Endocytosis of a Glycosylphosphatidylinositol-anchored Protein via Clathrin-coated Vesicles, Sorting by Default in Endosomes, and Exocytosis via RAB11-positive Carriers

Christoph G. Grünfelder,* Markus Engstler,† Frank Weise,* Heinz Schwarz,‡ York-Dieter Stierhof,§ Gareth W. Morgan,‖ Mark C. Field,‖ and Peter Overath*¶

*Max-Planck-Institut für Biologie, Abteilung Membranbiochemie, D-72076 Tübingen, Germany; †Ludwigs-Maximilians-Universität München, Department Biologie I, Bereich Genetik, D-80638 München, Germany; ‡Max-Planck-Institut für Entwicklungsbiologie, D-72076 Tübingen, Germany; §Zentrum für Molekularbiologie der Pflanzen, Universität Tübingen, D-72076 Tübingen, Germany; and ‖Wellcome Trust Laboratories for Molecular Parasitology, Imperial College London, Department of Biological Sciences and Centre for Molecular Microbiology and Infection, London, SW7 2AY, United Kingdom

Submitted October 10, 2002; Revised December 3, 2002; Accepted 2003
Monitoring Editor: Jennifer-Lippincott-Schwartz

Recently, proteins linked to glycosylphosphatidylinositol (GPI) residues have received considerable attention both for their association with lipid microdomains and for their specific transport between cellular membranes. Basic features of trafficking of GPI-anchored proteins or glycolipids may be explored in flagellated protozoan parasites, which offer the advantage that their surface is dominated by these components. In Trypanosoma brucei, the GPI-anchored variant surface glycoprotein (VSG) is efficiently sorted at multiple intracellular levels, leading to a 50-fold higher membrane concentration at the cell surface compared with the endoplasmic reticulum. We have studied the membrane and VSG flow at an invagination of the plasma membrane, the flagellar pocket, the sole region for endo- and exocytosis in this organism. VSG enters trypanosomes in large clathrin-coated vesicles (135 nm in diameter), which deliver their cargo to endosomes. In the lumen of cisternal endosomes, VSG is concentrated by default, because a distinct class of small clathrin-coated vesicles (50 – 60 nm in diameter) budding from the cisternae is depleted in VSG. TbRAB11-positive cisternal endosomes, containing VSG, fragment by an unknown process giving rise to intensely TbRAB11- as well as VSG-positive, disk-like carriers (154 nm in diameter, 34 nm in thickness), which are shown to fuse with the flagellar pocket membrane, thereby recycling VSG back to the cell surface.

INTRODUCTION

To a large extent, the specific composition of cellular membranes is achieved and maintained by the selective transport of integral membrane components by way of vesicular carriers, which shuttle between different cellular compartments. Paradigmatic are two positive sorting mechanisms: First, trans-membrane proteins in a donor membrane can be specifically incorporated into coated membrane domains, which bud off as transport vesicles and fuse with the target membrane (Kirchhausen, 2000); and second, certain lipids or lipid-anchored proteins are proposed to form liquid-ordered microdomains in the fluid-disordered bulk phase of a membrane and these domains are then considered to be selectively incorporated into carrier vesicles (Simons and van Meer, 1988; Simons and Ikonen, 1997; Brown and London, 2000). Positive selection implies concurrent negative selection or sorting by default, because the concentration of components excluded from coated vesicles or lipid microdomains increases as positively selected proteins and lipids are removed. The remaining membrane can thus be a source for vesicular carriers specifically enriched in unselected components.
We are interested in the cellular sorting of glycosylphosphatidylinositol (GPI)-anchored proteins, a class of cell surface proteins thought to occur in all eukaryotic cells, but studied mainly in mammalian cells, yeast, and kinetoplastid protozoa (McConville and Ferguson, 1993; Ferguson, 1999; Chatterjee and Mayor, 2001). The phosphatidylinositol residue confers on these proteins properties of “big lipids,” whereas the protein moiety allows for their ready detection by immunoochemical techniques and, in some cases, by their enzymatic or ligand-binding properties. Therefore, these proteins have become popular for studies on cellular sorting, and depending on the cell type and the membrane compartments investigated, both positive and negative sorting scenarios have been proposed (Bretscher et al., 1980; Bamezai et al., 1992; Keller et al., 1992; Simons and Ikonen, 1997; Harder et al., 1998; Muñiz et al., 2001; Sabharanjak et al., 2002).

We have chosen the mammalian stage of the parasitic protozoan *Trypanosoma brucei* (see Figure 1 for relevant ultrastructural features of the organism), as a model for the sorting of GPI-anchored proteins for three reasons. First, this and related flagellates abundantly express GPI-anchored proteins. *T. brucei* is covered by a dense coat of the GPI-anchored variant surface glycoprotein (VSG), which comprises 10% of the total cellular protein (Overath et al., 1994; Cross, 1996). Sorting is effective, because the concentration of VSG at the cell surface is 50 times higher than in the endoplasmic reticulum (ER) (Grüntfelder et al., 2002), where the protein is synthesized and the lipid anchor is attached. The concentration in membranes of the Golgi complex and of endosomes is 2.7 and 10.8 times higher than in membranes of the ER, respectively. Thus, the equivalent of the membrane area of the flagellar pocket is internalized every 2 min (Coppen et al., 1987). This observation suggests that VSG is rapidly endocytosed and recycled (Seyfang et al., 1990) and implies effective sorting in the endosomal system.

In this study, we focus on membrane structures involved in uptake and recycling of VSG in *T. brucei*. We show that VSG enters trypanosomes at the flagellar pocket in clathrin-coated vesicles and then associates with endosomes. VSG seems to be concentrated by default in cisternal endosomes by budding of a distinct class of clathrin-coated pits (CCPs) and vesicles, which are depleted in VSG. The cisternal endosomes give rise to VSG-rich, TbRAB11-positive exocytic carriers, which fuse with the flagellar pocket membrane. The results are discussed in comparison to sorting and membrane recycling of GPI-anchored proteins in mammalian cells, in which exocytic carriers recycling from endosomes to the plasma membrane have so far not been characterized (Sönntigsen et al., 2000).

**MATERIALS AND METHODS**

**Immunofluorescence**

Bloodstream stage *Trypanosoma brucei* (strain MITat 1.2) were cultivated in HMI-9 + 10% fetal bovine serum, at 37°C in an atmosphere of 5% CO₂ in air. Cells were harvested at a density of 8 × 10⁶ cells/ml by centrifugation at 1400 × g for 10 min. After three washes with ice-cold trypansomal dilution buffer (TDB; 5 mM KCl, 80 mM NaCl, 1 mM MgSO₄, 20 mM Na₂HPO₄, 2 mM NaH₂PO₄, 20 mM glucose, pH 7.4), the cell density was adjusted to 1 × 10⁶ cells/ml and incubation with 1 mM sulfo-NHS-SS-biotin, 1 mM 7-amino-4-methylcoumarin (AMCA)-sulfo-NHS (both from Pierce) was performed. After 15 min of incubation on ice with 60% HMI-9, 10% fetal calf serum, 50 mM glutathione, pH 9.0. After two final washings with TDB, cells were fixed with 4% paraformaldehyde in phosphate-buffered saline (PBS) for 24 h at 4°C. Throughout the procedure, the cell viability was monitored by microscopy. Permeabilization of trypanosomes was achieved by 15-min incubation in 500 µl of 0.1 M Na₂HPO₄/1 M glycine, pH 7.2, followed by 5-min incubation in 1 ml of 0.05 M Na₂HPO₄/0.5 M glycine/0.1% Triton X-100, pH 7.2. Biotinylated, endocytosed VSG was detected with Alexa Fluor 488-conjugated goat anti rabbit antibodies diluted 1:200 in PBS/1% bovine serum albumin. For immunodetection of TbRAB11A (Jeffries et al., 2001) and clathrin heavy chain (Morgan et al., 2001), respectively, affinity-purified rabbit antibodies diluted 1:200 in PBS/1% bovine serum albumin were used. As secondary reagent Alexa Fluor 488-conjugated goat anti rabbit antibody (1:2000; Molecular Probes) was used.

Image acquisition was performed with a motorized Axioskop2 widefield microscope equipped with a 63×/1.4 numerical aperture oil DIC objective (both from Carl Zeiss, Thornwood, NY), a 2.5× optics, and the CoolSnap HQ cooled (~30°C) charge-coupled device camera (Sony IXC285 interline, progressive-scan charge-couple device). For acquisition of three-dimensional (3D)-images, a
PIFOC objective z-stepper was driven by the piezo-amplifier E662 LVVPZT (Physik Instrumente, Karlsruhe, Germany). The microscopic setup was integrated and controlled using the scripting features of the IPlab for Macintosh software (version 3.5; Scanalytics, Fairfax, VA; for details, see http://www.scanalytics.com/). 3D images were acquired using the optimal sampling density derived from the optical setup and the respective fluorophor (z-step size ≥50 × 100 nm). The point spread function of the microscope was measured using fluorescent 0.17-µm microspheres (PS-Speck microscope point source kit; Molecular Probes). After image acquisition, the raw data were exported to the Huygens system software (version 2.1.8; Scientific Volume Imaging, B.V., Hilversum, The Netherlands; for details, see http://www.sviv.nl/), and digital deconvolution was performed using the “maximum likelihood estimation” algorithm (>80 iterations). The restored image data set was visualized and analyzed with the Imaris software package, featuring the Full 3D and Surpass modules (version 3.1 for IRIX; Bitplane AG, Zurich, Switzerland (for details, see http://www.bitplane.ch/products/imaris/ and http://www.bitplane.ch/products/imarisurpass/).

For animated presentation of 3D data, AMCA-stained cell surfaces were processed using the nonlinear morphological gradient filter implemented in the IPlab software. In this way a 3 × 3 eroded version of the AMCA channel was created and merged with the nonprocessed red and green channels, allowing visualization through the cell surface of endocytosed VSG and TbRAB11 or clathrin heavy chain, respectively. Colocalization analysis of multichannel 3D data series was performed with the colocalization software (version 1.0 for IRIX; Bitplane AG; for details, see http://www.bitplane.ch/products/colocalization/).

**Immunoelectron Microscopy**

Logarithmically growing trypanosomes of variant MITat 1.2 were either cryoimmobilized by high-pressure freezing as described previously (Grünfelder et al., 2002) or chemically fixed for Tokuyasu cryosectioning. Cryo-fixed cells were freeze substituted in 0.5% glutaraldehyde + 0.5% osmium tetroxide (OsO₄) in acetone from −90°C to −40°C for 39 h (Figure 2A) or in 0.5% OsO₄ in acetone from −90°C to −40°C for 36 h, followed by 0.25% OsO₄ and 0.25% gallic acid in acetone for 30 h at −40°C (Figure 3D). After washing with acetone, cell were embedded in Epon. For cryosectioning, chemical fixation was performed in suspension for 5–30 min at room temperature, followed by 1.5–2 h on ice by using final concentrations of 4% formaldehyde in 0.1 M HEPES (pH 7.2) or 2% formaldehyde in 0.1 M HEPES supplemented with 0.05% glutaraldehyde. After fixation, cells were centrifuged, embedded in 1 or 2% low melting temperature agarose (SeaPlaque, Rockland, ME) in PBS, cut into small blocks, washed twice in PBS, and infiltrated with 20% polyvinylpyrrolidone/1.8 M sucrose in 0.1 M NaHPO₄. Specimen blocks were mounted on copper stubs, frozen in liquid nitrogen and trimmed and sectioned (at 60–70 nm) with a Leica Ultracut S/FCS cryo-ultramicrotome. The grids with the thawed cryosections were floated on ice cold PBS for storage before immunolabeling.

Immunolabeling of ultrathin Epon- and cryosections was performed according to standard protocols (Sterhofs et al., 1999). The following antibodies were used: rabbit anti-VSG MITat 1.2 antisera (1:70; Grünfelder et al., 2002), rabbit antibodies against T. brucei clathrin heavy chain (1:50), TbRAB11A (1:70), and biotin (affinity purified; 1:100; Loxo, Dossenheim, Germany). In single labeling, the primary antibodies were detected by 6 or 15 nm Protein A gold complexes (PAG-6 and PAG-15). For double labeling (Figures 3E, inset; 5, D–F), the first antibody was generally detected by PAG-10 or PAG-6; after a block with a rabbit normal IgG (1:400), the second antibody was detected by PAG-10 or PAG-15. In one case (Figure 3E, inset), clathrin was first labeled with rabbit anti-TbCLH antibodies and PAG-6, whereas VSG was subsequently detected using biotinylated rabbit anti-VSG IgG followed by a monoclonal mouse anti-biotin antibody (1:100; Vector Laboratories, Burlingame, CA) and goat anti-mouse IgG-18-nm gold conjugates (1:20; Jackson Immunoresearch Laboratories, West Grove, PA). After labeling, the cryosections were embedded in 1.4% methylcellulose, 0.3% uranyl acetate. Sections were analyzed in Philips CM10 and Philips 201 electron microscopes at 60 kV. Images were reproduced on baryte papers (final magnification, 80–to 100,000-fold), scanned at 300 dpi resolution, and digitally arranged to the plates by using the Color Photopaint program, version 10.

**RESULTS**

We previously concluded from an analysis of the steady-state distribution of VSG in the mammalian stage of T. brucei that sorting of this protein occurs in multiple intracellular steps (Grünfelder et al., 2002). From these steps, the VSG membrane concentration gradient of approximately 5 between the cell surface and an averaged estimate for endosomes seemed to be most readily accessible for further analysis. Using recently described, specific antibodies against the clathrin heavy chain and the small GTPase TbRAB11 (Morgan et al., 2001, 2002a,b; Jeffries et al., 2001), we have investigated endocytic and recycling membrane structures, which are involved in VSG trafficking from and to the flagellar pocket.

**Distribution of Clathrin and Endocytosed VSG**

An overview of the intracellular distribution of clathrin in comparison with endosomal VSG was obtained by 3D immunofluorescence analysis. The cell surface was first double-labeled in the cold with cleavable sulfo-succinimidyl-2-(biotinamido)-ethyl-1,3-dithiopropionate (sulfo-NHS-SS-biotin) and noncleavable sulfo-succinimidyl-7-aminocarbony1-4-methylcoumarin-3-acetic acid (AMCA-sulfo-NHS). After endocytosis to steady state (3 min at 37°C; unpublished data), surface-associated biotin was removed by incubation with glutathione. Although AMCA-labeled VSG was endocytosed with the same kinetics as biotinylated VSG as shown by immunoelectron microscopy by using anti-AMCA antibodies (our unpublished data), the AMCA fluorescence was quantitatively quenched within the endocytic compartment. This fortuitous observation allowed us to visualize VSG on the cell surface, including the flagellar pocket membrane, as blue AMCA fluorescence, whereas endocytosed biotinylated VSG was detected through the red fluorescence of Alexa Fluor 594 conjugated to streptavidin. In combination with immunodetection of the clathrin heavy chain and Alexa Fluor 488-conjugated secondary antibodies, we were able to reconstruct a multichannel, three-dimensional distribution map of endosomal VSG- and clathrin-containing structures. Digitally deconvolved 3D data were analyzed using (animated) maximum intensity projection (Figure 2A), volume rendering (transparency blending) (Figure 2B), and shadow projection (Figure 2C; see legend for the advantages of the different visualization techniques).

As expected, endocytosed VSG filled part of the cellular space between the flagellar pocket and the nucleus (Figure 2A, labeled FP and N). The overall VSG distribution within this area, however, did not look diffuse, but formed a coherent compartment with sharp borders. In addition, clathrin-containing structures colocalizing with endocytosed VSG were found in distinct parts of the endocytic compartment. As shown in Figure 2B and C, clathrin was not only associ-
ated with VSG-positive structures close to the flagellar pocket, but was also detectable in areas distant from the site of endocytosis. Volume-rendering (Figure 2B) and shadow projections (Figure 2C) indicated that a significant amount of clathrin was located at the periphery of the endocytic compartment. In quantitative terms, 42.7 ± 3.2% of the endocytosed VSG colocalized with clathrin and 65.8 ± 4.3% of clathrin colocalized with VSG (n = 15), implying that a significant fraction of clathrin-associated structures did not carry VSG. The fidelity of our method was validated by documenting the complete absence of a colocalization signal between 1) clathrin and AMCA-conjugated VSG located on the cell surface, and 2) clathrin and specific markers for the ER and the lysosome (our unpublished data). The further characterization of the relationship of the two proteins in membrane trafficking required the resolution of the electron microscope.

**Endocytosis of VSG by Clathrin-coated Vesicles**

For optimal ultrastructural preservation, trypanosomes were subjected to high-pressure freezing and freeze-substitution followed by Epon embedding. The electron micrographs shown in Figure 3, A and D, clearly document the VSG coat on the cell surface, on the membrane facing the flagellar pocket as well as the inner aspect of coated pits (arrowheads) and vesicles (arrows; average diameter, ~135 nm). On all of these structures, 1) the fine-structural appearance, 2) the electron density, and 3) the thickness of the coat, as determined by a quantitative evaluation of perpendicular sectioned membrane profiles, were very similar. All subsequent experiments were performed with cryosections. Figure 3E depicts a flagellar pocket from cells surface-biotinylated at 0°C.
and briefly heated to 26°C to allow endocytosis. Biotinylated VSG was clearly present in CCPs (Figure 3E, inset) and a pinched-off vesicle (CCV) but not in the exocytic carrier vesicle (EXC; see below). Therefore, these experiments prove that the extensively documented spiny coat on the cytoplasmic side of endocytic vesicles of *T. brucei* (Vickermann, 1969; Langreth and Balber, 1975; Webster and Grab, 1988) is indeed composed of clathrin as previously suggested (Webster and Shapiro, 1990; Morgan et al., 2001). Based on criteria listed above, the VSG coat included in these vesicles is indistinguishable from that at the flagellar pocket membrane. We designate these vesicles as class I clathrin-coated vesicles.

On the average, sections through the flagellar pocket region showed three to four profiles of class I coated pits and vesicles. The abundance of these structures reflects the high rate of endocytosis in this organism. Preliminary experiments suggest that the endocytosed, biotinylated VSG is first delivered to cisternal structures located near the lysosome/nucleus and then moves via endosomes back to the flagellar pocket (Engstler and Grünfelder, unpublished data).

**Clathrin-coated Vesicles Depleted in VSG Bud from Endosomes**

Endosome-associated clathrin, was first described in mammalian cells by Killish et al. (1992) and later in more detail by Stoorvogel et al. (1996). In their recent study, Morgan et al. (2001) showed by cryoimmunogold electron microscopy that clathrin immunoreactivity is widely distributed among membrane structures in the posterior region of *T. brucei* in agreement with the fluorescence microscopic analysis (Figure 2). Herein, we present the relationship between endosome-associated clathrin and VSG.

On the electron microscopic level, the endocytic compartment of *T. brucei* consists of a prominent system of narrow, sheet-like cisternae (EC, for endocytic cisterna; 30.7 ± 0.3 nm in thickness), which in cross sections, have an extended (Figure 4 A), or more rarely, circular shape (Figure 6, A and D, circular endosomal cisterna; cEC). The cisternae contain biotinylated VSG taken up by endocytosis from the cell surface (Overath et al., 1994; our unpublished data). A common feature of endosomal cisternae are coated, budding structures at their rims (Figure 4, A and B, arrows) but occasionally also on their planar face (Figure 4C, arrow), which could be labeled by anti-clathrin antibodies. Clathrin-coated vesicles (Figure 4, A–C, and F, arrowheads) distinctly smaller in size (50–60 nm in diameter) than those arising at the flagellar pocket (cf. Figure 4D) were regularly found near the endosomes. We designate these vesicles class II clathrin-coated vesicles. In addition, a clathrin coat was present at the rim (Figure 4E, arrow) or along the external face of the outermost trans-Golgi cisterna (Figure 4F, arrows), a pattern observed only on some Golgi profiles (see unlabeled Golgi complex in Figure 4B).

Remarkably, the coated vesicles of class II budding from endosomal cisternae turned out to be depleted in VSG relative to the core of the cisternae (Figure 5, A and B). This is particularly evident in the endosomal structure shown in Figure 5A, which shows multiple endosomal buds essentially free of VSG-specific immunogold label, whereas the endosomal core is strongly positive for VSG. In principle, this pattern could be an artifact, if the antigen in the buds would be buried below the section’s surface and, therefore, not accessible to antibodies and Protein A gold. Two observations argue against this possibility. First, in the very thin cryosection shown in Figure 5C, the bud was decorated by a single gold grain (demonstrating that it is accessible) but its labeling density was weak compared with the core of the cisterna. Second, the double-labeling experiments shown in Figure 5, D–F, clearly indicated that VSG was not detectable
in the clathrin-positive buds (arrows) and vesicles (arrowheads), although the cisternal core was strongly positive. We conclude that class II vesicles budding from endosomes are depleted of VSG relative to the cisternal core structure, from which they originate.

**VSG Is Recycled to the Flagellar Pocket by TbRAB11-positive Carriers**

Because VSG is rapidly taken up by endocytosis but, at the same time, is metabolically very stable, it has been suggested that >95% of the internalized protein is recycled to the cell surface (Seyfang et al., 1990). This implies that VSG/membrane endocytosis is compensated by exocytosis and, in first approximation, the rate of transport must be the same in both directions. In mammalian cells, the small GTPase Rab11 is an established marker for recycling endosomes (Ullrich et al., 1996; Sönning et al., 2000) and recently, a homolog, TbRAB11, has been characterized in *T. brucei* (Jeffries et al., 2001). Therefore, this marker offered the opportunity to investigate structures involved in recycling of VSG.

On the fluorescence microscopic level, anti-TbRAB11 antibodies showed binding to well-defined, continuous subareas of the endocytic compartment (Figure 2, D–F; Jeffries et al., 2001). In all cells analyzed, a distinct signal was obtained juxtaposed to the flagellar pocket membrane (Figure 2E, arrowhead). This is likely the site where recycling, VSG-containing, exocytic carriers (see below) fuse with the plasma membrane. Interestingly, a high percentage of TbRAB11 colocalized with VSG distant from the flagellar pocket, and we have always detected Rab11-positive structures close to the lysosome (Figure 6, A and B). Maximum intensity and shadow projections of 3D images indicated that virtually all Rab11-positive structures carry VSG (Figure 2, D and F). Volume rendering showed that a significant fraction of Rab11-positive structures was embedded

---

**Figure 4.** Association of clathrin with endosomal cisternae and the Golgi complex. Cryosections were labeled with anti-clathrin antibodies and PAG-6. Clathrin-mediated budding (A and B, arrows) is frequently observed at the rims of the EC, but sometimes also on planar regions (see arrow pointing at a bent cisterna in C). The pinched-off vesicles (A–C, E, and F, arrowheads) are smaller than CCVs derived from the flagellar pocket membrane (cf. D and Figure 3, B, C, and E). The EXCs (D) are not labeled. Although most Golgi profiles (see G in B) are not labeled, in some images a clathrin coat is present at the ends of (E, arrow) or along the external face of the outermost trans-Golgi cisterna (F, arrows). BZ, budding zone between ER and cis-Golgi; FL, flagellum; K, kinetoplast. Bars (A–F), 250 nm.
within the endocytic compartment (Figure 2E). Quantitative colocalization analysis disclosed that 27.8 ± 2.7% of all endocytosed VSG colocalized with TbRAB11, whereas 89.2 ± 6.5% of TbRAB11 colocalized with VSG (n = 17).

On the electron microscopic level, both extended and circular profiles of the cisternae could be labeled on their cytosolic face by anti-TbRAB11 antibodies (Figure 6A, arrowheads), and a quantitative evaluation of 400 randomly selected sections showed that 51.7% of all endosomal profiles were positive for this protein. Some label was also associated with the trans-Golgi cisternae (Figure 6B, arrowhead). The most strongly labeled TbRAB11-positive, intracellular structures were flat, disk-shaped vesicles, which we have named EXCs. They are sausage-shaped in cross sections (Figures 6, E and F, and 7A; 153.6 ± 5.9 nm in length, 33.7 ± 0.5 nm in thickness) but reveal their plate-like structure when sectioned in-plane (Figure 6D, asterisk; 150 ± 3.8 nm in diameter) or tangentially (Figure 7D, asterisk). Exocytic carrier vesicles were located in abundance near the flagellar pocket (Figure 6, E and F) but were also found near endosomal cisternae close to the lysosome (Figure 6A). They approach the flagellar pocket in an orientation perpendicular to the plane of the target membrane (Figure 7, A and C) and then fuse (Figure 7, B and D). Importantly, the clathrin-coated pits budding from the flagellar pocket membrane did not react with anti-TbRAB11 antibodies (Figure 7, C and D).

Together with the endocytic vesicles, exocytic carriers had the highest VSG concentration of all intracellular structures (Figure 6F) and in the steady state, they were strongly positive for surface-derived biotinylated VSG (our unpublished data). An important question was whether the VSG density (molecules/membrane area) in the exocytic carriers was as high as or even higher than on the flagellar pocket membrane/cell surface. By counting the number of gold grains per membrane length on ultrathin cryosections, we estimated that the ratio in label density between the exocytic carriers and the plasma membrane is ~0.5 rather than ~1. This ratio may be an underestimate for experimental reasons. First, on cryosections (but not on plastic sections), the cell surface was labeled along the whole section thickness (confirmed by inspection of stereo images), whereas cytoplasmic structures were only labeled at the section surface. Therefore, this edge effect can result in an artificial overestimate of VSG density at the cell surface relative to that on intracellular membranes. Second, at high antigen density, the labeling efficiency is limited by electrostatic repulsion and/or steric hindrance of Protein A gold complexes (Griffiths, 1993). Thus, it is conceivable that these effects are more pronounced in the tightly opposed VSG layers of the flat exocytic carriers than on the single-layered plasma membrane, resulting in an underestimate of antigen density in the carriers.

Figure 5. Clathrin-coated vesicles budding from endosomes are deficient in VSG. Cryosections were labeled with anti-VSG antibodies and PAG-6 (A–C) or double labeled with anti-clathrin and anti-VSG antibodies and PAG-6, PAG-10, or PAG-15 as indicated (D–F). (A) An EC with numerous clathrin-coated buds has been sectioned perpendicularly in the upper part and in plane in the lower part. Whereas the lumen of the cisterna is strongly labeled for VSG, the buds are nearly VSG free. (B) The buds (arrows) as well as an adjacent region of the cisterna (arrowheads) are clathrin-coated but free of VSG. The latter region may indicate an incipient clathrin-coated bud. (C) A very thin cryosection with a single gold grain inside the terminal bud (arrow) at the rim of the cisterna. (D–F) Budding structures on cisternae marked by arrows are positive for clathrin and negative for VSG, whereas the reverse pattern is observed for their sheet-like core region. Adjacent, free clathrin-coated vesicles (E and F, arrowheads) are also free of VSG. SC, surface coat. Bars (A, B, E, and F), 125 nm; 250 nm (C); and 500 nm (D).
DISCUSSION

Clathrin-mediated Budding in T. brucei Occurs at the Flagellar Pocket, on Endosomes and a trans-Golgi Cisterna

The variant surface glycoprotein enters the cells through class I clathrin-coated pits and vesicles. For several reasons, we believe that this is the dominant and most likely the only endocytic pathway in T. brucei. First, because the flagellar pocket membrane is limited in area (3–7% of the total cell surface; Grünfelder et al., 2002) and well defined with its balloon-like shape, particularly in high-pressure-frozen cells, it is unlikely that alternative endocytic invaginations would have gone unnoticed, considering the large number of ultrastructural studies on trypanosomes performed in this and other laboratories. Second, class I clathrin-coated pits and vesicles at and near the flagellar pocket are found in abundance, and this correlates with the distinct, overlapping VSG/clathrin-containing region close to the flagellar pocket in fluorescence microscopic images (Figure 2, A–C). Therefore, these vesicles are formed at a high frequency, consistent with the rapid rate of receptor-mediated and fluid phase endocytosis by the parasite (Coppens et al., 1987). Third, if surface-biotinylated cells are allowed to endocytose for short times, intracellular VSG<sub>biotin</sub> is readily and only observed in this class of coated vesicles, strongly suggesting that VSG enters the cell solely by this pathway. The further investigation of this route, i.e., by a biochemical characterization of isolated coated vesicles (Shapiro and Webster, 1989), is likely of considerable interest considering that their composition may be unusual; T. brucei apparently lacks a homolog for the adaptin, AP2, the heterotetrameric complex associated with clathrin-mediated endocytosis in mammalian cells (Morgan et al., 2002b).

As judged by immunofluorescence (Figure 2, A–C), most of the cellular clathrin is located at a distance from the flagellar pocket in the direction of the nucleus. The majority of this clathrin colocalizes with biotinylated VSG taken up to

Figure 6. TbRAB11 marks endosomal cisternae and exocytic carrier vesicles rich in VSG. Cryosections were labeled with anti-TbRAB11 (A–E) or anti-VSG antibodies (F) and PAG-6. (A) Extended or circular profiles of endosomal cisternae (EC and cEC, respectively) near the lysosome (L). Some cisternae (arrowheads) and EXCs are labeled by anti-TbRAB11 antibodies. (B) The TbRAB11 protein is localized on the cytoplasmic side of endosomal membranes and occasionally, on cisternae of the Golgi complex (G, arrowhead). (C and D) TbRAB11-positive endosomes with several budding clathrin-coated pits (arrows). The asterisk marks an in-plane cross-sectioned EXC. (E) In contrast to the cisternae, the exocytic carrier vesicles are uniformly and strongly labeled for TbRAB11. (F) Together with the endocytic vesicles, the exocytic carriers have the highest VSG-concentration of all intracellular structures (compare the label density of the exocytic carriers [higher magnification in the inset] with that of perpendicularly sectioned regions of the VSG surface coat [SC]). Bars (A–F), 250 nm; inset (F), 125 nm.
the steady state by endocytosis; however, there is no overlap with VSG for approximately one-third of the clathrin. This distribution is consistent with the immunoelectron microscopic results. Cisternal endosomes positive for VSG in their core region display clathrin-coated budding structures designated class II, which are essentially free of VSG (Figures 4 and 5). Moreover, in the vicinity of the cisternae, class II clathrin-coated vesicles are found in abundance, which are also deficient in VSG and are, therefore, most likely derived from these cisternae. The target membrane for the class II coated vesicles remains unknown (compare legend to Figure 1).

In mammalian cells, abundant endosomal clathrin-coated buds and derived vesicles with a diameter of 60 nm, i.e., similar in size to those observed herein, and generally negative for the adaptor complex AP2, were described by Stoorvogel et al. (1996); the authors considered that these coated vesicles are involved in recycling from early endosomes to the plasma membrane or to the trans-Golgi network (TGN). In contrast, Futter et al. (1998) showed that coated buds on endosomal tubules contained a subunit characteristic for the adaptor complex AP1; these buds were proposed to be responsible for concentrating the transferrin receptor being packaged for basolateral sorting in MDCK cells. A recent study in HeLa cells by van Dam and Stoorvogel (2002) suggests that the pathway from recycling endosomes to the plasma membrane is mediated, at least in part, by endosome-derived clathrin-coated vesicles. Thus, the target membrane for these vesicles likewise remains to be defined.

There seems to be a third clathrin-dependent budding process in *T. brucei* occurring at the outermost trans-Golgi cisterna (Figure 4). Because tubular-vesicular structures defining the TGN in mammalian cells have not been observed in this organism, this cisterna may be the TGN equivalent (Field et al., 2000; McConville et al., 2002a), and, like in mammalian cells, clathrin may be involved in the sorting of lysosomal proteins (Alexander et al., 2002).

![Figure 7](image)

**RAB11-positive Carriers Mediate Transport of VSG from Endosomes to the Flagellar Pocket**

About half of all endosomal cisternae can be designated recycling endosomes because they contain the small GTPase TbRAB11. Some of these cisternae may be interconnected (Webster, 1989) and by analogy to mammalian cells (Sönntighsen et al., 2000), they may be organized as a mosaic of domains, which recruit different RAB proteins to subregions of the same structure. We propose that these cisternae are the source of the characteristic disk-like vesicles designated exocytic carriers, which 1) are strongly positive for TbRAB11 and VSG, 2) have a similar thickness and structural appearance as the cisternae in cross section, and 3) are generally observed in the vicinity of cisternae. Because of their abundance particularly close to the flagellar pocket, we consider these carriers as the dominant and likely the only vesicular structures that transport membrane from the endosomal compartment to the flagellar pocket. This implies that the flow of VSG/membrane entering the cell via class I clathrin-coated vesicles is balanced by a reverse flow by way of TbRAB11-positive exocytic carriers. Kinetic experiments on the rates of VSG-endocytosis and recycling support this view (unpublished data). They show that clathrin-containing structures become VSGbiotin-positive before the TbRAB11-carrying structures (Figure 3E) arguing against an endocytic function of the TbRAB11-positive carrier vesicles. To balance the endocytic membrane flow via class I CCVs, the EXCs must be assumed to fuse with the flagellar pocket rather than deliver the VSG by a kiss and run mechanism.
This view is strongly supported by the phenotype of cells where clathrin-function has been abolished using the RNA interference technology. There, the size of the flagellar pocket increases dramatically, which is consistent with an inhibition of inward but unhindered outward membrane flow (Engstler, unpublished data).

**Trafficking and Sorting of VSG**

The VSG at the surface coat has a purity of ≥95%, and the density of this protein at the surface is 50-fold higher than on the membrane of the ER. Therefore, VSG is very efficiently separated from other proteins as well as laterally concentrated. This concentration gradient is achieved in two (ER → Golgi complex → flagellar pocket membrane) or three steps (ER → Golgi complex → endosomes → flagellar pocket membrane). If exocytosis exclusively occurs via TbrAB11-positive exocytic carriers as suggested above, then newly synthesized VSG may be routed directly or indirectly from the Golgi complex to TbrAB11-positive recycling endosomes and carriers to the flagellar pocket. VSGs endocytosed at the flagellar pocket reaches these recycling endosomes probably via TbrAB5-positive early endosomes, which seem to be preferentially located close to the nucleus rather than in the direct vicinity of the flagellar pocket (Pal et al., 2002; Engstler and Grünfelder, unpublished data).

The multiple steps at which VSG is sorted and concentrated may involve both positive and negative selection mechanisms. Based on the observations presented in this article, we can only comment on two of these steps. First, we suggest that inclusion of the VSG coat in class I clathrin-coated pits and vesicles is essentially a passive process, which involves no gross change in VSG concentration. However, without an accurate quantification in terms of molecules/membrane area, it remains possible that the VSG concentration in the vesicles (e.g., due to steric hindrance) is in fact somewhat lower than at the flagellar pocket membrane. Second, we provide a rather strong argument for sorting of VSG by default in the recycling endosomes. The class II clathrin-coated vesicles budding from VSG-positive endosomal cisternae are strongly depleted in VSG, implying that VSG is concentrated by negative selection in the cisternae (Figure 5). We previously noted (Grünfelder et al., 2002) that the membrane concentration of VSG in cisternal structures increases two- to fourfold in a direction from the nucleus to the flagellar pocket and that the concentration is particularly high in circular cisternal profiles located close to the flagellar pocket. These observations are consistent with a gradual increase in the lateral VSG density as more and more membrane depleted in VSG is removed. As expected, the VSG concentration in the exocytic carriers is high and approaches the surface density, but for the technical reasons discussed above, may be underestimated relative to the cell surface.

Our observations can be compared with studies on trafficking and sorting in mammalian cells and yeast. As pointed out by Sabharanjak et al. (2002), nonselective uptake, i.e., without significant changes in membrane density, is the most parsimonious mechanism for GPI-anchored protein internalization by various routes in mammalian cells, given their uniform, diffuse distribution at the cell surface. There is now agreement that GPI-anchored proteins are not enriched in clathrin-coated pits/vesicles or in caveoli relative to the plasma membrane (reviewed in Chatterjee and Mayor, 2001). However, a recently described cdc42-regulated, clathrin- and caveolin-independent pinocytic pathway in COS and Chinese hamster ovary cells has been suggested to recruit GPI-linked proteins from the cell surface into a specific peripheral tubular-vesicular, early endosomal compartment, where they seem to be (secondarily?) enriched. Thereafter, they are delivered to pericentriolar recycling endosomes, where they are retarded by a sphingolipid- and cholesterol-dependent mechanism relative to bulk membrane constituent before being routed back to the plasma membrane by so far undefined carriers (Mayor et al., 1998; Chatterjee et al., 2001; Sabharanjak et al., 2002). These authors invoke sphingolipid- and cholesterol-rich lipid microdomains to explain both the selective uptake of GPI-anchored proteins into the specific endocytic compartment as well as their transient retention in recycling endosomes. In a recent article, Fivaz et al. (2002) concluded that the differential sorting and fate of GPI-anchored proteins in endosomes of baby hamster kidney and Chinese hamster ovary cells depends on their residence time in lipid microdomains. In the Golgi/trans-Golgi network of polarized Madin-Darby canine kidney and nonpolarized PtK2 cells, lateral segregation and subsequent incorporation into distinct transport carriers has been shown for a trans-membrane and a GPI-anchored cargo (Keller et al., 2001). Finally, sorting of GPI-anchored proteins into defined carriers has been demonstrated upon exit of the ER in yeast (Muñiz et al., 2001). Together, these studies suggest that sorting of GPI-linked proteins can occur in endosomes as well as at several stages in the biosynthetic pathway.

**Is Cellular Sorting the Key to GPI-Anchor Function?**

The question why there is a glycolipid anchor as a mode for attaching proteins to membranes has puzzled investigators for two decades (Ferguson, 1999; Chatterjee and Mayor, 2001). The “raft” hypothesis (Simons and Ikonen, 1997) suggests a function in cellular sorting by postulating that GPI-anchored proteins have a tendency to associate with highly dynamic liquid-ordered lipid domains formed in the liquid-disordered lipid bulk phase of a membrane. As summarized in a recent article by Anderson and Jacobson (2002), no consensus about the size, shape, and location of these domains at the surface of mammalian cells has emerged. In its second part, the hypothesis proposes that such domains are specifically incorporated into transport carriers formed, for example, at the trans-Golgi network for selective delivery of GPI-anchored proteins to the cell surface. The difficulty with this proposition is that it implies a sorting mechanism, which specifically identifies GPI-linked proteins and/or lipid-ordered domains from the opposite, cytosolic face of a membrane and ensures their preferential inclusion into vesicular carriers. Based on the observations on endosomal sorting reported in this study, we would like to suggest an alternative scenario. GPI-anchored proteins can be specifically excluded from structures budding from a given intracellular membrane, thereby being sorted and laterally concentrated by negative selection. This may still involve association with fluid-ordered lipid domains; however, these domains would not be included into budding structures. The excluded regions enriched in the GPI-anchored and, likely, other components such as sphingolipids or mul-
Sorting of GPI-anchored Proteins

tipass trans-membrane proteins such as transporters would then give rise to carriers such as the exocytic vesicles described in this study. In conclusion, the essence of the glycolipid anchor function would be that it is neglected by coats, which are involved in positive sorting of trans-membrane proteins. In trypanosomes, multiple negative sorting steps may operate to achieve the high VSG concentration at the cell surface, which is vital for survival in the mammalian vascular system providing a highly dynamic shield against antibodies toward buried invariant surface proteins as well as against attack by complement.

ACKNOWLEDGMENTS

We thank Sabine Geier for expert technical assistance and Drs. Michael Boshart (Munich, Germany), Anne Spang, and Suzanne Gokool (both from Tübingen, Germany) for reading the manuscript and helpful discussions. M.E. thanks Michael Boshart for continuous support in the laboratory. P.O. acknowledges funding by the Fond der Chemischen Industrie (Frankfurt, Germany), and M.B. by the Deutsche Forschungsgemeinschaft. G.W.M. and M.C.F. are supported by the Wellcome Trust (London, United Kingdom).

REFERENCES


