into EBs to start a new infection cycle (Dautry-Varsat et al., 2005; Hybiske and Stephens, 2007). Sn has a longer life cycle than other chlamydial species with a replication plateau at day 3 post infection and the release of EBs by host cell lysis after 5–12 days post infection (Kahane et al., 1993; 1999; 2002). Long-term growth is likely connected to inhibition of host cell death induced by Simkania infection (Karunakaran et al., 2011).

The biogenesis of the chlamydial inclusion and interaction with the host cell have been intensively investigated (for a review, see Fields and Hackstadt, 2002). Once inside the cell, Chlamydia resides within a membrane-surrounded host-derived vacuole within the cytoplasm (Hackstadt et al., 1997). Shortly after, the vacuole is transported to the microtubule organising centre in a dynein...
dependent manner likely through interaction with the dynactin subunit p150(Glued) (Clausen et al., 1997; Hackstadt, 2000; Grieshaber et al., 2003). Individual inclusions from C. trachomatis fuse to form a single large vacuole, whereas many other Chlamydioides develop multiple small inclusions (Clausen et al., 1997). Chlamydia directly modifies the inclusion by secretion of inclusion membrane proteins (Inc) that also extend to the cytosolic face of the inclusion membrane (Rockey et al., 1997; Hackstadt et al., 1999; Scidmore and Hackstadt, 2001). The action of bacterial proteins in the chlamydial inclusion membrane is assumed to, e.g. prevent endolysosomal fusion, avoid host immune responses and support fusion with sphingomyelin and cholesterol-containing exocytic vesicles from the Golgi apparatus (Scidmore et al., 1996b; Ocju, 1997; Hackstadt et al., 1999; Carabeo et al., 2003). It is currently unknown how well conserved basic molecular mechanisms behind these processes are within the order Chlamydioidae.

Sn forms a continuous membrane system spanning the host cell, and this is clearly distinct from the single vacuolar structure of Chlamydia (Mehlitz et al., 2014). Interestingly, Sn shapes extensive endoplasmic reticulum (ER) contact sites, which are reminiscent of inclusion-ER contact sites also termed pathogen synapses (Derre et al., 2011; Dumoux et al., 2012; Mehlitz et al., 2014). The recent discovery of Inc proteins within other members of the Chlamydioidae, like Parachlamydia, Waddlia and Simkania, and their low degree of conservation makes it highly likely that the interaction with the host is much more diverse than previously anticipated (Fields and Hackstadt, 2002; Heinz et al., 2010; Collingro et al., 2011). Only three putative Inc proteins are conserved among Chlamydioidae, and just a few members can enter and multiply in mammalian cells, indicating a dependency on the host cell type or different molecular mechanisms. The identification and characterisation of Inc or Inc-like proteins in the Sn vacuole could reveal the unique molecular interaction with the host ER.

The ER forms a highly branched membrane network arranged in tubes, sheets and cisternae enclosing a fluid-filled interior. Parts of the ER membrane directly associate with the nuclear envelope and extend into the cytosol spanning to the plasma membrane (PM) (English et al., 2009). The ER can be divided into smooth and rough ER (SER and RER). RER can easily be identified through lining ribosomes and is associated with multiple cellular functions including protein synthesis, folding or quality control as well as Ca²⁺ homeostasis and regulation of apoptosis (Brekenridge et al., 2003; Rizzuto et al., 2009; Braakman and Bulleid, 2011). The SER is responsible for vesicle formation, budding and fusion processes, is free of ribosomes and forms a highly complex tubular structure (Shibata et al., 2006; Bravo et al., 2013). In eukaryotic cells, the ER interacts with the Golgi apparatus through bidirectional vesicle trafficking, co-ordinating cellular protein and lipid transport (Ghaemmaghami et al., 2003).

Proteins on donor organelles, e.g. vSNAREs (vesicle soluble N-ethylmaleimide sensitive factor attachment receptor) guide the fusion with the target organelle via interaction with tSNAREs (target SNAPes) (Bonifacino and Glick, 2004). An example of this mechanism is provided by COPI and II vesicles, where COPII vesicles are transported from the ER to the Golgi (anterograde transport) and COPII vesicles are involved in the retrograde backhaul (Brandizzi and Barlowe, 2013). Also non-vesicular mechanisms have been reported for the transfer of proteins and lipids between organelles. Arf1/COPII complexes and the ER to Golgi ceramide transfer protein CERT mediate such non-vesicular transport (Hanada et al., 2003; Hanada, 2010; Wilfling et al., 2014). This type of transport bridges short distances, e.g. between ER and Golgi membranes, and is required for chlamydial inclusion development (Derre et al., 2011; Elwell et al., 2011; Dumoux et al., 2012). In contrast, data for evolutionary related organisms like Simkania are missing. Simkania forms numerous ER-vacuole contact sites (Mehlitz et al., 2014) and might thereby facilitate uptake of metabolites or lipids required for growth.

Chlamydia requires host-derived lipids like sphingomyelin (SM) and cholesterol for development (Hackstadt et al., 1995; Scidmore et al., 1996a,b; Carabeo et al., 2003). The inclusion membrane thereby constitutes a critical intracellular barrier as these lipids have to be transported into the inclusion to be accessible for the bacteria. As the inclusion membrane is impermeable to small molecules (Heinzen and Hackstadt, 1997), it is thought that Chlamydia acquires lipids from vesicles from the secretory pathway that fuse with the inclusion membrane (Carabeo et al., 2003; Heuer et al., 2009; Derre et al., 2011; Elwell et al., 2011). This includes the interception of SM-containing Golgi-derived exocytic vesicles destined for the PM or fusion with multivesicular body-derived vesicles (MBV) (Hackstadt et al., 1995; Scidmore et al., 1996b; Beatty, 2006; 2008; Robertson et al., 2009). The latter could also play a role in delivering nutrient into the inclusion that are then taken up by Chlamydia via bacterial transporters, permeases and translocases based on active transport processes (Saka et al., 2011). Golgi fragmentation induced by C. trachomatis has been shown to be important for SM acquisition during the later stages of infection and may be required for subsequent fusion with exocytic vesicles by a mechanism involving Rab6 and Rab11 (Heuer et al., 2009; Rejman Lipinski et al., 2009).

Here, we established an ER-Simkania negevensis containing vacuole (SnCV) isolation protocol to define the...
composition of the organelle by label free LC-MS². Our data suggest the depletion of components of endosomal trafficking, ER to Golgi as well as exocytic transport in the SnCV. Interestingly, proteins associated with recycling endosomal as well as Golgi to ER transport were enriched in the SnCV, suggesting a role of these pathways in nutrient uptake by *Simkania*. In line with this, interfering with retrograde transport affected *Simkania* replication and SnCV formation. SnCV formation and ceramide acquisition were severely affected by blocking retrograde trafficking underlying the importance of this route for intracellular adaptation of *Simkania*.

**Results**

**Purification of ER-SnCV membranes**

*Simkania negevensis* containing vacuoles are intimately associated with the ER of their host cell (Mehlitz *et al.*, 2014). It is unknown how the ER-SnCV interaction is established and whether it affects the composition of the ER. To investigate the effect of infection on the proteome of the ER membranes, the organelle had to be isolated from control and infected cells (Huber *et al.*, 2003). We therefore established the isolation of crude ER- and ER-SnCV-membranes from control and *S. negevensis* infected cells (see Fig. 1A and Experimental procedures for more details). Individual isolation steps were analysed by light-microscopy to confirm cell swelling and mechanical rupture (Fig. 1B). Sequential centrifugation steps led to the fractionation into pre-nuclear (PNF), pre-mitochondrial (PMF) and crude microsomal ER-fraction (CMF) (Fig. 1A). The quality of the ER-SnCV membrane purification was controlled by immunoblotting by detecting marker proteins for cytoskeleton (ß-Actin), mitochondria (Sam50) and nuclei (Lamin B1) (Fig. 1C). Calnexin an ER membrane protein was detected within PMF and CMF fractions indicating the strong association of the ER with organelles like mitochondria (Fig. 1C). A subset of soluble ER proteins like PDI and ERP72 were found in the PMF and only to a low extend in the CMF, which could be due to association with other organelles (Fig. 1C). KDEL (Lys-Asp-Glu-Leu) ER retention signal containing proteins were selectively enriched in the CMF (Fig. 1C). Bacteria were strongly depleted from the CMF as was monitored by detecting the bacterial heat shock protein GroEL (Fig. 1C). Residual GroEL was most likely derived from broken bacteria, as intact bacteria were hardly detectable in the CMF by transmission electron microscopy (TEM) (Fig. 1D). Interestingly, we observed only a fraction of total bacterial proteins within the CMF fraction likely representing bacterial proteins localised to the SnCV membrane (Fig. 1C). These results show the isolation of crude ER-SnCV membranes from *Simkania* infected cells as well as the modulation of the ER composition through *Simkania* infection.

**Proteome analysis of crude ER-SnCV membranes (CMF)**

We then analysed the crude ER-SnCV membrane fraction (CMF) of three biological replicates by LC-MS² to identify *Simkania* infection-mediated changes of ER composition and identified 1178 human and 302 *Simkania* proteins in the infected samples (Fig. 2A and Supplementary Tables S1 and S3). One hundred twenty-three out of 302 (41%) identified bacterial proteins were hypothetical proteins with no known function or homology upon database searches respectively (Supplementary Table S3). Inc proteins are highly species-specific proteins and can be predicted applying the following criteria: (i) hypothetical protein, (ii) no Sec-signal and (iii) bi-lobed hydrophobicity profile (Collingro *et al.*, 2011). Twenty-three out of 123 hypothetical proteins (19%) fulfilled these criteria supporting the co-purification of ER and SnCV (Supplementary Table S3). We next grouped the remaining 179 proteins according to known function or subcellular localisation: 16% of the proteins were ribosomal components, 11% were localised in the bacterial cytoplasm and 9% were membrane proteins or belonged to RNA-polymerase, proton transport, cell surface, helicase or acetyl-CoA carboxylase complex (Supplementary Table S3). More interestingly, 20% of the proteins were associated with type III or Sec-system according to Uniprot and Effectors databases (Supplementary Table S3). We did not detect any components of the *Simkania* type IV secretion system. Comparison of identified proteins with published pan-genome data (Collingro *et al.*, 2011) showed that 11 out of the 302 identified proteins are *Simkania* specific factors (11/11 hypothetical, 4/11 located on the plasmid pSN), 16/302 proteins showed homologs in *Chlamydia* and 12/302 proteins are virulence associated, e.g. LPS synthesis proteins or macrophage infectivity potentiator (Supplementary Tables S4 and S5).

The potential subcellular localisation of the identified human proteins was predicted by database searches using WEB-based Gene SeT AnaLysis (WEB-Gestalt) toolkit (Wang *et al.*, 2013) (Fig. 2B). Most of these proteins were predicted by GO annotation as part of macromolecular complexes, membrane enclosed lumen and membrane of the nucleus, cytosol, mitochondria, cytoskeleton, endomembrane system and ER (Fig. 2B). We then performed an enrichment analysis using WEB-Gestalt and GO analysis against the human genome as a reference set and standard settings with the exception of top10 significance level and minimum of 10 genes per category. Enrichment analysis confirmed isolation of ER-targeted proteins in comparison with the total genome.
A. Cell culture, 12 x 6 well plates, HeLa, 80% confluent
   Collect and wash, 5 min centrifugation, 4°C, 600 x g
   Swelling in 0.5 x hypotonic buffer 20 min, 4°C, 5 min centrifugation, 4°C, 600 x g
   Resuspend in 0.5 x isotonic buffer
   Cell-lysis, 10 min ultrasonic bath, 4°C, Dounce Homogenizor 10 strokes, 200 rpm, 4°C

Sequential centrifugation
PNF, 15 min, 4°C, 1,000 x g
PMF, 15 min, 4°C, 12,000 x g
CMF, 60 min, 4°C, 100,000 x g

B. Swelling
C. PMF
D. CMF

Fig. 1. Purification of Simkania-containing vacuole (SnCV) and ER membranes.
A. Flowchart showing purification of ER-SnCV-membranes. ER-SnCV-membranes were purified from infected HeLa cells by successive swelling and lysis steps followed by three sequential centrifugation steps.
B. Phase contrast microscopy of swelling and lysis steps described in A. Cells were equally swollen and visibly broken by mechanical lyses.
C. Immunoblot analysis of purified membrane fractions. Composition of the purified ER-SnCV-membranes is shown by various marker proteins. Purification led to isolation of mainly ER membrane associated (Calnexin and KDEL) and bacterial proteins (anti-Sn) in the final fraction (CMF). Soluble ER proteins (Erp72 and PDI) and markers for other cell compartments like mitochondria (Sam50), nuclei (Lamin B1) and cytoskeleton (ß-Actin) are excluded from the CMF.
D. Transmission electron microscopy of infected PMF (left) and CMF (right) fractions. While the PMF still contained both intact Simkania and mitochondria, the CMF is devoid of intact bacteria, indicating that purified bacterial proteins might be membrane-associated secreted proteins. Min = minutes, PNF = pre-nuclear fraction, PMF = pre-mitochondrial fraction, CMF = crude microsomal fraction.
validating the crude ER-SnCV fraction (CMF) used for the analysis (Supplementary Fig. S1). Most of the other significantly enriched categories were found to be involved in, e.g., RNA processing and translation (Supplementary Fig. S1).

The proteome data were further classified by function, like, e.g., protein-, nucleic acid-, ion and metabolic binding and various enzymatic functions (Fig. 2C). Categorisation by cellular process indicated that the identified proteins are involved mainly in metabolic processes but also cellular component organisation and responses to various stimuli (Fig. 2D). We next tested for differential expression in spectral counts between infected and non-infected samples after adjusting for sample effects using the edgeR statistical package. A large number of proteins with a nominal P-value < 0.05 could be identified (Supplementary Table S2). Comparing the expression of infection-associated genes across all three samples indicated that infection in sample 3 did not work (Supplementary Fig. S2). We focused our further analysis on samples 1 and 2, which show similar infection-induced regulatory patterns (Fig. 3A).

Six proteins (2’-5’-oligoadenylate synthase 3, Heterogeneous nuclear ribonucleoproteins A2/B1, Voltage-dependent anion-selective channel protein 1, ATP synthase subunit beta, Heterogeneous nuclear ribonucleoprotein A0 and ATP synthase subunit alpha) were differentially regulated after adjusting P-values for multiple testing (Supplementary Table S2). To allow for explorative network analysis, we lowered the threshold of adjusted P-value to < 0.2 now taking into account 51 differentially expressed genes (Supplementary Tables S2 and S6). Ingenuity analysis categorised these proteins into 21 significant categories, with the top canonical pathways being involved in mitochondrial dysfunction, oxidative phosphorylation, RNA charging, telomere extension by telomerase and acetyl-CoA biosynthesis III (Fig. 3B). Network connectivity analysis of these 51 proteins resulted in three major networks (Fig. 3C–E). We observed a large number of mitochondrial proteins (F1 ATPase, ATP5A1, ATP5B, ATP5O, ATP5F1, ATP synthase, VDAC1, TOMM40, UOCRC2) depleted from the ER/SnCV fraction (Fig. 3C). Similarly, anterograde transport components (Rab1A, Rab14, Sec22B) appeared to be depleted in infected cells (Fig. 3C).

Cytoskeletal components and regulators (ARHGEF2, MSN) especially cytokeratins appear to be de-regulated (enriched: KRT6A, Cytokeratin, KRT5, de-regulated: KRT8, KRT18) while myosins are depleted (MYH10, Myosin 2) in ER/SnCV from infected cells (Fig. 3C–E). Taken together, these results provide an overview of the proteins regulated in the ER and associated organelles of Simkania-infected cells.

Some differentially regulated proteins obtained by LC-MS² and statistical testing were validated by immunoblotting (Fig. 3F and G). Interestingly, we observed an enrichment of CCT2 (cytosolic T-complex 1 subunit beta, TCP-1-B), which forms a complex with CCT1 (cytosolic T-complex 1 subunit alpha, TCP-1-A) in ER/SnCV fractions (Fig. 3G). ARHGEF2 (Rho guanine nucleotide exchange factor 2, GEF H1) was also found to be enriched in purified membranes from infected cells (Fig. 3G) confirming our mass spectrometry approach (Fig. 3A, C–E, Supplementary Tables S1 and S2). Enrichment of ARHGEF2 and CCT2 at day 3 post infection was not apparent in whole cell lysates of time course experiments, validating that enrichment/depletion analysis is not merely due to overall expression changes within the cell. In contrast, we also tested for vesicle-associated membrane protein 2 (VAMP2), voltage-dependent anion channel 1 (VDAC1) and translocase of outer mitochondrial membrane 40 (TOMM40) and could confirm depletion on day 3 post-infection both on ER/SnCV as well as whole cell level (Fig. 3F and G). These results demonstrate that identification and quantification of these proteins by mass spectrometry and statistical prediction of protein regulation for day 3 post infection correlated with protein levels in ER/SnCV membranes detected by immunoblotting.

Endosomal trafficking is required for progeny formation

Chlamydia has developed mechanisms to avoid lysosomal degradation and acquire lipids via interception of Golgi-derived sphingomyelin-rich vesicles (Hackstadt et al., 1995; Scidmore et al., 2003). As acquiring lipids by modulation of cellular trafficking is absolutely essential for Chlamydia and proteins associated with anterograde transport appear to be depleted from the ER/SnCV fraction (Fig. 3C), we systematically searched our proteomics data for factors involved in vesicular trafficking. We detected several additional major regulators of endosomal trafficking (Supplementary Tables S1 and S7). STRING-based network analysis (von Mering et al., 2003) of these hits showed that endosomal and exocytic transport as well as COPII-dependent and -independent ER to Golgi transport were depleted from the ER/SnCV fraction upon infection with Simkania (Fig. 4A and corresponding values in Supplementary Table S7). In contrast, proteins of the endosomal recycling pathway were mainly unchanged, whereas COP I transport was partially enriched (Fig. 4A). Clathrin-dependent Golgi to ER transport proteins were enriched in ER/SnCV (Fig. 4A).

This pattern of regulation indicated that Simkania might require retrograde transport via clathrin- and Golgi-associated vesicles for nutrient/metabolite acquisition and growth. To test this assumption, we performed infectivity assays in cells, which were treated with inhibitors of retrograde trafficking (Retro-1, Retro-2 and VP-184). Retro-1 and Retro-2 have been selected on their activity to interfere with retrograde transport of ricin toxin and Shiga...
Bacterial AB5-toxins, like Cholera- and Shiga toxins, are variations in SnCV formation. Inhibitors of retrograde trafficking cause phenotypic variations in SnCV formation. Bacterial AB5-toxins, like Cholera- and Shiga toxins, are transported along the retrograde route from the PM to the trans-Golgi network (TGN) to reach the ER (Sandvig and van Deurs, 2002). Retro-1 and -2 specifically block the early endosome-to-Golgi transport of these toxins without affecting essential endogenous cargo proteins, protein biosynthesis or morphology of the ER (Stechmann et al., 2010; Noel et al., 2013). As all tested inhibitors of retrograde transport have had an effect on primary and progeny infection for Simkania (Fig. 4B–E), we inspected the subcellular structure of infected cells by TEM (Fig. 5). A strong inhibition of Simkania replication was observed at a concentration of 75 μM for Retro-1, -2 and 25 μM for VP-184 (Fig. 4B–E). SnCVs formed normally in DMSO-treated control cells (Fig. 5A). The SnCV was smaller in Retro-1-treated cells, and less sub-vacuoles were visible (Fig. 5B). Retro-2 affected the SnCV morphology as sub-vacuolar membranes were highly enlarged and contained excessive membrane material, probably as a consequence of defective membrane fusion and vacuole formation (Fig. 5C). Parts of the enlarged sub-vacuoles contained just few bacteria located at the inclusion boundary, which may indicate defective Simkania replication. Additionally, sub-vacuoles were surrounded by multiple membranes at several areas indicative of a role of Retro-2 inhibited pathways in SnCV formation and membrane fusion defect. VP-184 seemed to have an effect on cell growth and development because infected cells appear smaller than control cells (Fig. 5D). In line with this, SnCV size and morphology was also affected in about half of the cells (Fig. 4C). Surprisingly, sub-vacuoles contained small transparent vesicles of unknown origin and mitochondria were associated with vesicular structures similar to lipid droplets. Again cellular effects caused by VP-184 strongly influence SnCV formation and Simkania replication. The discovery of vesicular structures inside the SnCV and at the mitochondria indicated a possible shift in lipid metabolism. In summary, all three Retro compounds have unique effects on the SnCV morphology suggesting that retrograde transport is essential for normal development of Simkania.
Fig. 3. Network analysis of proteome data.
A. Heatmap of sample adjusted spectral counts (samples 1 and 2), of edgeR differentially expressed proteins only (raw P-value < 0.05 to 0.2).
B. Pathway enrichment of 51 proteins with adjusted P-values ≤ 0.2 analysed with the Ingenuity software (see Supplementary Table S2). Bars show the −log (P-value) for the enrichment of proteins of a given group within the list of selected proteins using a right-tailed Fisher’s exact test. Orange squares show the ratio of proteins detected (within the selected list) among all proteins associated with that group.
C–E. Proteins were clustered into networks of protein–protein interaction based on Ingenuity analysis. Network clustering show e.g. (C) downregulation of mitochondrial proteins and anterograde transport, de-regulation of cytoskeletal components, (D) upregulation of mRNA charging and (E) various metabolic functions. The intensity of green and red molecule colours indicates the degree of depletion or enrichment respectively. Symbols reflect protein class e.g. enzyme and connections depict the way proteins interact (see Supplementary Fig. S5 for additional information).
F. Immunoblot analyses of crude purified ER-SnCV (CMF) membrane fractions. Calnexin and glycerinaldehyde-3-phosphate dehydrogenase (GAPDH) were unchanged in ER-SnCV membranes after infection; T-complex 1 related (TCP1/CCT2) and Rho guanine nucleotide exchange factor 2 (ARHGEF2/GEF H1) were enriched; voltage dependent anion channel 1 (VDAC1) and translocase of outer mitochondria membrane 40 (TOMM40) were depleted.
G. Immunoblot analyses of whole cell lysates from 3 day time-course experiments of infected and control cells. Calnexin and GAPDH are unchanged in whole cell lysate at 3 days post infection. ARHGEF2/GEF H1 is weakly enriched, TCP1/CCT2 is unchanged, whereas VDAC1, TOMM40 and VAMP2 (vesicle-associated membrane protein 2) are depleted during infection. Actin was used as loading control.

(Hackstadt et al., 1995; Scidmore et al., 1996b; Beatty, 2006; 2008; Robertson et al., 2009). To test if inhibitors of retrograde transport directly affect cellular transport processes to the SnCV, we analysed lipid transport using the fluorescent ceramide derivative C6-NBD-ceramide in live cell imaging experiments (Fig. 6A–F, Supplementary Figs S3 and S4). This fluorescent ceramide provides a vital stain for the Golgi apparatus (Lipsky and Pagano, 1985b; Pagano et al., 1989; Rosenwald and Pagano, 1993) and is also transported to Chlamydia inclusions (Hackstadt et al., 1995). NBD-labelled-glucosylceramide and -sphingomyelin are transported from the PM retrograde to the Golgi complex across early and recycling endosomes (Babia et al., 2001). C6-NBD-ceramide transport to Golgi (mRFP-labelled) and SnCV was monitored over a time period of 30 min at 1 frame/30 s (Fig. 6A–C) in the absence or presence of inhibitors of retrograde transport or the previously described Golgi disrupting agent Brefeldin A (BFA) (Hackstadt et al., 1995; Scidmore et al., 1996a; Babia et al., 2001) (Fig. 6D–F, Supplementary Figs S3 and 4). As expected, the C6-NBD-ceramide transfer to mRFP-fluorescent Golgi followed a kinetic similar to carrier-mediated transport. Signal intensity at the Golgi was lower during infection but followed a similar kinetic (Fig. 6A). Fluorescent ceramide that was not taken up by the Golgi and/or the SnCV was detected in the cell cytosol (Fig. 6B and C). Analyses of untreated ER-KDEL fluorescent cells showed that C6-NBD-ceramide transport ends at the Golgi apparatus and does not reach the ER (Supplementary Fig. S3A). C6-NBD-ceramide transport was faster and signal intensity increased at the Golgi in VP-184-treated non-infected cells compared with control (Fig. 6D and Supplementary Fig. S3E). In contrast, Simkania-infected cells showed a strongly reduced C6-NBD-ceramide transport to the Golgi in the presence of VP-184 (Fig. 6E, Supplementary Fig. S4B). In accordance with this observation, VP-184 also strongly reduced the C6-NBD-ceramide signal at the SnCV (Fig. 6F, Supplementary Fig. S4B). C6-NBD-ceramide slightly accumulated in the Golgi in Retro-2-treated cells and infection strongly increased this effect (Fig. 6D and E, Supplementary Figs S3D and S4D). Thus, Retro-2 seemed to optimise retrograde lipid-trafficking to the Golgi per se but especially during infection. This was also observed at the level of C6-NBD-ceramide SnCV transport, which was increased in Retro-2-treated cells (Fig. 6F, Supplementary Fig. S4D). Retro-1 had the similar although milder effect as Retro-2 (Fig. 6D–F, Supplementary Figs 3C and 4C). BFA causes disassembly of the Golgi apparatus and thereby is expected to block C6-NBD-ceramide uptake by the Golgi. We observed Golgi fragmentation 30 min after BFA treatment that was followed by reduced fluorescent lipid uptake and Golgi transport in both non-infected and infected cells (Fig. 6D and E, Supplementary Figs 3B and 4A). Interestingly, the SnCV showed reduced but not completely abolished C6-NBD-ceramide levels in the presence of BFA (Fig. 6F). This result indicates possibly a secondary lipid-transport route to the Simkania inclusion. Thus, the Retro compounds affect C6-NBD-ceramide transport to the Golgi and to the SnCV, which may explain their effect on SnCV morphology and bacterial development.

Discussion
Proteomic analysis of isolated pathogen containing vacuoles has been successfully applied to many different bacteria to characterise these unique cellular compartments (Herweg et al., 2015). Proteome analysis of the Chlamydia-containing vacuole revealed the recruitment of host cell PM receptors and retromer complexes (Aeberhard et al., 2015; Mirrashidi et al., 2015; Subbarayal et al., 2015). Here, we applied a proteomics approach to investigate the composition of the SnCV and the host ER with the aim to identify changes in protein composition that could give us hints on the intracellular accommodation of Simkania. A crucial step towards this aim was the crude purification of the SnCV together with ER membranes, which was probably possible due to a close association of
these membranes and ER contact sites. Although direct information on trafficking pathways cannot be deduced from these proteomics data, the composition of the compartment can be estimated. We could use these semi-quantitative data and protein interaction predictions to identify complex networks of interacting proteins including transport processes with infection-dependent enrichment or depletion.

Intracellular pathogens frequently escape from the phagolysosomal route to prevent lysosomal degradation. Although initially believed to be exceptions, pathogen-containing compartments with ER or Golgi features (Desjardins, 2003) have been meanwhile demonstrated for the important pathogens Brucella (Anderson and Cheville, 1986; Pizarro-Cerda et al., 1998), Legionella (Horwitz, 1983; Swanson and Isberg, 1995; Tilney et al., 2001) and Chlamydia that replicate in vacuoles receiving lipids from the TGN (Heinzen et al., 1996). Legionella and Brucella interfere with the biogenesis of the early phagolysosome modulating their vacuolar composition to mature into ER-derived compartments (Desjardins, 2003). Like many other bacteria, all Chlamydiae develop and multiply within a unique vacuole, which is not acidified (Fris, 1972; Lawn et al., 1973; Wyrick and Brownridge, 1978; Heinzen et al., 1996; Al-Younes et al., 1999; Rockey et al., 2002). However, also Chlamydia-like organisms, such as Parachlamydia acanthamoebae, Simkania negevensis and Waddlia chondrophila grow in similar vacuoles in amoebae and cells of higher animals (Evérott, 2000). Similar to Legionella pneumophila and Brucella abortus, C. trachomatis uses host cell Golgi- and/or ER-derived vesicles to generate specialised membrane-bound compartments for the replication. We previously showed that Simkania establishes novel ER interaction sites at which the SnCV is extended along the ER within 3 days post infection (Mehlitz et al., 2014). It is likely that Simkania uses host cell Golgi- and/or ER-derived lipids and vesicles to generate the SnCV compartment.

Positive-strand RNA viruses are known to remodel intracellular membranes to create mini-organelles where RNA amplification and virion assembly take place (Miller and Krijnse-Locker, 2008). Flaviviruses like dengue virus and West Nile virus have been shown to replicate within membrane invaginations originating from the ER (Welsch et al., 2009; Gillespie et al., 2010). Hepatitis C virus most likely uses specialised vesicles for RNA replication that are derived primarily from the rough ER, early and late endosomes, COP vesicles, mitochondria and lipid droplets (Romero-Brey et al., 2012). We identified analogue cellular components in the proteome data indicating that vacuole formation processes are similar for Simkania. Interestingly, most proteins identified here are part of clathrin and COPI vesicle transport pathways (retrograde) that were enriched during infection whereas most exocytic and endosomal as well as COPII or other ER to Golgi transport proteins (anterograde) were depleted. This was the first hint that retrograde transport plays an important role in SnCV formation. Additionally, silencing COPB2 (COPI) and AP2B1 (clathrin) expression reduced the production of Simkania infectious progeny.

We demonstrated here a strong effect of Retro compounds on SnCV maturation and Simkania development. These inhibitors have initially been identified as compounds that prevented the cytotoxic effect of toxins that are transported via the retrograde route (Stechmann et al., 2010; Ming et al., 2013; Noel et al., 2013). Although the exact molecular targets of these compounds have not been identified yet, it is likely that the retrograde transport
Fig. 5. Effects of Retro compounds on SnCV formation during Simkania infection. Transmission electron microscopy of HeLa229 cells 3 days post-infection. Cells were treated with 75 μM DMSO, Retro-1, -2 and 25 μM VP-184 during infection. The SnCV (red) (Bravo et al.), associated to mitochondria (green) (Kahane et al.), spreads the cytoplasm near the nucleus (blue). A. The DMSO control shows a normal SnCV highly filled with bacteria that are covered with mitochondria. B. Retro-1 seems to affect SnCV size indicated by fewer small sub-vacuoles. C. Retro-2 seems to affect SnCV morphology and bacteria replication. Sub-vacuoles that are highly enlarged contain just few bacteria that are located at the SnCV boundary (blue arrows) and/or contain excessive membrane material (orange arrows). Some sub-vacuoles are surrounded by multiple membranes (yellow arrows). In total, the SnCV appears to be bigger in size. D. VP-184 affects cell development and SnCV size. Sub-vacuoles contain freestanding transparent vesicles of unknown origin (yellow arrows). Some mitochondria are associated with transparent droplets of unknown origin (white arrows).
Fig. 6. Effects of Retro compounds on Golgi-dependent sphingomyelin transport to the SnCV. Time-courses of life cell imaging from intracellular C6-NBD-ceramide transport in Simkania-infected or control HeLa Golgi-mRFP cells are shown. (A, D–E) Calculation of fluorescence intensity of C6-NBD-ceramide trafficking to the Golgi apparatus and (F) to the SnCV within 30 min (2 images/frame). Curves are representative of n = 3 biological experiments and m ≥ 20 cells/experiment. (B–C) Time-course images of C6-NBD-ceramide transport in untreated Simkania- and non-infected HeLa cells. Red = Golgi; green = C6-NBD-ceramide. Brefeldin A = BFA. Images are representative of n = 3 independent experiments.
A. C6-NBD-ceramide transport to the Golgi rises exponentially. Simkania-infected cells (orange) absorb more slowly and less of the fluorescent ceramide compared to non-infected cells (blue). The maximum gain is reached at around 28–30 min.
B. Time-course images of C6-NBD-ceramide transport in non-infected cells. Images of 2.5 min time lags starting at 0 min are illustrated.
C. Time-course images of C6-NBD-ceramide transport in Simkania-infected cells. Images of 2.5 min time lags starting at 0 min are illustrated.
D. C6-NBD-ceramide to Golgi transport in non-infected and Retro compound-, BFA- or untreated cells was analysed. Inhibitors were added 30 min prior to imaging (for more details, see Experimental procedures section). VP-184-treated cells (violet) showed a high increase that slightly dropped down at 12–13 min. Retro-2-treated cells (grey) showed a slight increase compared with the control (blue). Retro-1-treated cells (red) (Bravo et al.) showed a slight decrease. In BFA-treated cells (yellow) the transport was really weak.
E. C6-NBD-ceramide to Golgi transport in Simkania-infected and Retro compound-, BFA- or untreated cells was analysed. In Retro-1- and -2-treated cells (red, grey) the fluorescent lipid transport to the Golgi was highly increased. In VP-184 and BFA-treated cells (violet, yellow), the transport was weaker but still detectable.
F. C6-NBD-ceramide to SnCV transport in Simkania-infected and Retro compound-, BFA- or untreated cells was analysed. In Retro-1- and -2-treated cells (red, grey) the fluorescent lipid transport to the SnCV was highly increased. VP-184-treated cells (violet) showed a slight increase that dropped down nigh the Sn-control (orange). In BFA-treated cells (yellow) the transport was weaker but still detectable.
is already blocked early at the level of endosomal to TGN trafficking. We showed that Retro compounds influence intracellular SnCV formation and Simkania development. Electron microscopy studies indicated a strong effect on the SnCV morphology that appears to be unique for every compound. This diversity may reflect the inhibition of different targets by Retro compounds and—in line with this assumption—also different pathways of vesicular transport that all are required for SnCV formation. In the process of inclusion formation, Chlamydiae actively interrupt the classical endocytic pathway very early (Scidmore et al., 1996a) and enter the cells’ exocytic pathway that remains poorly defined (Hackstadt et al., 1995; 1997; Rzomp et al., 2003). In this study, we concentrated on retrograde trafficking as proteins controlling clathrin vesicle and COPI-Golgi to ER transport appeared enriched during Simkania infection. Retrograde transport involves proteins and lipids that initially are transported back from the PM and endosomes to the Golgi apparatus. Recycling endosomes serve here as a sorting organelle for retrograde transport (Bonifacino and Rojas, 2006). Subsequently proteins and lipids are transported within the TGN to the ER (Sandvig and van Deurs, 2005), which is mediated by COPI vesicles (Orci et al., 1997). Many viruses use similar routes to the ER for their assembly (Brandenburg and Zhuang, 2007). Recently, it was reported that Retro-2 can inhibit retrograde transport of polyoma-, papilloma- and adeno-associated viruses suggesting an overlap in the host factors used by bacterial toxins and viruses (Nelson et al., 2013; Carney et al., 2014; Nonnenmacher et al., 2015). Our data using Retro compounds and RNA interference during Simkania infection support that the SnCV is dependent on retrograde transport. It is likely that retrograde transport is required for nutrient acquisition during Simkania infection. Therefore, compounds like Retro-1 or -2 might be useful to hinder infection by Simkania and maybe Chlamydia as well, as they also require early endosome-to-Golgi and Golgi to ER transport for infection.

Metabolite delivery to the SnCV appears to be one of the processes that are influenced by Retro-1 and -2. Treatment with Retro-1 and -2 increased the amount of 1,2-(NBD)-ceramide in the Golgi and the SnCV, probably due to the interference of both inhibitors with the transport of 1,2-(NBD)-ceramide from the TGN, which could result in retention in the Golgi and increased uptake by the SnCV. VP-184 had the most dramatic effect by causing strong retention in the Golgi only in non-infected cells and blocked 1,2-(NBD)-ceramide transport to the Golgi after infection. In infected cells, the effect of VP-184 was similar to BFA, a Golgi-disrupting compound, and lipid accumulation in the Golgi was not detected at all, pointing to an infection-dependent block of lipid transport at the Golgi under these conditions. 1,2-(NBD)-ceramide transport to the SnCV seemed to be affected by this block at later time points. The finding that BFA prevented accumulation of 1,2-(NBD)-ceramide in both the Golgi and the SnCV adverts to a role of the Golgi in lipid transport to the SnCV. The chlamydial inclusion is not permeable to molecules as small as 520 Daltons (Heinzen and Hackstadt, 1997) and therefore, all larger metabolites and proteins have to cross the membrane barrier either by direct transport or by delivery via vesicle involving both human and bacteria proteins in the inclusion membrane. We used 1,2-(NBD)-ceramide, a vital stain for the Golgi apparatus that has been used to study sphingolipid trafficking in viable cells (Lipsky and Pagano, 1985a; b; Pagano et al., 1989; Rosenwald and Pagano, 1993; Zhong et al., 2001). Within the cis or medial Golgi apparatus, 1,2-(NBD)-ceramide is processed to sphingomyelin (SM) or glucosylceramide, like endogenous ceramide (Lipsky and Pagano, 1985a), and delivered to the PM by a vesicle-mediated pathway. By interruption of exocytic lipid transport, C. trachomatis (Scidmore et al., 1996b), C. psittaci (Rockey et al., 1996) and C. pneumoniae (Wolf and Hackstadt, 2001) intercept the fluorescent SM that subsequently is incorporated into the cell wall of the intracellular bacteria. This modification of intracellular lipid trafficking by induced fusion of the inclusion with a subset of exocytic vesicles leads to the acquisition of a significant amount of the exported SM (Fields and Hackstadt, 2002). Simkania like other Chlamydia species (Fritsche et al., 2000) lack genes for SM biosynthesis and therefore likely intercept SM by exploiting the host cell ceramide transport. Prolonged treatment with BFA causes disassembly of the Golgi apparatus and induces redistribution of Golgi, secretory and membrane proteins into the ER. BFA has been shown to abrogate SM transport to the chlamydial inclusion (Scidmore et al., 1996b). We observed that BFA generally blocks 1,2-(NBD)-ceramide uptake, indicating an early inhibition of ceramide transport already at the PM. Despite advanced Golgi fragmentation in BFA-treated cells, small amounts of ceramide were incorporated into the SnCV, suggesting that Simkania gains the main part of ceramide directly from the Golgi but uses also other pathways for lipid acquisition. The enhanced accumulation of ceramide in the SnCV could be directly stimulated by the blocked transport to or out of the Golgi (treatment with Retro-1, Retro-2, perhaps also VP-184) resulting in increased transfer of ceramide to other transport pathways, e.g. clathrin-derived vesicles. This is supported by similar findings on retrograde toxin transport in Vero cells (Schapiro et al., 1998) as well as enrichment of clathrin-associated proteins like AP2A1 and AP2B1 found in our LC-MS² analysis. BFA treatment has no effect on Chlamydia replication (Scidmore et al., 1996b), suggesting that C. trachomatis acquires SM by additional routes that involve BFA-insensitive and/or non-vesicular trafficking pathways.
(Derre et al., 2011; Elwell et al., 2011; Dumoux et al., 2012). These may involve bacterial effector proteins as in case of Salmonella, where the SPI-2 effector SseL binds to a lipid-binding protein and may direct host lipid-transport to the Salmonella-containing vacuole (Auweter et al., 2012).

We show here that early events in retrograde transport seem to be essential for SnCV formation, Simkania development and ceramide acquisition. Because we do not know the targets of the Retro compounds, the affected trafficking pathways remain to be defined. We cannot exclude COPI-dependent and independent pathways of retrograde transport, the ERGIC as first anterograde and retrograde sorting station or the influence of depleted anterograde transport proteins during infection. We hypothesise that the enrichment of clathrin or Golgi to ER transported proteins indicates a direct bypass of important nutrient trafficking pathways to the SnCV during infection. Further studies are necessary to understand the influence of retrograde and anterograde transport on Simkania development.

**Experimental procedures**

**Cell lines and bacteria**

HeLa229 (ATCC CCL-2.1) were grown in RPMI1640 medium (Glutamax, 10% FBS, w/o HEPES) (Invitrogen). Stable HeLa229 cell lines were established to constantly label the Golgi apparatus (B4Gat1 in a pCMV6-AC-mRFP cloning vector, OriGene) and the ER (KDEL in a pDsRed2-ER expression vector). Selection for B4Gat1-mRFP or KDEL-DesRed2 was done with G418 (400 μg/ml). *S. negevensis* strain Z (ATCC VR-1471) was prepared as described previously (Mehlitz et al., 2014). Briefly, HeLa229 cells were grown to 50–70% confluence and were inoculated with *S. negevensis* in RPMI1640 with 5% FBS, for 6 h at 35°C in a humidified incubator at 5% CO2. Medium was replaced by infection medium (RPMI1640, Glutamax, 5% FBS, w/o HEPES), and growth was allowed for 3 days. Cells were mechanically detached, and bacteria were released using 2–5 mm glass beads (Carl Roth). Low speed supernatant (600 × g, 4°C and 5 min) was subjected to high-speed centrifugation (20 000 × g, 4°C and 30 min) to pellet bacteria. Bacteria were washed twice with 5 ml SPG (250 mM sucrose, 50 mM sodium phosphate, 5 mM glutamate, pH 7.4), aliquoted and stored at −80°C in SPG.

**ER-SnCV purification**

HeLa229 cells were seeded in 6-well cluster plates (12 plates/experiment) and grown to 70% confluence before infection with Simkania at multiplicity of infection (moi) 1.0 (see also Fig. 1A for experimental scheme). Plates were centrifuged for 1 h at 4°C, 910 × g and inoculated for 5 h at 35°C in a humidified incubator at 5% CO2. Medium was replaced by infection medium and growth was allowed for 3 days. Growth medium was removed, and cells were released with a rubber policeman into ice-cold PBS followed by one wash step ice cold PBS (5 min, 4°C, 600 × g). PBS was replaced by 600 μl 0.5 × hypotonic buffer (10 mM HEPES, 0.5 mM EGTA, 12.5 mM KCl, 0.125 M sucrose, protease and phosphatase inhibitors, pH 7.6), and cells were incubated for 20 min at 4°C in a rotary shaker. Cells were pelleted (5 min, 4°C, 600 × g) and resuspended in 0.5 × isotonic buffer (5 mM HEPES, 0.5 mM EGTA, 12.5 mM KCl, protease and phosphatase inhibitors, pH 7.6) and sonicated at 4°C for 10 min in a sonication bath. Prelysate was homogenised with a Dounce Homogenizer (glass cylinder and pestle), 10 strokes at 200 r.p.m. Crude microsomal fraction containing ER and SnCV were prepared by sequential centrifugation: (i) pre-nuclear fraction (PNF) – 1,000 × g at 4°C for 10 min; (ii) pre-mitochondrial fraction (PMF) – 12,000 × g at 4°C for 15 min and (iii) crude microsomal fraction (CMF) – 100,000 × g at 4°C for 1 h in an Optima MAX-XP ultra-centrifuge using a MLA-80 fixed angle rotor (Beckmann Coulter).

**Reagents and antibodies**

Chemicals were obtained from Sigma-Aldrich (Germany) unless otherwise stated. Primary antibodies used: Calnexin (Cell Signaling, C5C9), PDI (Cell Signaling, C81H6), Erp72 (Cell Signaling, D70D12), KDEL (Enzo, 10C3), Lamin B1 (Santa Cruz, 6216), Sam50 (kindly provided by Vera Kozjak-Pavlovic), β-Actin (Sigma-Aldrich, A5411), CCT2 (Cell Signaling, 3561), GEF H1 (55B6) (Cell Signaling, 4076), VDAC1 (Calbiochem, 529532), TOMM40 (Santa Cruz, sc-11414) and VAMP2 (D601A) (Cell Signaling, 13508). Antibodies against GroEL (anti-Sn-GroEL) or whole bacteria (anti-Simkania) were prepared as previously described (Mehlitz et al., 2014). Carbocyanine-labelled secondary antibodies were purchased from Dianova. Nuclear staining was done with 4',6-diamidino-2-phenylindole (DAPI). Fluorescent sphingolipid chased from Dianova. Nuclear staining was done with 4',6-diamidino-2-phenylindole (DAPI). Fluorescent sphingolipid NBD-C2-ceramide (6–(N–[7–nitrobenz–2–oxa–1,3–diazol–4–yl] amino) hexanoyl) sphingosine) complexed to BSA was obtained from Molecular Probes (Invitrogen, N-1154). G418 was obtained from InvivoGen. Protease and phosphatase inhibitors were obtained from Roche.

**Immunoblotting and immunofluorescence staining**

Western blotting was performed according to standard procedures (Mehlitz et al., 2010), and signals were detected with a Chemo Cam Imager (Intas). Quantification was performed using FIJI (ImageJ) and Excel (Microsoft). For immunofluorescence, cells were seeded in 12-well cluster plates with or without coverslips and were infected in a humidified incubator at 35°C and 5% CO2 with mois indicated in the respective experiments. Cells were fixed at indicated time points with 4% PFA-sucrose for 25 min, washed once with PBS and stained while gently shaking. Subsequently cells were permeabilised with 0.2% Triton X-100 in PBS for 45 min and washed 3 × with PBS for 15 min at RT. Cells were blocked with 2% goat serum in PBS for 60 min and were stained with primary antibodies diluted in 2% goat serum in PBS for 24 h at 4°C. After 3 × washing with PBS for 15 min, samples were incubated with secondary antibodies for 1 h in blocking solution at RT in the dark. Cells were washed 3 × with PBS and counterstained with DAPI (1 μg/ml) in PBS for 15 min. Cov-
Table 1. Non-linear gradient operating parameters for peptide elution.

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<th>Duration (Ming et al.)</th>
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Electron microscopy

Samples were fixed with 2.5% glutaraldehyde (50 mM sodium cacodylate pH 7.2; 50 mM KCl; 2.5 mM MgCl2) at 4°C for 1 h and buffered with 50 mM sodium cacodylate (pH 7.2) at 4°C. Then samples were incubated with OsO4 for 2 h at 4°C. Cells were dehydrated, embedded in Epon812 and ultrathin sectioned at 50 nm. Sections were stained with 2% uranyl acetate in ethanol, followed by staining with lead citrate and analysed on a Zeiss EM10 (Zeiss, Germany).

LC-MS/MS

For LC-MS\(^2\) measurements, three biological replicates of ER-SnCV purified samples were generated and analysed. Proteins were separated by 1D-SDS-PAGE, the gel lanes were excised in 10 equidistant pieces and subjected to trypsin digestion as described before (Otto et al., 2010). For LC-MS/MS analyses, in-house self-packed columns were prepared and used with an EASY-nLC II system (Thermo). In brief fused silica emitter tips with an inner diameter of 100 μm and an outer diameter of 360 μm were prepared by using a P-2000 laser puller (Sutter Instruments, Novato, CA). The resulting emitter tip was then packed with Phenomenex Aeris C18 reversed phase material (3.6 μm particles) in custom build pressure bomb to obtain a 20 cm nano-LC column.

The peptides were loaded onto the column with 10 μl buffer A (0.1% acetic acid) at a constant flow rate of 500 nL/min without trapping. Subsequently, peptides were eluted using a non-linear 180 min gradient from 1% to 99% buffer B (0.1% acetic acid in acetonitrile) with a constant flow rate of 300 nL/min and injected online into the mass spectrometer. Voltages between the emitter capillary and the orifice were set to 2.5 kV by using liquid junction. Afterward, the column was washed with 99% buffer B for 5 min and equilibrated with 99% buffer A for 18 min. The gradient is described in Table 1.

MS and MS/MS data were acquired with a LTQ Orbitrap Classic (Thermo). After a survey scan at a resolution of R = 30 000 within a scan range (m/z) of 300–2000 in the Orbitrap with activated lockmass correction, the five most abundant precursor ions were selected for fragmentation. Singly charged ions as well as ions without detected charge states were not selected for MS/MS analysis. CID fragmentation was performed for 30 ms with normalised collision energy of 35, and the fragment ions were recorded in the linear ion trap.

After mass spectrometric measurement, MS data were subjected to database searching via Sorcerer using Sequest (version 27, revision 11; SageN, Milpitas, CA, USA) without charge state deconvolution and deisotoping performed. Database searching was performed with a database of combined entries of NCBI Simkania negevensis NC 015713 and NC 015710 and UniProtKB Homo sapiens (version 2/12). The combined database was used as target/decoy databases with a list of common contaminants added (45580 entries). Sequest was used assuming trypsinisation with a fragment ion mass tolerance of 1.00 Da and a search tolerance of 10 ppm for the overview scans. Oxidation of methionine was specified in Sequest as a variable modification. Scaffold (version 3.5.1, Proteome Software, Portland, OR, USA) was used to filter and validate MS/MS-based peptide and protein identifications. Peptide identifications were accepted when spectra exceeded Xcorr values of 2.2, 3.3 and 3.8 for doubly, triply and quadruply charged peptides with deltaCN values of more than 0.1. Protein identifications are based on at least 2 unique peptides. Proteins that contained similar peptides and could not be differentiated based on MS/MS analysis alone were grouped to meet the principle of parsimony.

Analysis of proteomics data

Proteomics data were analysed via Web-based Gene SeT Analysis Toolkit (WEBgestalt) (Wang et al., 2013). Enrichment analysis was performed using GO Analysis against the Homo sapiens genome. WEBgestalt settings were hypergeometric statistical method and multiple test adjustment according to Benjamini and Hochberg (1995). Significance levels allowed only the top 10 categories at a minimum protein number of 10 per category. Proteomic count data was filtered using Scaffold as described above (LC-MS/MS). Differential expression was analysed using the edgeR method (Robinson et al., 2010) as implemented in the statistical package mssTest (Gregori et al., 2013) in the software R. Given the uncertainty of normalisation factor estimation due to the relatively small in number and possibly selected set of proteins detected we did not normalise counts across samples by default. Estimated normalisation factors for paired samples showed that those were rather small (< 0.1-fold difference for all pairs). We used a model incorporating each experiment as blocking factor to perform paired tests. As a sensitivity analysis, we repeated the analysis using edgeR’s global normalisation method, which resulted in the same list of differentially expressed genes (adjusted P < 0.2) except for proteins HNRPC, K1C18, K2C8, SFXN1, AHNK, PABP1, ATPO and SEPT2. Heatmaps were generated after a variance stabilising transformation as provided by package DESeq (Love et al., 2014), adjusting for experiment effects using the same model.
as above. Trafficking proteins were further analysed via ‘Search Tool for the Retrieval of INteracting Genes/proteins’ (STRING) (von Mering et al., 2003) allowing standard setting for predictive methods (Neighbourhood, Gene Fusion, Co-occurrence, Co-expression, Experiments, Databases and Textmining) at medium confidence (0.4).

**Infectivity assays**

The 40 000 cells were seeded in 12-well cluster plates, inhibitor-treated and infected as indicated in the respective experiment. Cells were processed for immunoblotting, immunofluorescence staining or infectivity assay. For infectivity assays cells were either fixed and stained at indicated time points (inclusion formation/primary infection) or bacteria were released via one freeze thaw cycle (−70°C/37°C) followed by mechanical release through pipetting and transfer to fresh HeLa229 cells (1:25–1:50, progeny/infectivity). Cells were centrifuged for 1 h at 35°C and medium exchanged to infection medium. Progeny was fixed at day 3 post infection and processed for staining or harvested for immunoblotting. Infectivity assays were imaged on an automated fluorescence microscope Leica DMIR. Numbers and average sizes of the SnCV as well as host cell numbers were determined via GroEL and DAPI staining, and images were analysed and quantified using FIJI (ImageJ) and Excel (Microsoft).

**siRNA transfections**

SiRNA against STX5 (GS6811), COPB2 (GS9276), AP2B1 (GS163), All-Stars negative control (SI03650325) were purchased from Qiagen. ARHGEF2 siRNA was obtained from Dharmacon (M-009883-01-0005). Transfection was done using HiPerFect transfection reagent (301705) as described in the manufacturer’s instructions (Qiagen).

**RNA isolation and qRT-PCR**

RNA was isolated using TRI reagent solution (Ambion, AM9738) according to manufacturer’s instructions. Instead of BCP, we used 200 µl chloroform. Total RNA concentration was determined by using a NanoDrop fluorospectrometer (Thermo Scientific). Synthesis of first strand cDNA was performed on 1–2 µg total RNA by using RevertAid RT Reverse Transcription Kit (Thermo Fisher, K1691). qRT-PCR was performed on 20 µl reaction volumes in triplicates using SYBR green master mix (Quanta, 733–1386) according to the manufacturer’s instructions. qRT-PCRs were performed on a green master mix (Quanta, 733–1386) according to the manufacturer’s instructions (Qiagen). qRT-PCR was performed on 1–2 (Thermo Scientific). Synthesis of first strand cDNA was performed by using a NanoDrop fluorospectrometer (Thermo Scientific). RNA isolation and qRT-PCR were determined by using a NanoDrop fluorospectrometer (Thermo Scientific). Synthesis of first strand cDNA was performed on 1–2 µg total RNA by using RevertAid RT Reverse Transcription Kit (Thermo Fisher, K1691). qRT-PCR was performed on 20 µl reaction volumes in triplicates using SYBR green master mix (Quanta, 733–1386) according to the manufacturer’s instructions. qRT-PCRs were performed on a green master mix (Quanta, 733–1386) according to the manufacturer’s instructions (Qiagen).

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**Supporting information**

Additional supporting information may be found in the online version of this article at the publisher’s web-site.