Dual control of caveolar membrane traffic by microtubules and the actin cytoskeleton

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Summary
Live cell, time-lapse microscopy was used to study trafficking of caveolin-1-GFP in stably expressing CHO cells. Multiple cytological and biochemical tests verified that caveolin-1-GFP was a reliable marker for endogenous caveolin-1. At steady state, most caveolin-1-GFP was either at the cell surface associated with invaginated caveolae or near the centrosome in caveosomes. Live cell fluorescence imaging indicated that while much of the caveolin-1-GFP in caveolae at the cell surface was relatively sessile, numerous, highly motile caveolin-1-GFP-positive vesicles were present within the cell interior. These vesicles moved at speeds ranging from 0.3-2 μm/second and movement was abolished when microtubules were depolymerized with nocodazole. In the absence of microtubules, cell surface invaginated caveolae increased more than twofold and they became organized into linear arrays. Complete depolymerization of the actin cytoskeleton with latrunculin A, by contrast, triggered rapid and massive movements of caveolin-positive structures towards the centrosomal region of the cell. The caveolar membrane system of CHO cells therefore appears to be comprised of three caveolin-1-containing compartments. These include caveolae that are confined to the cell surface by cortical actin filaments, the peri-centrosomal caveosomes and caveolar vesicles, which we call ‘cavicles’, that move constitutively and bi-directionally along microtubules between the cell surface and caveosomes. The behavior of cavicles suggests that they function as transport intermediates between caveolae and caveosomes.

Movies available online

Key words: Caveolin, Recycling endosome, Caveolae, Caveosome

Introduction
A detailed understanding of how microtubules are involved in membrane trafficking is rapidly emerging, but the role of actin filaments in this process remains relatively unknown. Microtubules serve as tracks for long range, rapid movements of transport intermediates between donor and acceptor compartments in the secretory and the endocytic pathways (Bloom and Goldstein, 1998). In addition, microtubules control the movement of organelles such as mitochondria and peroxisomes and the steady state distribution of the ER, the Golgi apparatus, endosomes and lysosomes. Actin filaments also appear to be involved in organelle and vesicle movements particularly along axons (Kuznetsov et al., 1992) and in microvilli (Fath and Burgess, 1993) but in general these movements occur both within a shorter range and have slower velocities than microtubule based movements.

Interestingly, there are several recent studies that show the movements of some organelles and vesicles are controlled by both microtubules and actin filaments. These include movement of phagosomes in macrophages (Al-Haddad et al., 2001), of lysosomes in hepatoma cells (Cordonnier et al., 2001), of mitochondria and other vesicles in neurons (Bridgman, 1999; Morris and Hollenbeck, 1995) and of pigment granules within the extensions of pigment producing cells (Rodionov et al., 1998; Rogers and Gelfand, 1998; Wu et al., 1998). The distribution of pigment granules in melanocytes provides a striking example of this coordinate control by both of these cytoskeletal arrays. Long distance, bi-directional movements of the granules occur along radially arranged microtubules, and require microtubule motor proteins of the kinesin and dynein families. In contrast, more spatially restricted movements occur along actin filaments and make use of the actin motor myosin Va (Rogers and Gelfand, 2000). Myosin Va also appears to be required to tether pigment granules to actin filaments located in the cell periphery and this is the mechanism responsible for determining the steady state distribution of the granules (Wu et al., 1998). Both microtubule and actin filaments may also play roles in the intracellular movement of endocytic membrane generated from clathrin coated pits (Apodaca, 2001). Actin seems to be important for events that take place during the budding process (Kamal et al., 1998; Lamaze et al., 1997) while microtubules appear to be responsible for the movement of newly budded endocytic vesicles to different sites within the cell (Kamal et al., 1998; Valetti et al., 1999). By contrast, little is known about how the cytoskeleton controls the dynamics of caveolae and the transport intermediates that might communicate with them.

Caveolae, or plasmalemmal vesicles, were originally described as flask-shaped, plasma membrane invaginations found in gall bladder epithelial and endothelial cells (Palade, 1953; Yamada, 1955) but are now known to be present in a broad variety of cell types (Anderson, 1998). Numerous
morphological studies have documented that caveolae mediate the movement of molecules across endothelial cells (Simionescu, 1983). Transcytosis appears to involve membrane invagination, vesicle budding and movement of vesicles filled with cargo to the opposite side of the cell. The principle method used to identify caveolae in these studies is their shape. A morphologic membrane signature works well in thin endothelial cells where vesicles travel only short distances to traverse the cell. In most cells, however, caveolae-derived transport intermediates rapidly mix with structures of similar morphology originating from other membrane compartments. As a consequence, it is very difficult to track the movement of caveolae-derived membranes in most cells.

The standard method for mapping the traffic of intracellular membranes is to follow the movement of a visible marker for that membrane. A well-accepted marker for caveolae in fibroblasts is caveolin-1 (Rothenberg et al., 1992). This integral membrane protein, a component of the cytoplasmic, filamentous coat that decorates the cytoplasmic surface of fibroblast and endothelial cell caveolae. Attachment of green fluorescence protein (GFP) to the C-terminal end of the molecule does not appear to interfere with its localization to caveolae (Pelkmans et al., 2001). Previous studies suggest that caveolin-1-GFP collects at sites of cell-cell contact (Volonte et al., 1999) and along the cleavage furrow during cytokinesis (Kogo and Fujimoto, 2000). Caveolin-1-GFP has also been used to show that SV-40 virus is internalized by caveolae and the caveosome, and in transfection of CHO cells (Pelkmans et al., 2001).

Here we describe caveolin-1 dynamics in CHO cells stably transfected with caveolin-1-GFP. We observed the fluorescent fusion protein in caveolae and the caveosome, and in constitutively motile vesicles and tubules that evidently serve as transport intermediates between them. Furthermore, because perturbation of either microtubules or actin filaments caused rapid and profound changes in the dynamics and localization of caveolin-1-GFP, we conclude that these two structurally distinct components of the cytoskeleton act coordinately to regulate trafficking of the caveolar membrane system.

Materials and Methods
Construction of caveolin-1-GFP and generation of stable caveolin-1-GFP-expressing cell lines
GFP was attached to the C-terminal end of caveolin-1. HindIII restriction sites were added to both 5' and 3' ends of the human caveolin-1 cDNA by PCR using the Expand High Fidelity PCR system (Roche Molecular Biochemicals, Indianapolis, IN). The PCR product was digested with HindIII and inserted into the HindIII site of the EGFP-N1 eukaryotic expression vector (Clontech, Palo Alto, CA). The proper orientation and sequence of caveolin-1-GFP were verified by sequencing. The polypeptide content (Brewer, 1994) was used to transfect CHO K1 cells with the caveolin-1-GFP construct. Transfected cells were selected by growing in 0.8 mg/ml G418 (Invitrogen, Carlsbad, CA) for 14 days. Individual clones were isolated and screened for expression of the caveolin-1-GFP by fluorescence microscopy, expanded and then frozen for future use. The transfected CHO cells used in each experiment were maintained in DMEM (Invitrogen, Carlsbad, CA) supplemented with 10% FCS plus and 40 µg/ml proline in a humidified CO2 (5%) incubator at 37°C. New batches of cells were established from frozen stocks every 10-12 weeks. Normal human fibroblasts were grown as previously described (Goldstein et al., 1983).

Live cell imaging
For live cell recordings, cells were sub-cultured at least 2 days before each experiment into 35 mm dishes which had been modified with a hole in the bottom to which a #1 thickness glass coverslip was glued (Mat Tek Corp, Ashland, MA). The dishes were placed in a humidified Plexiglas chamber and maintained at 37°C and 5% CO2 throughout each experiment. We did not see any effect of either the number of days in culture or cell density on caveolin-1-GFP movement. Cells were imaged using a Leica TCS-SP laser scanning confocal microscope with a 100× 1.4 NA, plan-achromatic lens using a computer controlled 488 nm argon laser to excite GFP or a 568 nm krypton laser to excite Alexa red dyes. Signals were collected at 1.6 second intervals using a photomultiplier tube, digitized (512×512 pixels) and converted into images using Leica TCS software. FRAP (fluorescence recovery after photobleach) was performed by exposing 3-10 bleach spots in areas of interest and bleaching with the 488 laser line at full power for 2 seconds. The Leica TCS software was used to quantify fluorescent intensities in photobleached regions, which were corrected for overall bleaching by comparison with areas that were not exposed to the full power of the laser.

Immunofluorescence and immunogold electron microscopy
Cells plated onto coverslips in 12-well dishes were fixed either with 4% paraformaldehyde (PFA) in PBS at room temperature or with methanol at −20°C. Cells fixed with PFA were permeabilized with 0.2% TX-100. Non-specific binding sites were blocked by incubating cover slips for 30 minutes in PBS containing 1% fish skin gelatin (Sigma, St Louis, MO). Fixed cells were incubated with primary antibodies diluted in PBS containing 1% fish skin gelatin for 30 minutes, washed with PBS several times and then incubated with the appropriate secondary antibodies conjugated to the specified Alexa dyes (Molecular Probes, Eugene, OR) for 30 minutes before mounting with Agua Poly/mount Polysciences). Images were collected using the Leica TCS-SP laser scanning confocal microscope and processed using Adobe Photoshop. Images usually consisted of from 10-16 optical slices in order to span the entire depth of the cell. Mouse anti-transferrin receptor (Zymed, San Francisco, CA) was used to label recycling endosomes in CHO cells. Rabbit anti-mannosidase II was kindly provided by Kelly Moreman. For electron microscopy (EM) analysis, cells were processed for indirect immunogold detection of either GFP or caveolin-1 and quantified as previously described (Smart et al., 1994).

Isolation of Triton-X-100-insoluble membranes, velocity sedimentation and immunoblotting
Triton-X-100-insoluble membranes were isolated as previously described (Liu and Anderson, 1995) with the following changes. Briefly, two 100 mm dishes of confluent CHO cells stably expressing caveolin-1-GFP were washed twice with ice cold buffer A (12.5 mM MES, 75 mM NaCl, pH 6.5). The cells were then scraped from the dish and solubilized in 1 ml of buffer B (1% Triton X-100 in buffer A containing 10 µM leupeptin, 500 µM benzamidine, 10 µg/ml soybean trypsin inhibitor, 1 µg/ml pepstatin A, 200 µg/ml phenylmethylsulfonyl fluoride and 1 mM EDTA) and incubated on ice for 20 minutes. The sample was then homogenized with a dounce homogenizer 20 times, mixed with 1 ml of 2.5 M sucrose in buffer A and loaded on the bottom of a 12-mL ultracentrifuge tube. The sample was overlaid with 8 ml of a 10-30% sucrose gradient in buffer A and centrifuged at 29,000 g for 21 hours at 4°C in a SW41 rotor. Equal fractions (700 µl) were collected and the protein was precipitated by adding TCA and deoxycholate (final concentrations 7% and 0.015%, respectively) from concentrated stock solutions. The pellets were washed once with acetone, solubilized in sample buffer (Laemmli, 1970) and separated by SDS-PAGE. Proteins were transferred to PVDF membranes at 20-30 volts overnight in transfer buffer (25 mM
Caveolin-1-GFP is a reliable marker for endogenous caveolin-1

To examine the intracellular trafficking of caveolin-1, we fused the C-terminal end of wild-type caveolin-1 to green fluorescence protein (GFP) and established a stable cell line expressing this protein. Several criteria were used to establish that this chimeric protein behaved like endogenous caveolin-1. One of the properties of caveolin-1 is that it is resistant to extraction in cold Triton X-100. Enriched fractions of detergent insoluble membranes were isolated from cells, fractionated on sucrose gradients and analyzed by immunoblotting with anti-caveolin-1 antibody (Transduction Laboratories, Lexington, KY) and the immune complexes collected by adding 50 µl of protein G-agarose (Sigma, St Louis, MO) for 1 hour. The precipitates were washed 3 times with buffer C, solubilized in sample buffer containing 10 mM DTT by boiling and separated by SDS-PAGE on a 5-20% gradient gel. Gels were treated with Amplify (Amersham Biosciences, Piscataway, NJ) and dried for autoradiography. Films were developed after 7 days.

Results

Caveolin-1-GFP behaves like endogenous caveolin-1. CHO cells stably transfected with caveolin-1-GFP were grown for 2 days until confluent. (A) Triton X-100-insoluble caveolae were separated from soluble membranes on sucrose gradients and analyzed by immunoblotting with anti-caveolin-1 (Fig. 1A). The low-density membranes migrated towards the top of the gradient (fractions 6-12), as judged by the presence of caveolin-1 (Fig. 1A). The distribution of caveolin-1-GFP was indistinguishable from that of endogenous caveolin-1, indicating that the chimeric protein was also associated with detergent insoluble caveolae (Cav-GFP, Fig. 1A). Comparison of the immunoreactive signals for caveolin-1-GFP and caveolin-1 demonstrated that the fluorescent fusion protein was expressed at a low level compared to caveolin-1. Quantification of immunoblots from this and similar experiments indicate that the molar ratio of caveolin-1-GFP to endogenous caveolin-1 ranged from 0.2 to 0.3 in the stably transfected CHO cells.

Fig. 1. Caveolin-1-GFP behaves like endogenous caveolin-1. CHO cells stably transfected with caveolin-1-GFP were grown for 2 days until confluent. (A) Triton X-100-insoluble caveolae were separated from soluble membranes on sucrose gradients and immunoblotted with anti-caveolin-1. Caveolin-1-GFP (Cav-GFP) co-fractionated with endogenous caveolin-1 (Caveolin-1). The intensity of the bands for the two proteins indicates that caveolin-1-GFP constitutes a minor amount of the total caveolin-1 pool. (B) Cells were incubated for 2.5 hours in the presence of 3H-palmitate to label acylated proteins. Caveolin-1 was immunoprecipitated, separated by PAGE and processed for autoradiography. Both the endogenous protein (caveolin-1) and the caveolin-1-GFP (cav-GFP) were covalently labeled with 3H-palmitate. (C) Caveolin was extracted from cells using a combination of Triton X-100 and deoxycholate to solubilize the caveolar membranes and then analyzed by velocity gradient sedimentation. Both the endogenous caveolin (caveolin-1) and caveolin-1-GFP (cav-GFP) form oligomers that migrate to the bottom of the gradient.

of caveolin-1 to oligomerize, another important property of native caveolin-1 (Monier et al., 1996). This was indicated by
co-fractionation of the fusion protein with caveolin-1 containing high density complexes on sucrose gradients of membranes solubilized with a mixture of Triton-X-100 and deoxycholate (Fig. 1C). Immunofluorescence microscopy (Fig. 2A) and immunogold EM (Fig. 2B) were also used to determine the distribution of caveolin (compare the caveolin distribution in the non-expressing cell at the top with one that is expressing the GFP construct below it). (B) Cells were grown for 2 days in 6-well tissue culture dishes and incubated for 1.75 hours in the presence of 10 µM nocodazole, which was used to enhance our ability to recognize caveolae at the cell surface of these cells. The samples were then fixed and processed for immunogold localization of GFP. Bars, 20 µm (A); 0.25 µm (B).

Caveosomes are clustered near the microtubule organizing center in close proximity to recycling endosomes in CHO cells

Conventional immunofluorescence of normal human fibroblasts has shown that caveolin-1 is frequently concentrated in patches at the edge of the cell and in linear arrays over the cell surface (Rothberg et al., 1992). Confocal imaging of caveolin-1-GFP in CHO cells showed a similar pattern (Fig. 2A). In addition to the typical surface staining of caveolin however, confocal optical sections taken through the cell showed that there were numerous scattered puncta of caveolin-1 throughout the cytoplasm and a prominent accumulation of caveolin positive membranes at the center of the cell (Fig. 2A, arrows). This peri-nuclear structure did not co-localize with the Golgi apparatus markers, mannosidase II (Fig. 3D-F) or GM130 (data not shown). Instead, it accumulated near the microtubule-organizing center (MTOC), which was visualized by anti-tubulin staining (Fig. 3A-C). The recycling endosome is also located near the MTOC in CHO cells. To compare the distribution of the caveolin-1-GFP compartment with recycling endosomes, cells were stained with anti-transferrin receptor antibody. Fig. 3G-I shows that caveolin-1-GFP and the transferrin receptor were highly concentrated within the same peri-centrosomal region. Caveolin-1-GFP generally covered a broader area than the transferrin receptor and at high magnification appeared to be largely distinct, although tightly intertwined with the compartment stained by the transferrin receptor antibody. Caveolin-1 positive membranes and recycling endosomes, therefore, share an overlapping distribution in these cells.

Motility of caveolin-1-GFP

The trafficking of caveolin-1-GFP was observed in living cells by confocal, time-lapse microscopy and was found to be remarkably dynamic. At least three different types of movement were seen. Many of the caveolin-1-GFP patches at the cell surface were relatively immobile (yellow arrows, Fig. 4A), exhibiting only localized tethered movements. Occasionally a patch of caveolin-1-GFP appeared to move away from the cell
surface, as if it were budding from the membrane, but these events were rare and difficult to capture. Since EM showed that cell surface caveolin-1-GFP was primarily associated with caveolae membranes (Fig. 2), we conclude that a substantial proportion of the caveolae in these cells are sessile. The possibility remains, however, that these seemingly immotile caveolae internalize and recycle locally without being detected by time lapse fluorescence microscopy (Anderson et al., 1992).

In addition to being localized at the cell surface, caveolin-1-GFP was observed in vesicle-like structures that commonly exhibited rapid movement along curvilinear paths within the cytoplasm. These movements were most visible near the lower surface of the cells just below the nucleus where the surface area is large. The velocities and distances traveled by these caveolar vesicles, or ‘cavicles’, were quite variable. Eighteen recordings in 15 different cells, were used to determine the apparent instantaneous velocities of individual cavicles and these ranged from 0.3-2.0 μm/seconds with uninterrupted movements that ranged from 2-16 μm. These rapid movements occurred in all directions, and frequently were interrupted by short pauses. A dramatic example of one of these long-range movements is shown in Fig. 4A (separation of white arrows). Many other examples are shown in the accompanying movies (http://jcs.biologists.org/supplemental) (Fig. 4C).

In cells expressing caveolin-1-GFP, we also routinely saw thin tubules decorated with caveolin-1-GFP (yellow arrows, Fig. 4B). These tubules were very dynamic and formed networks that would break, reform and undergo rippling movements (Fig. 4B; see also movies online). The tubules did not appear to be induced by overexpression of caveolin-1-GFP because they were observed in cells expressing both high and low levels of the fusion protein. These structures were not evident in conventionally fixed, untreated cells, however, implying that the tubules do not survive chemical fixation. We do not know the origin of these tubules, but they were present both at the cell periphery and in association with caveolin-1-GFP positive membranes that accumulate near the centrosome in CHO cells.

These observations suggest that caveolae can pinch off or elongate from the plasma membrane and travel into the cell, but ordinarily only a minor fraction of the total caveolae exhibit such activity during short time intervals. To determine whether the caveolin-1-GFP located in the peri-centrosomal region of the cell was exchanging with the other pools of caveolin-1-GFP we performed fluorescence recovery after photobleaching experiments (FRAP). The peri-centrosomal region of the cell was completely bleached using the 488 laser at full power for 2 seconds and recovery monitored over a 5 minute and 40 second recovery period. Fig. 5 shows images taken before bleaching, immediately after bleaching (0 time) and 5 minutes after the bleach period. Recovery was apparent within approximately 1 minute and there was a clear accumulation of GFP in the same region by 3 minutes. Quantification of the photobleached area demonstrated that recovery was 100% when the decrease in the overall signal due to FRAP-independent bleaching was subtracted. Recovery was not affected by pretreatment of the cells with cyclohexamide indicating that caveolin-1-GFP was being transported from the other regions of the cell and was not due to new synthesis. The movie of the bleach and recovery can be viewed online (Fig. 5A).
Effects of disrupting microtubules

Both the speed of caveolin-1-GFP movements within the cell and the fact that they followed curvilinear paths suggested that cavicles were traveling along microtubules. This was confirmed by treating caveolin-1-GFP expressing cells with nocodazole to depolymerize microtubules. Almost all caveolin-1-GFP movement ceased under these conditions (Fig. 6A; see also movies online). Unexpectedly, nocodazole also caused caveolin-1-GFP to accumulate on the cell surface in linear arrays that were most apparent near the bottom surface of CHO cells (Fig. 6B). The caveolin-1-GFP in the linear arrays was nearly immobile, and even the tethered movements were reduced.

The linear arrays of caveolin-1-GFP that were amplified by nocodazole were reminiscent of stress fibers, which represent bundles of actin filaments. Accordingly, we used fluorescent phalloidin to compare the distributions of actin filaments and caveolin-1-GFP. Surprisingly, linear arrays of caveolin-1-GFP were not always aligned along or adjacent to stress fibers in nocodazole-treated cells (Fig. 7) and were occasionally found in areas where fluorescent phalloidin staining was weak or absent.

We also examined the effects of nocodazole on the behavior of endogenous caveolin-1 in normal human fibroblasts (Fig. 8). A significant amount of caveolin-1 is normally organized in linear arrays in these cells (Fig. 8A); however, the number and thickness of these arrays increased markedly after cells were exposed to nocodazole (compare Fig. 8A with 8C). Anti-tubulin staining showed that the microtubules had been disrupted by the nocodazole treatment (compare Fig. 8B with 8D). A similar redistribution of endogenous caveolin-1 into linear arrays was observed in NIH3T3 cells and rat embryo fibroblasts (data not shown). When normal human fibroblasts treated with nocodazole were examined by electron microscopy, we saw a more than twofold increase in the number of invaginated caveolae (Fig. 8E). These structures were positive for immunogold labeling with anti-caveolin-1 (data not shown). This increase was quantified using numerous micrographs and the data is presented in Table 1. We conclude that microtubules are required for long-range movement of caveolin-1 positive structures and in their absence both caveolin-1 and invaginated caveolae accumulate at the cell surface.

Effects of disrupting the actin cytoskeleton

The arrangement of caveolae in linear arrays, particularly in fibroblasts, has been described before and is believed to be due to an association of caveolae with the cortical actin cytoskeleton (Izumi et al., 1989). To determine if actin was involved in organizing the caveolin-1 into these linear arrays, we looked at the combined effects of nocodazole and drugs known to disrupt actin filaments (Fig. 9). Cells expressing caveolin-1-GFP were

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Fig. 4. Caveolin-1-GFP displays three different types of movement. Cells were plated 48 hours before use into culture dishes that had a glass coverslip glued to the bottom. Confocal sections were taken at 1.6 second intervals near the bottom of the cell, just under the nucleus in cells stably expressing caveolin-1-GFP. (A) Some of the caveolin-1-GFP in these cells showed only saltatory movements (yellow arrows). A 12.8 second recording showed, however, that other patches of caveolin-1-GFP in the same cell rapidly moved apart (follow white arrows). (B) A third population of caveolin-1-GFP was associated with dynamic tubular structures (yellow arrows), which can also be seen in the accompanying movie. (C) Stills from two videos showing examples of long range and saltatory movement in control cells. Bar, 10 μm (A); 5 μm (B). The movies that correspond to panels B and C can be viewed online (http://jcs.biologists.org/supplemental).
incubated in the presence (Fig. 9B,D,F) or absence (Fig. 9A,C,E) of nocodazole plus either media alone (Fig. 9A,B), cytochalasin D (Fig. 9C,D), which partially disassembles actin filaments, or latrunculin A (Fig. 9E,F) which depolymerizes actin filaments completely. Projection images of the entire stack of control cells showed a patchy distribution of caveolin-1-GFP with prominent accumulations in the center of the cell (arrows). Nocodazole treatment caused the peri-centrosomal accumulations of caveolin-1-GFP to disappear with a concomitant appearance of linear arrays at the cell surface (Fig. 9B). Cytochalasin D at low concentrations had little effect on the overall distribution of caveolin-1-GFP (Fig. 9C) but completely prevented the formation of linear arrays in nocodazole treated cells (Fig. 9D). Latrunculin A, by contrast, dramatically increased the amount of caveolin-1-GFP both in the center of the cell and in irregularly shaped structures dispersed throughout the cytoplasm. This was accompanied by a loss of caveolin-1-GFP from the cell surface (Fig. 9E). In cells exposed to both latrunculin A and nocodazole (Fig. 9F), caveolin-1-GFP was only found in dispersed, irregularly shaped structures and no longer accumulated at the center of the cell. The results of these experiments indicate that cortical actin filaments confine and organize caveolae near the cell surface while microtubules carry cavicles that have been released from actin tethers into the cell.

The effects of latrunculin A on the localization of caveolin-1-GFP could be better appreciated by following the marker at early times after cells were exposed to the drug (Fig. 10A). Within 1 minute of exposure to latrunculin A, there was a massive increase in the inward movement of the caveolin-1-GFP. Aggregates of caveolin-1-GFP positive structures began to separate from the plasma membrane and migrate towards the center of the cell (Fig. 10A, 1 min; see also movies online). By 10 minutes, most cells were rounding up but remained attached to the substratum by several elongated processes. They persisted in this state for several hours. Caveolin-1-GFP could be seen moving rapidly into and out of these processes as well as through other regions of the cell (Fig. 10A, 10 min; see also movies online). By 20-30 minutes there was an obvious accumulation of the caveolin-1-GFP in the center of the cell and cavicles could still be seen moving towards and away from this region (Fig. 10A, 30 min). These rapid movements occurred along microtubules because they were completely eliminated when cells were treated with both nocodazole and latrunculin A (Fig. 6C; see also movies online). Live cells were

| Treatment   | Number of cells | Number of caveolae | Length (cm) | Caveolae/ unit length | % increase |
|-------------|-----------------|--------------------|-------------|-----------------------|------------|
| None        | 142             | 593                | 0.2         | 0.3                   | 0          |
| Nocodazole  | 143             | 1102               | 0.15        | 0.73                  | 155        |

Table 1. Quantification of the nocodazole-induced increase in surface caveolae

Fig. 5. FRAP analysis showing that there is active exchange between caveolin-1-GFP located in the peri-centrosomal region and the other pools of caveolin. Cells stably expressing caveolin-1-GFP were plated 48 hours before viewing into culture dishes that had a glass coverslip glued to the bottom. Confocal images were taken at 1.6 second intervals for approximately 1 minute and then the entire peri-centrosomal region of the upper cell was bleached in 6 spots for 2 seconds each using the 488 laser at full power. Immediately following the bleach, images were again collected at 1.6 second intervals for another 5 minutes and 40 seconds. Panel A shows the prebleach image and peri-centrosomal accumulations of caveolin-1-GFP in two cells. The central accumulation in the top cell was then bleached (area within the white circle, B). Panel C shows that the accumulation at the center of the cell has recovered within 5 minutes, indicating that caveolin-1-GFP travels between the various caveolin positive compartments. The corresponding movie can be viewed online (http://jcs.biologists.org/supplemental).

Fig. 6. Nocodazole blocks the movement of caveolin-1-GFP in live cells and organizes caveolin-1-GFP into linear arrays. CHO cells stably expressing caveolin-1-GFP were grown for 2 days before being viewed by confocal microscopy, as described in Materials and Methods. (A) Still from video of CHO cells pretreated with 10 μM nocodazole for 1 hour before viewing. Long range movements have disappeared. (B) Immunofluorescence micrograph of CHO cells pretreated with nocodazole taken at the bottom surface of the cell. (C) Still from video of stably expressing CHO cells pretreated with 10 μM nocodazole for 1 hour before the addition of 1 μM latrunculin A. Within 1 minute the arrays have disappeared and everything appears to be immobile. Bar, 10 μm. The movies that correspond to panels A and C can be viewed online (http://jcs.biologists.org/supplemental).
first treated for 30 minutes with nocodazole in order to form arrays and then latrunculin A was added. Within 1 minute of the addition of latrunculin A the arrays had disappeared. In the presence of both drugs there was virtually no movement. Furthermore, latrunculin A did not induce caveolin-1-GFP to accumulate near the centrosome in cells that were treated with nocodazole (Fig. 6C, Fig. 8F).

The effect of latrunculin A on the distribution of caveolin-1 in untransfected normal human fibroblasts was also examined using immunogold electron microscopy (Fig. 10B). The anti-caveolin-1 immunogold was found associated with invaginated caveolae before treatment (not shown). By contrast, within minutes after adding latrunculin A, the immunogold label was found associated with arrays of 50-70 nm invaginations that extended from the surface into the cell (e.g. between black arrows). Also common were circular profiles of membrane, containing numerous invaginated caveolae (asterisk) that appeared to have detached from the surface. A typical example of one of these clusters is shown in the inset of Fig. 10B. These structures seen by EM correspond well to the large, irregularly shaped structures that were seen by fluorescence microscopy in caveolin-1-GFP expressing cells (compare Fig. 10A and 10B). We do not know if these membrane profiles are transport intermediates or simply regions of membrane that buckled in the absence of an actin support. Nevertheless, they always contained multiple, invaginated caveolae.
Discussion
Thanks to a few recent studies of cultured mammalian cells transiently expressing GFP-tagged versions of caveolin-1, a picture of caveolar membrane dynamics in individual cells has begun to emerge (Pelkmans et al., 2001; Thomsen et al., 2002). We have extended that approach by stably transfecting caveolin-1-GFP into a cell type, CHO, in which fluorescent caveolin-1 analogues have not been studied previously. Live cell, time-lapse microscopy of these cells revealed several novel aspects of caveolin-1 trafficking. First, we found that CHO cells, in contrast to other cell types that have been studied by similar approaches, contain an abundance of constitutively motile caveolar vesicles, or ‘cavicles’ that were revealed by FRAP experiments to serve as transport intermediates between plasma membrane caveolae and peri-centriolar caveosomes in the absence of specific extracellular stimuli. The finding that this form of trafficking was abolished by nocodazole indicated that microtubules serve as the tracks along which motile cavicles move. Second, the trafficking of cavicles from caveolae to caveosomes was observed to be abruptly upregulated when actin filaments were disassembled by exposure of cells to latrunculin A. The cortical network of actin filaments therefore appears to constrain either the formation of cavicles from caveolae, or the movement of newly formed cavicles towards the centrally located caveosomes. Thus, microtubules and actin filaments act coordinately to control membrane trafficking in the caveolar system. Finally, comparison of the distributions of caveosomes with other peri-nuclear endomembrane compartments emphasizes the existence of an extensive caveolar membrane system that is analogous to the endosomal membrane system and there may also be at least a partial merger of the caveosome with the recycling endosome. A model that summarizes many of these findings is presented in Fig. 11.

Fig. 9. Disruption of the actin cytoskeleton blocks the formation of the linear arrays of caveolin-1-GFP induced by nocodazole. CHO cells stably expressing caveolin-1-GFP were plated on coverslips and grown for 2 days, as described in Materials and Methods. Cells were incubated for 30 minutes in the absence (A,B) or presence of either 2.5 μM cytochalasin D (C,D) or 1 μM latrunculin A (E,F) and then incubated for an additional 60 minutes with (B,D,F) or without (A,C,E) 10 μM nocodazole, before fixing with paraformaldehyde and processing for visualization of GFP. Note that both cytochalasin D and latrunculin A blocked the redistribution of caveolin into linear arrays (compare B with D and F) and that nocodazole blocked the latrunculin A induced redistribution of caveolin-1-GFP to the cell center (compare E with F). Each image is the projection of a stack of confocal images that extend through the entire cell. Bar, 20 μm.

Recycling endosomes and the caveosomes
In CHO cells we found that caveosomes accumulated at the MTOC. This observation was reminiscent of the accumulation of caveolin containing membranes near the centrosome of A431 cells exposed to okadaic acid plus hypertonic medium (Parton et al., 1994). Not all cells accumulate caveolin-1 near the centrosome, however, because we did not observe such accumulations in human fibroblasts (data not shown). Centrosomal caveolin-1-GFP in CHO cells has a very similar distribution to that of the transferrin receptor-rich recycling endosomes (Maxfield and Yamashiro, 1987). This region of a CHO cell is difficult to resolve by immunofluorescence microscopy but careful examination of multiple stacked confocal images suggested that the two membrane systems are intertwined rather than in the same compartment. Nevertheless, the close apposition of caveosomes and recycling endosomes and their partial co-localization at the light microscope level may provide opportunities for direct communication between the caveolar and the clathrin-mediated endocytic systems.

Biochemical evidence for communication between these two membrane systems has already been reported, as anti-transferrin receptor IgG was able to immunoprecipitate both the receptor and caveolin-1 (Gagescu et al., 2000). Recycling endosomes appear to share other caveolae markers including...
flotillin-1, sphingomyelin and cholesterol (Gagescu et al., 2000). Moreover, the folate receptor, a GPI anchored protein that clusters in caveolae (Ying et al., 1992), has been found in recycling endosomes containing transferrin (Mayor et al., 1998). Perhaps this ligand entered cells through caveolae before being delivered to recycling endosomes, even though many other ligands found in recycling endosomes indisputably arrive there via the coated pit-dependent pathway. In support of this idea, our own studies looking at the uptake of rhodamine-tagged transferrin suggest that at early time points transferrin and caveolin are internalized into separate vesicles but at later time points the transferrin pathway can communicate with caveolin-1 containing compartments (data not shown).

In as much as the centrosome is the principal site of internal caveolin-1-GFP positive membranes in CHO cells, we favor the idea that this compartment corresponds to the recently described caveosome (Pelkmans et al., 2001). To date the principle markers for the caveosome are caveolin, and SV-40 virus. Caveosomes accumulated SV-40 virus that had been internalized by caveolae in CV-1 cells. The caveosomes in CV-1 cells were not clustered at the centrosome, but this difference may simply reflect cell type specific variability. A recent paper by Nichols (Nichols, 2002) also describes a distinct class of caveolin-1 positive endosomes that accumulate 10K dextran, cholera toxin, GPL-GFP and SV-40 virus. We found that our cavicles also contain 10K dextran and cholera toxin (data not shown) a finding that contrasts with the reported lack of dextran uptake in SV-40-infected CV-1 cells. One possible explanation for this difference is that during infection with the virus the caveosome is somehow modified to exclude other cargo. Further work is required before this compartment can be fully defined functionally and biochemically and to determine the extent, if any, to which caveosomes communicate with recycling endosomes.

**Dual control of caveolin-1-GFP traffic**

Our findings indicate that actin filaments and microtubules play opposing roles in regulating the steady state distribution of caveolin-1-GFP. Whereas the cortical actin cytoskeleton appears to confine caveolin-1-GFP at the cell surface, microtubules serve as tracks for the transport of caveolin through the cytoplasm. The speed of these microtubule based movements ranged from 0.3 μm/seconds to as high as 2 μm/second, which is consistent with dynein and kinesin directed movements of organelles and vesicles along microtubules. Although the microtubule motors responsible for caveolae motility remain to be defined, one leading candidate is the recently described minus end-directed kinesin, KIFC3. This motor reportedly moves detergent resistant membranes like caveolae along microtubules (Noda et al., 2001) Depolymerization of microtubules caused a redistribution of the caveolin-1-GFP located at the centrosome to punctate structures located throughout the cell as well as an increase in the total number of invaginated caveolae at the cell surface. In
contrast to the apparent trapping of caveolae at the cell surface in the absence of microtubules, elimination of actin filaments caused a marked decline in caveolin-1-GFP located near the cell surface and a corresponding increase in peri-centrosomal caveolin-1-GFP. This suggests that surface caveolae have the potential to bud from the plasma membrane and travel into the cell, but are prevented from doing so by the actin cytoskeleton.

How do actin filaments restrain the inward movement of caveolin-1? Two obvious possibilities are that cortical actin filament networks act as simple physical barriers to the detachment of caveolae from the plasma membrane or the inward movement of caveicles beyond the cortex. In either of these cases, local disassembly of cortical actin networks would be necessary to initiate inward transport of caveolae along microtubules. Another possibility is that cross-linking molecules attach caveolae or caveicles to actin filaments in the cell cortex. In this case, either removal of the cross-links or local depolymerization of actin filaments would allow caveolae to move inward. In vitro studies have shown that caveolae bud from isolated membranes in a process that requires ATP, GTP and cytosolic factors (Gilbert et al., 1999). A similar in vitro system has been used to study the budding of clathrin coated pits from purified fibroblast membrane (Lin et al., 1991). In this system release of a subpopulation of coated vesicles requires a proteolytic cleavage step to release the budding membrane from its spectrin/ankyrin linkage with the actin cytoskeleton (Kamal et al., 1998). Interestingly, only this subpopulation of vesicles is able to move along microtubules to a degradative compartment. A similar mechanism may regulate caveolae budding and caveicle traffic. Once formed from budded caveolae, caveicles may associate with motor molecules that propel them along microtubules. Actin regulation of the caveolar membrane system fits well with the growing evidence that spectrin and actin regulate membrane traffic in other membrane systems as well (Lippincott-Schwartz, 1998).

The behavior of caveolae and caveicles is highly reminiscent of the movements of pigment granules. In mammals, the pigment melanin that is deposited in the skin and hair is produced in melanocytes and transported to the cell periphery where it is secreted and then endocytosed by adjacent keratinocytes. In dilute mice, which lack myosin Va, the amount of secreted pigment is drastically reduced resulting in their characteristic light coat color (Wu et al., 1998). Time lapse video microscopy of wild type and dilute melanocytes has shown that fast, bi-directional, microtubule dependent transport of melanosomes is the mechanism responsible for long range transport of melanosomes, while accumulation at the dendritic tips where they are secreted is due to the myosin Va dependent capture of these organelles in the actin-rich periphery. This generates an actin-filament dependent polarized distribution of organelles that are transported bi-directionally along microtubules. Several unconventional myosins have now been implicated in organelle transport (reviewed by Tuxworth and Titus, 2000). These include a related, myosin V dependent mechanism that regulates pigment granule distributions in fish scale melanophore cells (Rodionov et al., 1998; Rogers and Gelfand, 1998), the involvement of myosin 1vα in the movement and distribution of lysosomes (Cordonnier et al., 2001), and a role for myosin VII in clathrin mediated endocytosis (Buss et al., 2001). It will be interesting to learn whether any myosins are similarly involved in establishing the steady state enrichment of caveolae at the plasma membrane, and of caveosomes near the centrosome.

Our data are in strong agreement with the results obtained by other investigators who have used caveolin-GFP (Pelkmans et al., 2001) or GFP-caveolin (Thomsen et al., 2002) to demonstrate that cell surface caveolae are relatively stationary. They are also consistent with a report that GFP-caveolin-1 is present on a population of relatively motile structures scattered throughout the cytoplasm (Thomsen et al., 2002). In other respects, however, our results seem to be at odds with those reported in prior similar studies. For example, caveolin-1-GFP was used to demonstrate that binding of SV-40 virus could trigger internalization of caveolar membranes from the surface of CV-1 cells, which was rarely observed in the absence of virus (Pelkmans et al., 2001). Although SV-40 can clearly induce uptake of caveolae, it is not clear what normally initiates this process in virus-free cells. Our data indicate that in CHO cells, at least, caveolae are internalized constitutively, albeit at lower rates than have been observed for clathrin coated pit uptake. Likewise, photobleaching of peri-centrosomal GFP-caveolin-1 in transiently transfected HeLa cells (Thomsen et al., 2002) was followed by fluorescence recovery that was much slower than we observed in CHO cells that stably expressed caveolin-1-GFP. Although the causes of this discrepancy are unknown, leading suspects include cell type-specific differences in caveolar membrane trafficking, functional differences between GFP-caveolin-1 and caveolin-1-GFP, and overexpression of GFP-caveolin-1 in the
transiently transfected HeLa cells. The last possibility seems especially likely, because the low level of FRAP that was observed in HeLa cells whose peri-centrosomal regions were photobleached was abolished by pretreating the cells with cycloheximide. This implies that GFP–caveolin-1 was accumulating in the Golgi complex of the HeLa cells as a result of overexpression. In contrast, we show here that caveolin-1–GFP in stably transfected CHO cells was expressed at low levels relative to endogenous caveolin-1, and that peri-centrosomal caveolin-1–GFP in the CHO cells was not associated with the Golgi, but rather with caveosomes that fortuitously accumulated near the MTOC. This allowed us to bleach caveosomes selectively and to show that rapid trafficking of caveolin-1–GFP to these organelles occurred continuously in the presence of cycloheximide. In CV-1 cells Pelkmans et al. (Pelkmans et al., 2001) also only saw caveolin-1–GFP in the Golgi apparatus in a few cells that were highly overexpressing the fusion protein. More work is needed to determine if these are cell-specific differences in caveolar membrane traffic, or simply reflect the influence of the probes used to monitor the movement of these membranes. Whatever the explanation proves to be, the caveolar membrane system is clearly more extensive and dynamic than has been previously appreciated.

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