Receptor-mediated Endocytosis of Transferrin and Recycling of the Transferrin Receptor in Rat Reticulocytes

CLIFFORD HARDING, JOHN HEUSER, and PHILIP STAHL
Department of Physiology and Biophysics, Washington University School of Medicine, St. Louis, Missouri 63110

ABSTRACT At 4°C transferrin bound to receptors on the reticulocyte plasma membrane, and at 37°C receptor-mediated endocytosis of transferrin occurred. Uptake at 37°C exceeded binding at 4°C by 2.5-fold and saturated after 20–30 min. During uptake at 37°C, bound transferrin was internalized into a trypsin-resistant space. Trypsinization at 4°C destroyed surface receptors, but with subsequent incubation at 37°C, surface receptors rapidly appeared (albeit in reduced numbers), and uptake occurred at a decreased level. After endocytosis, transferrin was released, apparently intact, into the extracellular space. At 37°C colloidal gold-transferrin (AuTf) clustered in coated pits and then appeared inside various intracellular membrane-bounded compartments. Small vesicles and tubules were labeled after short (5–10 min) incubations at 37°C. Larget multivesicular endosomes became heavily labeled after longer (20–35 min) incubations. Multivesicular endosomes apparently fused with the plasma membrane and released their contents by exocytosis. None of these organelles appeared to be lysosomal in nature, and 98% of intracellular AuTf was localized in acid phosphatase-negative compartments. AuTf, like transferrin, was released with subsequent incubation at 37°C. Freeze-dried and freeze-fractured reticulocytes confirmed the distribution of AuTf in reticulocytes and revealed the presence of clathrin-coated patches amidst the spectrin coating the inner surface of the plasma membrane. These data suggest that transferrin is internalized via coated pits and vesicles and demonstrate that transferrin and its receptor are recycled back to the plasma membrane after endocytosis.

Receptor-mediated binding and endocytosis of transferrin occur in many cell types (4, 17, 23, 24, 34, 38) and appear to be requisite steps in iron delivery under some conditions (7, 9). Transferrin uptake has been best studied in erythropoietic cells, where the synthesis of hemoglobin requires a large amount of iron. Expression of transferrin receptors in these cells peaks early in development and declines progressively during the maturation of erythroblasts and reticulocytes (23, 34, 36). Despite their lower level of receptor expression, reticulocytes are a convenient model system, since they may be easily isolated from the blood of anemic animals.

Transferrin binding is mediated by protease-sensitive receptors (8) but is not inhibited by glycosidase treatment of either transferrin or its receptor (8, 18, 22). Transferrin receptors have been identified and characterized by many research groups. The uptake of transferrin is both temperature and energy dependent, and transferrin endocytosis has been demonstrated by the use of EM-autoradiography (9, 21), ferritin- or horseradish peroxidase-conjugated transferrin (9, 31), and ferritin-conjugated antitransferrin antibodies (31). These techniques reveal that transferrin binds to the plasma membrane at 4°C and appears inside intracellular vesicles during incubation at 37°C. Receptor-mediated transferrin uptake may occur by mechanisms similar to the uptake of other ligands: namely, clustering in coated pits and endocytosis via coated vesicles.

In this paper we present further biochemical data concerning transferrin endocytosis and provide evidence for the recycling of the transferrin receptor. Additionally, we report the use of various morphologic techniques to examine the mech-
anism of endocytosis in these cells, and we further define the intracellular compartments through which internalized transferrin passes.

MATERIALS AND METHODS

Ligand: Transferrin from pooled human serum (Calbiochem-Behring Corp., San Diego, CA or Sigma Chemical Co., St. Louis, MO) was tested for purity by SDS PAGE. Differferin transferrin was used in all experiments and was produced by the method of Galbraith et al. (5). Inductions were performed by the chloramine T method as modified by Stahl et al. (30), except that 200 μg (20 μl) of chloroamine T was added to 200 μg of transferrin in a total volume of 130 μl. 15-nm colloidal gold beads were made by the citrate method (6).

The samples were placed on the method described by Heuser (10, 12). Briefly, the samples were placed on treated polylysine-coated coverslips (slips were coated by incubation in 2 mg/mL polylysine in buffer at 37°C). Fixed samples were prepared without freezing and stained for acid phosphatase (using cytidine monophosphate as substrate) (1) or for arylsulfatase B (using p-nitrophenyl sulfate) (2). For examination of surface-bound AuTf, fixed cells were adsorbed to small (5-mm square) polystyrene-coated coverslips (slips were coated by immersion in 2 mg/mL polylsine in distilled H2O for 45 min, followed by extensive washing in distilled H2O, rinsed in 15% MeOH in distilled H2O, and immediately quick-frozen. To examine the inside of the plasma membrane, reticulocytes were adsorbed to polylsine-coated coverslips, rinsed in Ca2+-free Ringer’s, scraped open in KHME (70 mM KCl, 5 mM MgCl2, 3 mM EGTA, 30 mM HEPES, pH 7.0) using a platinum wire, transferred to fixatives (115 mM KCl, 5 mM MgCl2, 3 mM EGTA, 30 mM HEPES, pH 7.0, 1% glutaraldehyde and 2% paraformaldehyde), washed in buffer several times, and embedded, and thin sectioned without freezing gave similar results. Silver replicas were cut on a Porter-Blum MT-2 ultramicrotome (Porter Instrument Co., Inc., Hatfield, PA) and stained for 20 min with 5% uranyl acetate in 50% MeOH, 50% distilled H2O, and for 2-5 min in 0.4% lead citrate in 0.15 N NaOH.

RESULTS

Biochemical Approaches to Transferrin Endocytosis

Reticulocytes were incubated in the presence of 125I-transferrin plus or minus excess unlabeled transferrin. At 4°C transferrin binding was found to be specific, saturable, and reversible. Binding was found to be essentially complete within 60 min; thereafter, an incubation time of 90 min was routinely used. Saturation of binding occurred at ~1.8 μg/ml (7.5 × 10^-8 M), and a Kd of 2 x 10^-8 M was estimated from a Scatchard plot. Extrapolation indicated that ~ 4.9 × 10^4 receptors were expressed on the surface of these cells. Disso- ciation at 4°C in the presence of 150 μg/ml unlabeled transferrin was found to occur with a t1/2 of 80 min.

At 37°C saturation of ligand uptake occurred within 25-35 min. After a 35-min uptake cells were incubated in the presence of unlabeled transferrin (2.0 mg/ml) at 37°C. Release of cell-bound 125I-transferrin occurred with a t1/2 of 8 min. The radioactivity released into the medium during the second incubation was 90% photostungastic acid-precipitable. This indicated that 125I-transferrin uptake at 37°C was reversible and that under these conditions virtually all of the bound ligand was released into the medium in an essentially intact form. At 37°C saturation of 125I-transferrin uptake occurred at ~ 1 μg/ml (8.8 × 10^-8 M). A Scatchard plot indicated a Kd of 5.1 x 10^-8 M and 1.2 x 10^-4 receptors/cell. Comparison with the previous Scatchard data showed that at 4°C-binding labeled 40% of the total receptors labeled at 37°C, indicating that ~ 60% of the receptors were inaccessible and may be intracellular. The requirement of an equilibrium state for Scatchard analysis appeared to be met by the fact that both uptake to a steady state and release of bound ligand were virtually complete within 20-30 min. The number of receptors expressed per cell varied considerably (generally 1-3 x 10^5/cell) between experiments, as would be expected with a receptor number dependent on developmental stage, but the 4°C binding level was generally 35-40% of the 37°C uptake in the same cell population. No specific transferrin binding or uptake was observed with mature erythrocytes.

To investigate the distribution of ligand after 37°C uptake, trypsinization at 4°C was employed to distinguish cell surface
ligand from intracellular ligand. Following 25 min of incubation at 37°C, ~ 20-40% of the bound ligand was found to be trypsin resistant, indicating that internalization had occurred. (Some variability between experiments may have been caused by a varying amount of loss of intracellular ligand due to cell lysis during the trypsinization.) While these results, similar to those of Octave et al. (24), indicate that a minimum of 20-40% of the ligand bound at 37°C was intracellular, the actual internal fraction probably exceeded this value. The trypsin-resistant fraction was found to increase with time during 37°C incubation until a plateau was reached after 25-35 min. Either internalization of receptor-ligand complexes ceased at this time, or, more likely, a steady-state equilibrium was achieved where internalization was balanced by externalization of ligand or receptor-ligand complexes.

To determine whether receptors could be externalized, cells were pretrypsinized at 4°C, washed, and then incubated with 125I-transferrin at 37°C. Trypsinization at 4°C destroyed at least 94% of the original surface receptor pool (Fig. 2). Nonetheless, the subsequent uptake occurred with a similar time course but saturated at ~ 40% of uptake by untrypsinized cells of the same population. This indicated that trypsin-resistant receptors, presumably intracellular, replaced destroyed surface receptors to mediate transferrin uptake. Furthermore, uptake after 4°C-trypsinization was not significantly affected by the presence of cycloheximide (50 μg/ml), indicating that this uptake was not dependent on the synthesis of new receptors.

The existence of distinct internal and surface receptor pools that communicate only above 4°C was further demonstrated. Cells were trypsinized at 4°C, washed to remove trypsin, incubated without transferrin at 37°C, and cooled to 4°C; binding of 125I-transferrin was then measured at 4°C (Fig. 1). Again, 4°C trypsinization destroyed the original pool of surface receptors, but subsequent incubation at 37°C allowed recovery of surface receptors to ~ 45-70% of pretrypsinization levels. These results confirm the existence of a trypsin-resistant (internal) pool of receptors that can replenish surface receptors destroyed by trypsin.

**Morphologic Approaches to Transferrin Endocytosis**

A number of techniques were used to visualize the process of transferrin endocytosis. AuTF was used as a morphologic probe for the transferrin receptor and the pathway of transferrin endocytosis. The specificity of AuTF binding was demonstrated by two methods. First, 125I-transferrin was conjugated to gold beads and its uptake was assayed as described above for unconjugated iodinated ligand. Fig. 3 demonstrates that uptake of 125I-labeled AuTF was inhibited by excess unlabeled, unconjugated transferrin (1 mg/ml final concentration). Inset: specific uptake as a function of the amount of colloid added to the cells. In addition, 100 μl of the supernatant from a second colloid centrifugation was similarly incubated with reticulocytes; the uptake observed in the absence (−) and presence (+) of unlabeled, unconjugated transferrin is indicated by the stippled bars on the right.

![Figure 1](https://example.com/figure1.png)  
**Figure 1** Uptake of 125I-transferrin by rat reticulocytes after the destruction of surface receptors by 4°C trypsinization. Control reticulocytes were incubated with 125I-transferrin (20 μg/ml) in Ringer’s at 37°C for various periods. Other cells were trypsinized at 4°C, washed, and then incubated at 37°C as above. Specific uptakes of control cells (open circles) and pretrypsinized cells (solid circles) are shown as a function of incubation time.

![Figure 2](https://example.com/figure2.png)  
**Figure 2** Recovery of 4°C binding during 37°C incubation following the destruction of surface receptors by 4°C trypsinization. Reticulocytes were trypsinized at 4°C, washed, incubated in Ringer’s at 37°C for various periods, and cooled to 4°C. The cells were then incubated with 125I-transferrin (6 μg/ml) for 90 min at 4°C. Specific binding is shown as a function of time of incubation at 37°C.

![Figure 3](https://example.com/figure3.png)  
**Figure 3** Specific uptake of 125I-transferrin conjugated to colloidal gold by reticulocytes at 37°C. 125I-transferrin-conjugated gold colloid was prepared, centrifuged, and suspended in Ringer’s. Aliquots (100, 50, and 25 μl) of this suspension were incubated with reticulocytes for 30 min at 37°C in a final volume of 200 μl. The first three pairs of bars show uptake in the absence (−) and presence (+) of unlabeled, unconjugated transferrin (1 mg/ml final concentration). Inset: specific uptake as a function of the amount of colloid added to the cells. In addition, 100 μl of the supernatant from a second colloid centrifugation was similarly incubated with reticulocytes; the uptake observed in the absence (−) and presence (+) of unlabeled, unconjugated transferrin is indicated by the stippled bars on the right.
and a 35-min chase at 37°C with cold, unconjugated transferrin, the cells were removed by centrifugation (1,000 g for 5 min); 71% of the phosphotungstic acid-precipitable radioactivity remaining in the supernatant was pelleted by centrifugation at 20,000 g for 75 min. This suggests that most of the released 125I-transferrin was still associated with gold beads and/or with membrane-bound vesicles released by multivesicular endosome (MVE) exocytosis (see below).

AuTf binding to the plasma membrane was demonstrated by incubating reticulocytes with AuTf at 37°C, washing with Ringer's at 4°C, and fixing with fixative, at 4°C. Fixed cells were then adsorbed onto polylysine-coated cover slips, quick-frozen, freeze-dried, and rotary-replicated. The resulting replicas retained surface-bound gold beads, strikingly illustrating the distribution of transferrin receptors on the plasma membrane. AuTf was found scattered randomly on the cell surface and clustered in membrane pits (Figs. 4 and 5). To study the distribution of intracellular AuTf, incubations were similarly performed, but cells were quick-frozen in a thick suspension. Some cells were quick-frozen without prior fixation. These samples were then freeze-substituted, stained, and embedded for thin sectioning. Fig. 7 shows thin sections of reticulocytes incubated with AuTf for 5 min at 37°C. AuTf is localized to the free surface of the plasma membrane, within coated pits and vesicles (~100 nm o. d.), inside small uncoated vesicles of varying size (<200 nm; generally ~100 nm diam), and inside tubular structures (usually 60–80 nm diam) (Fig. 8). Some vesicular profiles may represent cross sections of tubular compartments. The clathrin coat is sometimes difficult to discern in thin sections, since the reticulocyte cytoplasm stains so densely that it can mask the presence of a densely staining coat on the cytoplasmic surface of membrane structures.

When reticulocytes were incubated for 20 min or longer at 37°C, AuTf was also found within MVE (see Fig. 9). These structures are often 250–300 nm diam, but range from 120 to 800 nm diam and may be irregularly shaped. Included in this class are membrane-bound structures that contain distinct vesicular inclusions as well as others that contain more amorphous inclusions that stain with a density similar to that of the cytoplasm. In some cases vesicular inclusions may represent cytoplasmic protrusions into the lumen of the MVE; such protrusions were occasionally visualized. MVE were often heavily labeled, with one structure containing many gold beads. The beads frequently appeared to be associated with the membranes and inclusions of the MVE.

The enzyme content of MVE was explored to determine whether or not these structures contained lysosomal enzymes. Acid phosphatase staining was performed using cytidine monophosphate as the substrate (1). Figs. 10 and 11 demonstrate that this procedure stained reticulocyte lysosomes but did not stain MVE. Arylsulfatase staining also failed to stain MVE. Thus, MVE differ from some other multivesicular bodies, which often contain acid phosphatase and other lysosomal enzymes. In fact, 98% of intracellular AuTf was observed in acid phosphatase-negative compartments. Most lysosomal structures did not contain AuTf, but occasionally these structures did contain a small number of beads. These data indicated that transferrin processing occurred in nonlysosomal compartments.

Table I shows the time course of distribution of AuTf among various compartments. In our analysis, coated vesicles, small uncoated vesicles, and tubular structures represent one kinetic class; these structures all contained AuTf after 5 min of uptake at 37°C. (Differences in the rate of labeling between these compartments may exist but were not discernible by our methods.) MVE, on the other hand, were labeled only rarely in 5–10-min incubations, but by 35 min they contained 17% of the cell-bound AuTf. MVE were observed in reticulocytes after all periods of incubation at 37°C, and counts indicated that their abundance did not vary significantly between 5 and 35 min of incubation with AuTf at 37°C. Therefore, the increased percentage of beads residing in MVE was due to an increase in the density of MVE labeling rather than an increase in MVE number. For example, MVE in reticulocytes incubated with AuTf for 5–10 min at 37°C were sparsely labeled (0–3 beads/MVE), whereas the longer incubations produced much denser labeling (sometimes exceeding 70 beads in one MVE). Even after 35-min incubations, labeled MVE were much less common than other labeled compartments, but they contained a disproportionately large fraction of the beads owing to their heavy labeling.

To examine the recycling of AuTf, reticulocytes were incubated with AuTf for 35 min at 37°C, washed in 4°C Ringer's, and incubated in the presence of 2 mg/ml unconjugated transferrin for 25 min at 37°C. These cells were examined in thin sections, and cell-bound AuTf was quantitated to determine the fate of endocytosed AuTf. After 25 min of chase, the average number of beads bound per cell section decreased to 22% of the level found in control (unchased) samples, indicating that endocytosed AuTf was released from reticulocytes upon subsequent incubation. The cellular distribution of AuTf also changed (Table I). Beads were virtually non-existent inside coated pits, presumably owing to competition by unconjugated transferrin, and the chase also decreased the frequency of AuTf-labelling of small vesicles and tubules. Although the absolute level (beads per organelle per cell section) of MVE labeling was also decreased, the fraction of cell-bound beads inside MVE rose to 38%. This indicated that MVE represent a late stage in transferrin processing in the reticulocyte.

An additional observation in this experiment was the apparent exocytosis of AuTf-labeled MVE with the consequent release of the vesicular inclusions into the extracellular space (Figs. 12–17). Often vesicles similar to those found in MVE were found attached to the plasma membrane (Figs. 15 and 17). Furthermore, 54% of the AuTf bound on the free surface of the cells after the chase was associated with adherent vesicular inclusions, and another 21% occurred in clusters that looked as though they might have been the remains of a MVE exocytic event. These results suggested that at least some of the transferrin that accumulated inside MVE was eventually released by exocytosis.

The distribution of AuTf in reticulocytes described above has been confirmed by several techniques. After incubation with AuTf, reticulocytes were quick-frozen without prior fixation to control for fixation artifacts. The results were identical to those obtained with fixed cells, including the accumulation of AuTf within MVE and the apparent exocytosis of AuTf-labeled MVE. Freeze-fracture studies confirmed the intracellular distribution of AuTf determined in thin sections (Figs. 18 and 19). In intracellular AuTf was not visualized in this type of preparation unless the samples were OsO4-fixed before freezing. Otherwise, the fracture plane followed the membrane of intracellular vesicles and did not expose their lumen. In OsO4-fixed cells that were freeze-fractured and deep-etched,
AuTf remained in the replicas, appearing within cross-fractured intracellular vesicles. This demonstrates that gold beads may be used in freeze-fracture studies to explore the distribution of both intracellular and extracellular ligands. Importantly, this provided a control for the differential loss of gold beads from organelles in thin sections. Fig. 19 shows an MVE.
with its vesicular inclusions which was labeled with AuTf during a 20-min incubation at 37°C. AuTf appears to be bound to the inner walls and vesicular inclusions of these vesicles. When viewed in stereo, intracellular vesicles appeared as bowl-like depressions in a mesa of etch-resistant cytoplasm; they etched more than the surrounding cytoplasm because they contained much less protein.

During a 20-min incubation at 37°C, AuTf appears to be bound to the inner walls and vesicular inclusions of these vesicles. When viewed in stereo, intracellular vesicles appeared as bowl-like depressions in a mesa of etch-resistant cytoplasm; they etched more than the surrounding cytoplasm because they contained much less protein.

Due to the densely-staining nature of reticulocyte cytoplasm and other inherent limitations of the technique, thin sections do not provide good views of coated pit and vesicle formation in reticulocytes. More information concerning clathrin-coated structures can be obtained from platinum-carbon replicas (11). To further explore the morphologic basis of endocytosis, we used techniques developed earlier (10) to permeabilize or scrape open cells to remove the soluble cytoplasmic proteins. Following these treatments the samples were quick-frozen, freeze-fractured, deep-etched, and rotary-replicated. Triton extraction of reticulocytes largely removed the plasma membrane, leaving behind an erythrocyte-like cytoskeleton (devoid of microfilaments, microtubules, etc.). Clathrin-coated structures were observed within these samples. Saponin extraction left the plasma membrane more nearly intact but permeabilized the cell and released enough cytoplasmic proteins to reveal coated pits and vesicles. Soluble cytoplasmic proteins were also released by hypotonic lysis or were diluted by swelling the cells with hypotonic solutions; both treatments exposed coated pits on the plasma membrane. Finally, reticulocytes attached to polylysine-coated glass slips were scraped open, fixed, quick-frozen, and freeze-dried to reveal the proteins applied to the inner surface of the plasma membrane. Figs. 20 and 21 demonstrate the results of this approach, revealing coated structures surrounded by a background of fixed cytoskeletal proteins. Coated regions are usually 60-180 nm (mostly 80-140 nm) in diameter, slightly domed structures, viewed from this perspective. The size and conformation of coated pits may be restricted in this preparation, since the formation of coated pits may be altered by adherence to the cationic polylysine substrate. There is further variation in the size of clathrin patches, including very small clathrin aggregates. However, the large amounts of clathrin revealed by similar techniques in fibroblasts (11), hepatocytes (13), macrophages, lymphocytes, and bone marrow cells were not observed in reticulocytes. This may reflect a decreasing ability of maturing reticulocytes to perform receptor-mediated endocytosis.

**DISCUSSION**

The endocytosis of transferrin provides many interesting contrasts to other endocytic systems, including those of the mammalian mannose and glucose receptors. Unlike these systems, transferrin binding is apparently not oligosaccharide mediated, since glycosidase treatment of transferrin or its receptor does not reduce transferrin binding (8, 18, 22). The mannose receptor of macrophages (30) and the galactose receptor of hepatocytes (16, 37, 41) both target their ligands largely to lysosomes, where they are degraded. These ligands dissociate from their receptors, probably in a prelysosomal acid compartment, allowing the receptors to separately recycle back to the cell surface (32, 33). Transferrin, however, is not degraded and is released intact from cells after endocytosis, indicating that it may largely avoid lysosomes during its intracellular passage. This is substantiated by our morphologic results, which indicated that 98% of intracellular AuTf resides in acid phosphatase-negative compartments. Furthermore, transferrin may remain largely associated with its receptor inside the cell. At 37°C, 125I-transferrin uptake saturated and appeared to satisfy the equilibrium requirements for Scatchard analysis, unlike ligands that dissociate from their receptors and accumulate within the cell. In some systems lysosomotropic amines interfere with receptor recycling and may inhibit receptor-ligand dissociation (32). In contrast, 125I-transferrin uptake is relatively unaffected by these agents, while removal of the iron from transferrin is inhibited (20, 25, 26, 6a). Thus, intracellular transferrin-receptor complexes may enter a low pH environment, which is necessary for iron removal but does not affect the transferrin-receptor interaction (6a). The close association of AuTf with membrane surfaces in both thin-section and freeze-fracture studies also suggests a continued receptor-ligand interaction, although this association could be explained as nonspecific or as a fixation artifact. When reticulocytes were chased by exposure to free transferrin after incubation with AuTf, intracellular ligand was released. Small clusters of AuTf that remained attached to the cell surface after 35 min of chase may represent exocytosed receptor-ligand complexes. This suggests that, for transferrin, the receptor-ligand interaction remains intact.

![Figures 7-11](https://www.jcb.org/content/105/3/335/F10.large.jpg)
throughout endocytosis, leading to recycling of both receptor and ligand to the surface, in contrast to other receptor systems where receptor-ligand dissociation allows targeting of ligand to lysosomes and recycling of the unoccupied receptor. The fidelity of the transferrin-receptor interaction is fortunate, since this makes the ligand an unusually useful morphologic probe for its receptor.

The necessity of endocytosis for the removal of iron from transferrin continues to be questioned by some groups (6, 19, 27, 39, 40). Nonetheless, transferrin endocytosis has been well documented (9, 21, 31) and is confirmed by our results. Our morphologic data revealed that after 5–10 min of incubation at 37°C, most intracellular AuTf appeared inside small vesicular and tubular profiles. After 20–35 min of incubation, an increasing fraction appeared inside heavily labeled MVE. Transferrin-labeled structures of all types were mostly acid-phosphatase negative. MVE displayed a relatively slow time course of AuTf labeling, and these structures may represent a late stage in transferrin processing. If MVE represent an obligate step in transferrin/receptor recycling or iron removal, this would have important implications for the time course of the recycling process. Nunez et al. (23) calculated an iron uptake rate of 0.062 atom of iron/receptor for their orthochromatic erythroblast-reticulocyte fraction. This translates to one pair of iron atoms every 32 min for each receptor. Assuming removal of two iron atoms from each diferric transferrin internalized, this means that the average receptor cycle time is 32 min, which is kinetically compatible with passage of most receptor-ligand complexes through MVE. On the other hand, a shorter transferrin-receptor recycling time has been reported by other workers (35). This suggests that transferrin/receptor complexes may be recycled directly from the tubular and vesicular compartments without passing through MVE. Thus, MVE may be an optional detour rather than an obligate step in transferrin processing.

Recycling of transferrin-receptor complexes from MVE is supported by the apparent concomitant exocytosis of AuTf and vesicular inclusions derived from MVE. This may be observed in reticulocytes incubated with AuTf for 35 min at 37°C and after a subsequent 35-min chase with free transferrin. Exocytosis of vesicular inclusions could partially account for the loss of membrane surface area observed during reticulocyte maturation (3, 28). Zweig et al. (42) demonstrated that the cytoplasmic aspect of reticulocyte endosomes are free of spectrin by ferritin immunolabeling of ultrathin frozen sections, and they proposed that during reticulocyte maturation, these spectrin-free membranes are preferentially lost by intracellular degradation or by "exocytosis" (by which they mean inclusion in plasma membrane blebs that are removed in the spleen). We have demonstrated that MVE inclusions may be truly exocytosed (as opposed to the above process, which might be better termed "blebbing"), and this may provide an additional mechanism for the preferential loss of spectrin-free membrane. In some cases AuTf was associated...
Fig. 20: View of the inside of a scraped-open reticulocyte, revealing the presence of numerous clathrin-coated domes. Bar, 500 nm. × 41,000. Fig. 21: Higher power view of a clathrin-coated dome surrounded by the anastomosing spectrin network of the reticulocyte cytoskeleton. Bar, 100 nm. × 179,000.

Fig. 22 illustrates the possible routes of transferrin and its receptor during processing by the reticulocyte. After transferrin binds to its receptor, it is internalized via coated pits and vesicles. Subsequently, it appears in small vesicles and tubular structures. Removal of iron from transferrin may occur inside these structures and/or inside MVE. Recycling to the plasma membrane may occur from the small vesicle or tubule compartment. Transferrin passes into MVE and may then also recycle to the plasma membrane.

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