

vators of the ubiquitin-proteasome pathway, or a combination of the two have therapeutic potential for fALS. In conclusion, our results demonstrate that mutant SOD1 is degraded by at least two pathways, macroautophagy and the proteasome pathway, and that the clearance of mutant SOD1 by macroautophagy reduces its cell toxicity. These findings may provide insight into the molecular mechanisms of the pathogenesis of fALS.

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The Region-Specific Functions of Two Ubiquitin C-Terminal Hydrolase Isozymes along the Epididymis

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Abstract: We previously showed that *gad* mice, which are deficient for ubiquitin C-terminal hydrolase L1 (UCH-L1), have a significantly increased number of defective spermatozoa, suggesting that UCH-L1 functions in sperm quality control during epididymal maturation. The epididymis is the site of spermatozoa maturation, transport and storage. Region-specific functions along the epididymis are essential for establishing the environment required for sperm maturation. We analyzed the region-specific expression of UCH-L1 and UCH-L3 along the epididymis, and also assessed the levels of ubiquitin, which has specificity for UCH-L1. In wild-type mice, western blot analysis demonstrated a high level of UCH-L1 expression in the caput epididymis, consistent with ubiquitin expression, whereas UCH-L3 expression was high in the cauda epididymis. We also investigated the function of UCH-L1 and UCH-L3 in epididymal apoptosis induced by efferent duct ligation. The caput epididymides of *gad* mice were resistant to apoptotic stress induced by efferent duct ligation, whereas *Uchl3* knockout mice showed a marked increase in apoptotic cells following ligation. In conclusion, the response of *gad* and *Uchl3* knockout mice to androgen withdrawal suggests a reciprocal function of the two UCH enzymes in the caput epididymis.

Key words: apoptosis, epididymis, ubiquitin, UCH

Introduction

The mammalian epididymis is a highly convoluted tubule that connects the efferent ducts of the testis to the vas deferens [2, 8]. The epididymis is composed of three distinct compartments, caput (head), corpus (body)

and cauda (tail), each having a specific role in sperm maturation, sustenance, transport, and storage [2, 6]. However, the molecular basis for the maturation process remains largely unknown.

It has been suggested that the epididymis acts as a quality control organ to eliminate defective spermato-

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zoa before ejaculation [37]. The epididymis is an organ with voluminous protein traffic between the epithelium and lumen. Numerous proteins, secreted in an apocrine manner by the epididymal epithelium, are implicated in spermatozoa maturation [18]. Two major components of the ubiquitin-dependent proteolytic pathway, ubiquitin and UCH-L1 (PGP9.5), are expressed in epididymal tissue [10, 35]. Ubiquitin is present in human seminal plasma [26], and defective spermatozoa become ubiquitinated during epididymal passage [23, 37]. Our previous work showed that UCH-L1 associates with monoubiquitin and stabilizes its expression [31]. In addition, it has been suggested that UCH-L1 functions as a regulator of apoptosis via the ubiquitin pathway [13, 23, 25]. We found that testes of gracile axonal dystrophy (*gad*) mice, which lack UCH-L1, have reduced ubiquitin levels and are resistant to cryptorchid injury-mediated germ cell apoptosis [25]. Furthermore, our recent work demonstrated that the percentage of morphologically abnormal spermatozoa is significantly higher in *gad* mice, compared with wild-type mice [23].

Two mouse UCH isozymes, UCH-L1 and UCH-L3, are strongly but reciprocally expressed in the testis during spermatogenesis [25], suggesting that these proteins have distinct functions in the testis [23], even though they have high amino acid sequence identity and share significant structural similarity [21]. The functional regionalization of the epididymis is delineated at the molecular level by regional differences in gene expression [16–19]. Regional differences along the epididymis might be essential characteristics of the environment required for sperm quality control. Although it has been shown that UCH-L1 and UCH-L3 have reciprocal functions with respect to cryptorchid injury, their molecular functions in regulating sperm quality during epididymal passage are not fully understood. Thus, we examined the epididymal expression of UCH-L1 and UCH-L3 with regard to their involvement in the regulation of apoptosis. In addition, we assessed the reciprocal functions of these two proteins in the epididymis.

Materials and Methods

Animals

We used *gad* (CBA/RFM) [34] and *Uchl3* knockout (C57BL/6J) [21] male mice at 10 weeks of age. The

gad mouse is an autosomal recessive mutant that was obtained by crossing CBA and RFM mice. The *gad* line has been maintained by intercrossing for more than 20 generations [34]. *Uchl3* knockout mice were generated by the standard method [21] using homologously recombinant ES cells, and the knockout line has been back-crossed several times to C57BL/6J mice. Both strains are maintained at our institute. Animal care and handling were in accordance with our institutional regulations for animal care and were approved by the Animal Investigation Committee of the National Institute of Neuroscience, National Center of Neurology and Psychiatry.

Unilateral efferent duct ligation

Animals were either left intact to serve as controls or were unilaterally ligated at the efferent duct [9, 38]. Four mice in each group were anesthetized with pentobarbital (Abbott Laboratories, North Chicago, IL), and the testis and epididymis on the right side were exposed through a scrotal incision. The thin avascular attachment joining the initial segment of the epididymis to the tunica albuginea was cut to permit exposure of the efferent ducts coursing above and parallel to the vascular supply. A silk suture was passed by needle through the thin sheet of connective tissue between the ductules and the blood vessels, and the efferent ducts were occluded by ligation. Mice were sacrificed 2 or 4 days after ligation. Both epididymides were immersed in 4% paraformaldehyde for at least 24 hr before they were dehydrated and embedded in paraffin [22].

Histological and immunohistochemical assessment of the epididymis

The caput, corpus and cauda epididymides along the epididymal region embedded in paraffin were cut into 4- μ m sections and stained with hematoxylin and eosin. Light microscopy was used for routine observations. For immunohistochemical staining, the sections were incubated with 10% goat serum for 1 h at room temperature followed by incubation overnight at 4°C with a rabbit polyclonal antibody raised against peptides within UCH-L1 or UCH-L3 (1:1,000 dilution; peptide antibodies [24]) and ubiquitin (1:500; DakoCytomation, Glostrup, Denmark) in PBS containing 1% BSA. Sections were then incubated for 1 h with biotin-conjugated anti-rabbit IgG diluted 1:200 in PBS, followed by

Vectorstain ABC-PO (Vector Laboratories, Burlingame, CA) for 30 min at room temperature. Sections were developed using 3,3'-diaminobenzidine and counterstained with hematoxylin.

In situ apoptosis was detected by TUNEL (TdT-mediated nick end-labeling) staining with the DeadEnd Fluorometric TUNEL system (Promega, Madison, WI) according to the manufacturer's instructions, to identify apoptotic cells in situ via specific labeling of nuclear DNA fragmentation. Quantification was performed using four mice on each of postoperative days 0, 2 and 4. The total number of apoptotic cells was determined by counting the positively stained nuclei in each caput epididymis section [9]. Four sections from each mouse and 100 total circular tubules per group were processed.

Western blotting

Western blots were performed as previously reported [24]. Total protein (10 µg/lane) from each epididymal region including spermatozoa was subjected to SDS-polyacrylamide gel electrophoresis using 15% gels (Perfect NT Gel, DRC, Japan). Proteins were electrophoretically transferred to polyvinylidene difluoride membranes (Bio-Rad, Hercules, CA) and blocked with 5% non-fat milk in TBS-T (50 mM Tris base, pH 7.5, 150 mM NaCl, 0.1% (w/v) Tween-20). The membranes were incubated individually with primary antibodies to monoubiquitin (1:1,000; u5379, Sigma-Aldrich, St. Louis, MO), UCH-L1 and UCH-L3 (1:1,000 dilution; anti-peptide antibodies [24]), p53, Bax, and Bcl-xL (1:1,000 dilution; all from Cell Signaling Technology, Beverly, MA), and Bcl-2 (1:500; Transduction Laboratories, Franklin Lakes, NJ). Blots were further incubated with peroxidase-conjugated goat anti-mouse IgG or goat anti-rabbit IgG (1:5,000 dilution; Pierce, Rockford, IL) for 1 h at room temperature. Immunoreactions were visualized using SuperSignal West Dura Extended Duration Substrate (Pierce) and analyzed using a ChemiImager (Alpha Innotech, San Leandro, CA). Each protein level was normalized to α -tubulin following analysis with a ChemiImager using AlphaEase software.

Statistical analysis

The mean and standard deviation were calculated for all data (presented as mean \pm SD). Student's *t*-test was used for statistical analysis.

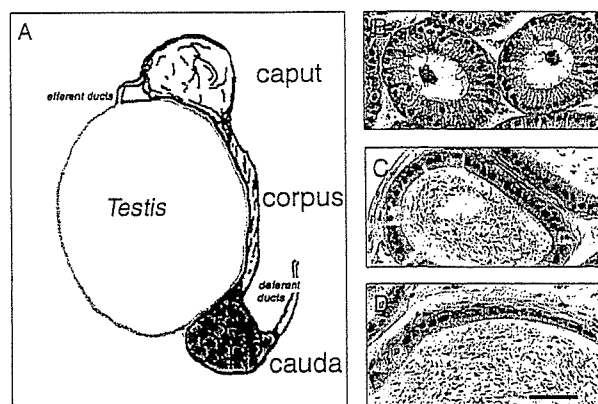


Fig. 1. A: Diagram of the epididymis. B-D: Morphology of the caput (B), corpus (C) and cauda (D) epididymidis from a wild-type mouse. Magnification, 200 \times . Scale bar, 40 μ m.

Results

Levels of UCH-L1 and UCH-L3 in individual epididymal regions

The epididymis is a single long, coiled tubule situated on the surface of the testis (Fig. 1A). The epididymal epithelium is composed of four major cell types, principal cells, basal cells, clear cells and narrow cells [7], and can be divided anatomically and functionally into three regions, the caput, corpus and cauda epididymis (Fig. 1B, C, D). We used western blotting to characterize UCH-L1 and UCH-L3 levels along the epididymis (Fig. 2). In wild-type mice, the level of UCH-L1 was highest in the caput epididymis and that of UCH-L3 was highest in the cauda epididymis. UCH-L1 and UCH-L3 were not observed in *gad* and *Uchl3* knockout mice, respectively (Fig. 2; comparison of UCH-L1 and UCH-L3 levels with those in wild-type control mice).

Immunohistochemistry of UCH-L1, UCH-L3 and ubiquitin in the epididymis

Under light microscopy, granular and diffuse UCH-L1 and UCH-L3 immunoreactivity was detected in many epithelial cells of the caput, corpus and cauda epididymis in wild-type mice (Fig. 3A, C). Granular immunoreactivity to ubiquitin was seen in the epithelial cells of the epididymis (Fig. 3B). The distribution of ubiquitin in the corpus and cauda epididymal epithelial cells was similar to that of the caput epididymis,

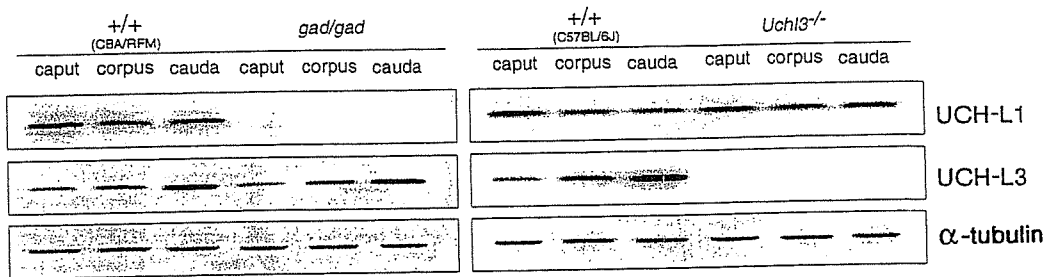


Fig. 2. Comparison of UCH-L1 and UCH-L3 expression by western blotting of caput, corpus and cauda epididymis lysates from two wild-type (CBA/RFM and C57BL/6J), *gad* and *Uchl3* knockout mice. Blots were reprobbed for α -tubulin, which was used to normalize the protein load. Images are representative of four independent experiments.

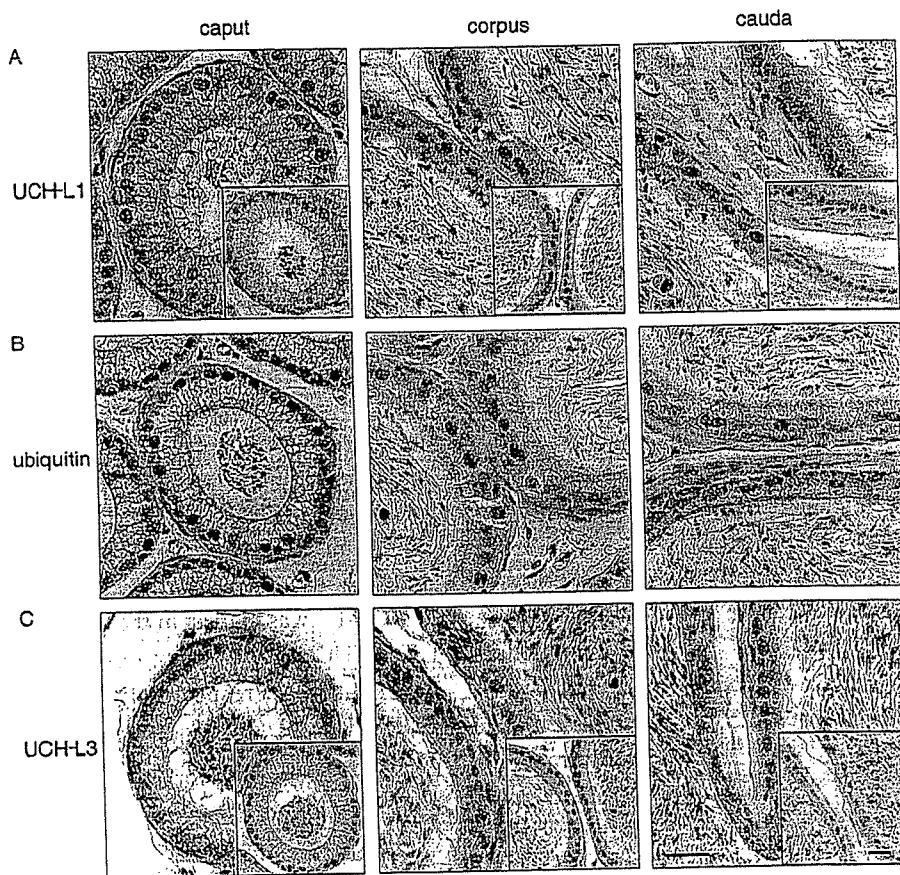


Fig. 3. Immunohistochemistry of UCH-L1, UCH-L3, and ubiquitin in the individual epididymal regions of wild-type mice. Each of the protein-positive cells in the caput, corpus and cauda epididymis is stained by DAB. The insets show that no cells are positive for UCH-L1 and UCH-L3 in the individual epididymal compartments from *gad* (A) and *Uchl3* knockout (C) mice, respectively. A: UCH-L1-positive cells. B: Ubiquitin-positive cells. C: UCH-L3-positive cells. Magnification, 400 \times . Scale bar, 20 μ m.

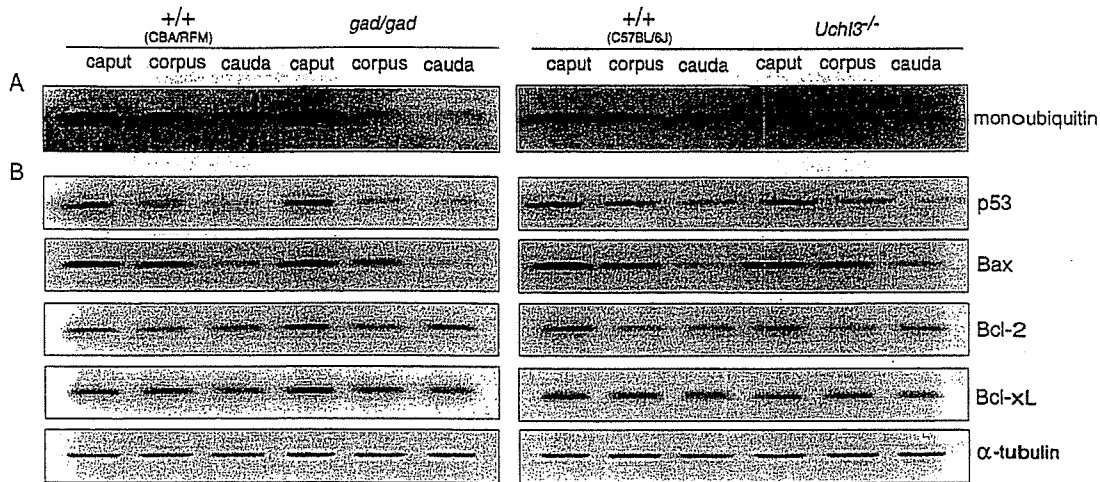


Fig. 4. Western blot analyses showing monoubiquitin and apoptotic proteins in the individual epididymal regions. Total protein (10 μ g per lane) was prepared from the caput, corpus and cauda epididymidis from two wild-type (CBA/RFM and C57BL/6J), *gad* and *Uchl3* knockout mice. The blots show the expression levels of monoubiquitin (A) and apoptotic proteins (p53 and Bax) and antiapoptotic proteins (Bcl-2 and Bcl-xL) (B). Blots were reprobed for α -tubulin, which was used to normalize the protein load. Images are representative of four independent experiments.

the ubiquitin staining in these epididymal regions was less intense (Fig. 3B). Immunoreactivity to both UCH-L1 and ubiquitin was intense in the caput epididymal epithelial cells, which was consistent with the expression level (Fig. 2 and Fig. 4A). Diffuse cytoplasmic immunoreactivity in the epididymal epithelial cells to UCH-L3 was intense in the cauda epididymis (Fig. 3C). As shown previously [24], no UCH-L1 or UCH-L3 immunoreactivity was found in the epididymal epithelial cells of *gad* and *Uchl3* knockout mice, respectively (Fig. 3A, C. inset).

Region-specific localization of ubiquitin and apoptotic proteins in the caput epididymis

We previously reported that UCH-L1 binds ubiquitin, and that the level of ubiquitin is decreased in *gad* mice [25, 31]. To determine whether UCH-L1 is associated with the ubiquitin level in the epididymis, we performed western blot analysis of the individual epididymal regions. The monoubiquitin level was markedly higher in the caput epididymis than in the corpus and cauda epididymis, and the low level of monoubiquitin in *gad* mice is consistent with our previous report [25] (Fig. 4A). The epididymis of *Uchl3* knockout mice did not show a difference in ubiquitin level compared with the corresponding wild-type controls.

To explore whether apoptotic phenomenon of spermatozoa in the caput epididymis is in accord with the high expression of apoptotic proteins, we used western blot analysis to verify the expression levels of p53 and Bcl-2 family proteins, which are associated with cell death [12, 28, 29]. The levels of p53 and Bax protein, considered to be proapoptotic, were strikingly high in the caput epididymis, consistent with the pattern of the monoubiquitin level (Fig. 4B). In the *gad* mouse, the levels of the antiapoptotic proteins, Bcl-2 and Bcl-xL, were markedly elevated compared in wild-type mice in the caput epididymis [23] as well as a possible increase in the corpus and cauda epididymis, whereas the levels of apoptotic proteins, p53 and Bax, were unchanged (Fig. 4B). However, we did not detect a difference in the analyzed protein levels between the epididymis of *Uchl3* knockout and wild-type mice.

Region-specific apoptosis in the epididymis following unilateral efferent duct ligation

Androgen deprivation by efferent duct ligation induces glandular epithelial cell death via an apoptotic mechanism [9, 38]. We previously showed that germ cell apoptosis differs between *gad* and *Uchl3* knockout mice following cryptorchid injury [25]. To detect apoptosis in the epididymis following efferent duct li-

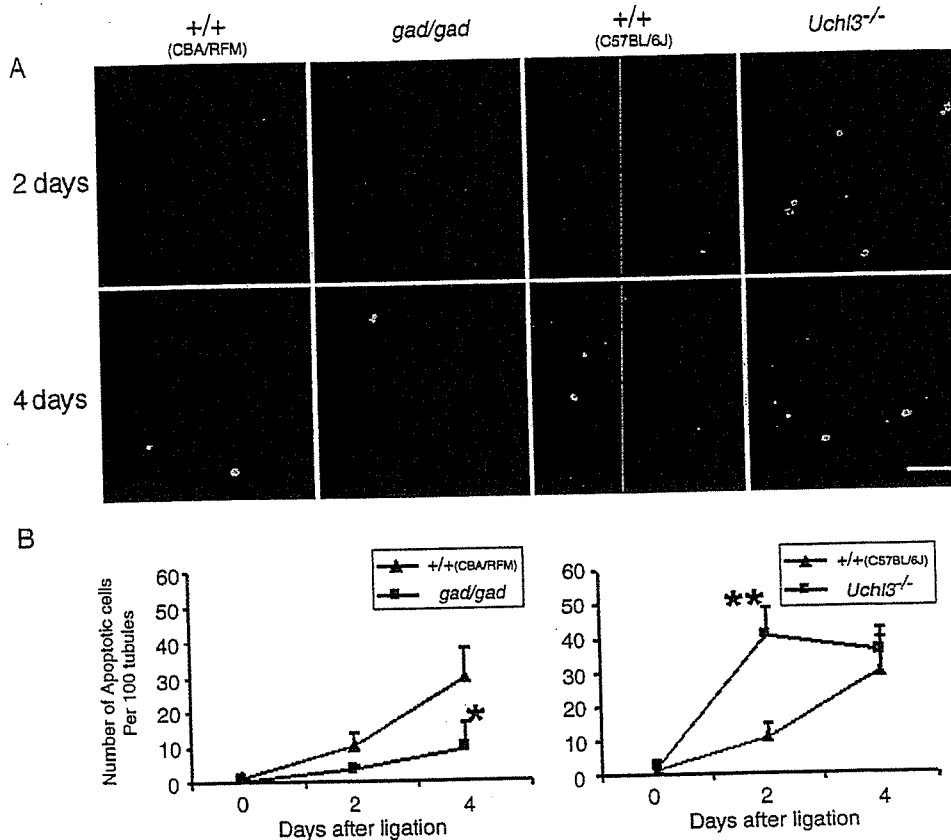


Fig. 5. TUNEL staining of apoptotic cells in the caput epididymis following unilateral efferent duct ligation. A: TUNEL staining in the caput epididymis cross-sections on days 2 and 4 after ligation. Green fluorescence, TUNEL-positive cells; red fluorescence, nuclei stained with propidium iodide. Magnification, 200 \times . Scale bar, 30 μ m. B: Quantitation of epithelial cell apoptosis in the caput epididymis following efferent duct ligation. The number of apoptotic epithelial cells in *gad* and wild-type mice is shown on the left. Each value represents the mean \pm SD; * P <0.05. The number of apoptotic epithelial cells in *Uchl3* knockout and wild-type mice is shown on the right. Each value represents the mean \pm SD; ** P <0.01.

gation, we used an *in situ* TUNEL assay to examine apoptosis in *gad* and *Uchl3* knockout mice on postoperative days 2 and 4 (Fig. 5). After efferent duct ligation, epithelial cell apoptosis was observed only in the caput epididymis (mostly the initial segment). The caput epididymis showed a time-dependent increase in epithelial cell apoptosis after efferent duct ligation and epithelial cell apoptosis was found mainly in the principal cells (Fig. 5A). Compared with wild-type mice, the caput epididymis of *Uchl3* knockout mice showed a marked increase in apoptotic epithelial cells on postoperative day 2, whereas *gad* mice resisted efferent duct-ligated epithelial cell apoptosis (Fig. 5A). By postoperative day 2, the number of apoptotic cells per 100

tubules increased with statistical significance (** P <0.01, n =4) in the caput epididymis of *Uchl3* knockout mice as compared with wild-type mice (Fig. 5B). However, *gad* mice showed resistance to ligation-induced apoptosis in the caput epididymis relative to wild-type mice by postoperative day 4 (* P <0.05, n =4) (Fig. 5B).

Discussion

After leaving the testis via the testicular rete, spermatozoa collect in the epididymis, where they undergo final maturation and storage [2, 36, 37]. During epididymal passage, ubiquitination may trigger apoptotic mechanisms that recognize and eliminate abnormal sper-

matozoa, and ubiquitination is believed to play an important role in controlling sperm quality to ensure the production of intact, functional spermatozoa [10, 27, 37]. Ubiquitination of abnormal spermatozoa predominantly occurs in the caput epididymis [37].

Previous studies have shown that two closely-related UCH isozymes, UCH-L1 and UCH-L3 have distinct expression patterns during spermatogenesis [24] and reciprocal functions following cryptorchid injury [25]. We have proposed that UCH-L1 might function as a regulator of apoptosis. Indeed, UCH-L1-deficient *gad* mice are resistant to apoptotic stress [13, 23, 25], and this apoptotic resistance leads to alterations in sperm motility and morphology as well as an increased number of defective spermatozoa in the epididymis of *gad* mice [23]. Our present study demonstrated that UCH-L1 and UCH-L3 have distinct expression patterns along the epididymis in wild-type mice (Fig. 2). We detected a high level of UCH-L1 in the caput epididymis, the main maturation organ, whereas the UCH-L3 level was high in the cauda epididymis, the main storage organ [10]. These region-specific variations in UCH-L1 and UCH-L3 level suggest that they have different functions in the epididymis. The regional differentiation of the epididymis, as suggested by region-specific gene expression, reflects different luminal environments between the regions [16, 19].

We also determined the expression pattern/level of the major component of the proteolytic pathway, ubiquitin, which has specificity for UCH-L1. UCH-L1 associates with monoubiquitin [31], and the monoubiquitin level is reduced in *gad* mice relative to wild-type mice [25, 31]. Predictably, monoubiquitin expression pattern showed similar patterns to UCH-L1 and the monoubiquitin level was reduced in the epididymis of *gad* mice, which had its highest level in the caput epididymis relative to the corpus or cauda epididymis in wild-type mice (Figs. 3 and 4A). Ubiquitin induction is important for regulating programmed cell death, which is a fundamental component of spermatogenesis [1, 23, 32]. Under specific circumstances, the caput epididymis contains a high level of ubiquitin, which may serve to maintain apoptotic mechanisms that eliminate abnormal spermatozoa [37]. This is consistent with the high levels of apoptotic p53 and Bax observed in the caput epididymis compared with the corpus and cauda epididymis (Fig. 4B). Protein p53

and Bax are classically thought to be involved in regulating apoptotic processes, and are targets for ubiquitination [5, 7, 29, 30]. The role of p53 in mediating apoptosis in the male genital tract has been demonstrated in several mice lines [28, 29, 42]. However, p53-independent apoptosis is suggested in the prostate and seminal vesicles by androgen withdrawal or in the rat epididymis by deprivation of luminal factors [3, 11, 14, 38]. Previous studies indicated that Bcl-2 family proteins are involved in the induction or prevention of apoptosis [12, 33, 39, 40]. In *gad* mice, in the present study, the levels of the antiapoptotic proteins, Bcl-2 and Bcl-xL, were markedly increased in the caput epididymis (Fig. 4B), although there was no difference in the levels of the apoptotic proteins, p53 and Bax, relative to wild-type mice. The high levels of Bcl-2 and Bcl-xL in the caput epididymis of *gad* mice is consistent with a previous report that the percentage of morphologically abnormal spermatozoa is significantly higher in *gad* mice [23]. Therefore, the variations of in the levels of Bax, and Bcl-2 and Bcl-xL combined in the caput epididymis probably maintain the regulation of apoptosis [4].

Our previous work focused on the reciprocal functions that UCH-L1 and UCH-L3 exhibit, a distinct feature in testicular germ cells following cryptorchid-induced apoptosis [25]. To characterize the distinct functions of UCH-L1 and UCH-L3 in the epididymis, *gad* and *Uchl3* knockout mice were examined after efferent duct ligation. The epididymal epithelium of the two mutant mice showed differences in apoptotic induction following efferent duct ligation (Fig. 5), after which the circulating androgen level decreases rapidly as a result of apoptotic cell death [9, 20, 38]. After duct ligation, the number of apoptotic cells increased in the caput epididymis of *Uchl3* knockout mice compared with wild-type mice, whereas *gad* mice showed relative resistance in this regard (Fig. 5B). In *gad* mice, the resistance to apoptotic stress can be explained by the high levels of Bcl-2 and Bcl-xL combined in the caput epididymis (Fig. 4B). The tissue androgen level is higher in the caput epididymis than in the corpus or cauda epididymis [15, 38]; thus, apoptotic cells showed in the caput epididymis rather than in the corpus and cauda epididymis following efferent duct ligation. These results may suggest that UCH-L1 and UCH-L3 have reciprocal functions in the caput epididymis fol-

lowing apoptotic stress induced by androgen withdrawal, as was shown with cryptorchid stress [25].

We cannot explain the profound apoptotic phenomenon observed in the present study in the caput epididymis of *Uchl3* knockout mice after efferent duct ligation by the balance of the Bcl-2 family proteins alone. Although our previous report showed that the Nedd8 expression level increased in the testis of *Uchl3* knockout mice [25], we found no difference in the present study (data not shown). The mechanism with regard to the antiapoptotic function of UCH-L3 requires further study. Our present study demonstrated that UCH-L1 and UCH-L3 have distinct expression levels along the epididymis as well as reciprocal functions in response to apoptotic stress induced by androgen withdrawal.

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Three-dimensional reconstruction of the membrane skeleton at the plasma membrane interface by electron tomography

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Three-dimensional images of the undercoat structure on the cytoplasmic surface of the upper cell membrane of normal rat kidney fibroblast (NRK) cells and fetal rat skin keratinocytes were reconstructed by electron tomography, with 0.85-nm-thick consecutive sections made ~100 nm from the cytoplasmic surface using rapidly frozen, deeply etched, platinum-replicated plasma membranes. The membrane skeleton (MSK) primarily consists of actin filaments and associated proteins. The MSK covers the entire cytoplasmic surface and is closely linked to clathrin-coated pits and caveolae. The actin filaments

that are closely apposed to the cytoplasmic surface of the plasma membrane (within 10.2 nm) are likely to form the boundaries of the membrane compartments responsible for the temporary confinement of membrane molecules, thus partitioning the plasma membrane with regard to their lateral diffusion. The distribution of the MSK mesh size as determined by electron tomography and that of the compartment size as determined from high speed single-particle tracking of phospholipid diffusion agree well in both cell types, supporting the MSK fence and MSK-anchored protein picket models.

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Introduction

The portion of the cytoskeleton that is closely associated with the cytoplasmic surface of the plasma membrane is often called the membrane skeleton (MSK; Heuser and Kirschner, 1980; Hirokawa and Heuser, 1981; Bennett, 1990; Luna and Hitt, 1992). The term MSK is useful partly because this part of the cytoskeleton is expected to differ from the bulk cytoskeleton in terms of its structure and protein composition, for its interactions with the plasma membrane in general and with specific molecules in the plasma membrane, and also because it plays important roles in a variety of membrane functions. It is involved in the localization of transmembrane proteins at specific sites in the cell membrane (Bennett and Chen, 2001; Pan et al., 2006) and in endocytosis and exocytosis (Gaidarov et al., 1999; Valentijn et al., 2000) in various cell types. It also provides the plasma membrane with the mechanical strength and resilience to withstand the stress and shear forces from the outside environment,

which is well established in the thick cortical actin layers in immune cells (Hartwig and Yin, 1988) and in the spectrin-actin network in red blood cells (Mohandas and Evans, 1994; Discher et al., 1995). Therefore, the MSK works as a part of the plasma membrane as well as a part of the cytoskeleton. It is a truly interfacial structure between the bulk cytoskeleton and the 2D bilayer of the plasma membrane.

Recently, a new function of the MSK has become apparent. It was proposed that a part of the MSK is directly and closely associated with the cytoplasmic surface of the plasma membrane, and this part induces partitioning of the cell membrane with regard to the translational diffusion of membrane molecules based on high speed single-particle tracking data on membrane proteins and lipids (Jacobson et al., 1995; Kusumi et al., 2005). In the short-time regime, these membrane molecules are temporarily confined within the compartments delimited by the MSK mesh, and, in the long-time regime, they undergo macroscopic diffusion by hopping between these compartments (MSK fence model). In the fence model, as a result of the collision of the cytoplasmic domains of transmembrane

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Abbreviations used in this paper: CCP, clathrin-coated pit; FRSK, fetal rat skin keratinocyte; MSK, membrane skeleton; NRK, normal rat kidney fibroblast.

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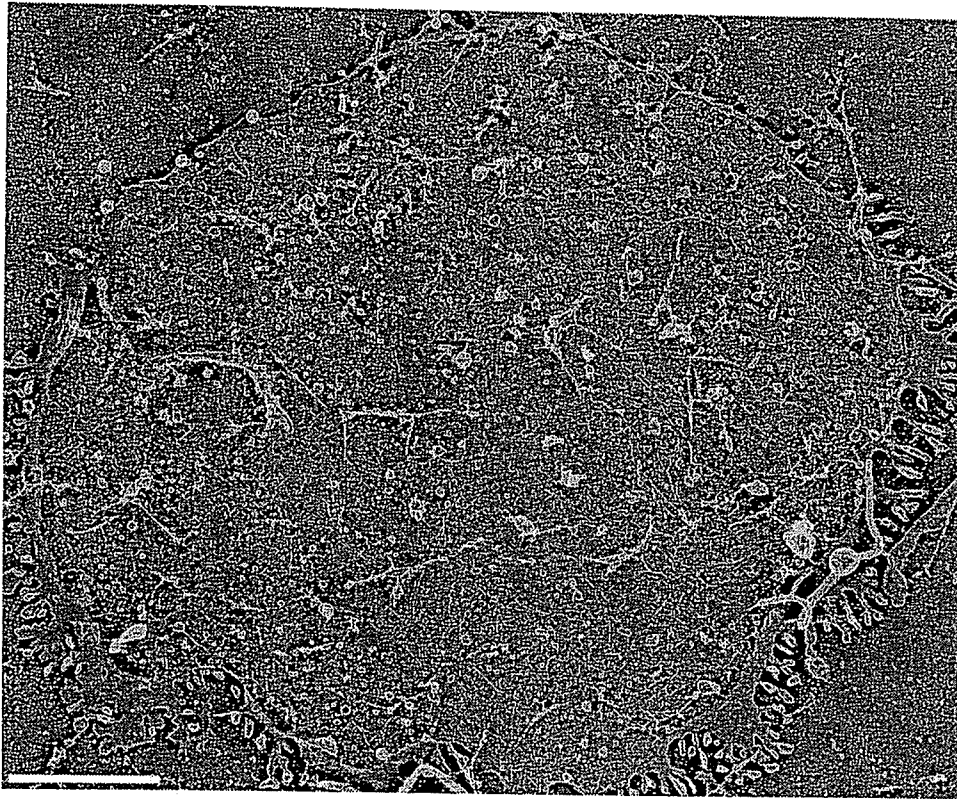


Figure 1. A bird's-eye view of the large cytoplasmic surface of the upper cell membrane (the membrane facing the buffer rather than the coverslip) of an NRK cell observed by rapid-freeze, deep-etch, freeze-replica EM. Bar, 2.5 μm .

proteins with the MSK, transmembrane proteins are temporarily confined in the MSK mesh (Sheetz, 1983; Tsuji and Ohnishi, 1986; Tsuji et al., 1988; Saxton, 1989, 1990; Sako and Kusumi, 1994, 1995; Jacobson et al., 1995; Kusumi and Sako, 1996; Saxton and Jacobson, 1997; Sako et al., 1998; Tomishige et al., 1998; Suzuki et al., 2005).

Lipid molecules also undergo hop diffusion, which might be explained by the anchored protein picket model (Fujiwara et al., 2002; Murase et al., 2004; Kusumi et al., 2005). In this model, various transmembrane proteins anchored to the actin-based MSK might effectively act as rows of pickets against the free diffusion of all of the molecules incorporated in the cell membrane as a result of steric hindrance and circumferential slowing (a hydrodynamic frictionlike effect, which propagates quite far from the immobile protein surface; without this effect, pickets will not be effective for blocking diffusion; Bussell et al., 1994, 1995) of the immobile picket proteins anchored to and lined up along the MSK. Lipid movement is affected only by pickets, whereas both pickets and fences would act on transmembrane proteins. These MSK picket-fence effects would be dramatically enhanced when the membrane receptor molecules form signaling complexes upon ligand binding as a result of receptor oligomerization and/or binding of the cytoplasmic signaling molecules to the receptor, leading to the trapping of signaling complexes in the MSK mesh, where the extracellular signal is received. This would enable spatial confinement and

regulation of the downstream signaling events (Kusumi and Sako, 1996; Iino et al., 2001).

Despite the importance of the MSK functions and the long history of its study using EM (Byers and Porter, 1977; Heuser and Kirschner, 1980; Hirokawa and Heuser, 1981; Heuser and Anderson, 1989; Hartwig and DeSisto, 1991; Rothberg et al., 1992), our knowledge of its structure and the overall distribution over the plasma membrane has been very limited. For example, we do not know whether the MSK exists everywhere on the cytoplasmic surface of the cell membrane, how extensive the spatial variations of MSK mesh size is, and whether and how MSK interacts with other structures in the plasma membrane such as clathrin-coated pits (CCPs), caveolae, and cell adhesion structures. Even the structure of the MSK of the human red blood cell ghost, a traditional paradigm for MSK studies, is not satisfactorily understood (Sheetz and Sawyer, 1978; Tsukita et al., 1980; Branton et al., 1981; Shen et al., 1986; Ursitti et al., 1991; Takeuchi et al., 1998).

In this study, to further advance our understanding of the MSK structure and function, we observed the undercoat structure on the cytoplasmic surface of the plasma membrane of cultured mammalian cells using rapid-freeze, deep-etch, immunoreplication EM. We paid special attention to the following three points.

First, we tried to consistently prepare and observe large plasma membrane fragments ($>10 \mu\text{m}$ in diameter) to facilitate inspections of very large plasma membrane areas. Almost all of

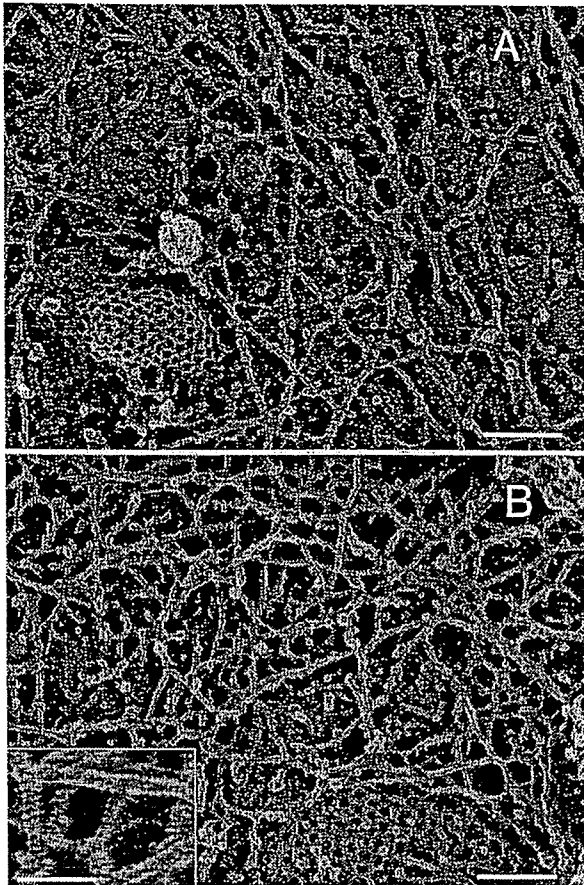


Figure 2. Magnified MSK images of an NRK and FRSK cell on the cytoplasmic surface of the upper membrane. (A) NRK cell; (B) FRSK cell. Clathrin-coated structures (A and B) and a caveola (A) show the cytoplasmic surface. The striped banding patterns with the 5.5-nm periodicity on individual filaments are characteristic of actin filaments. These images also reveal close links of the MSK actin filaments with the clathrin-coated structures and caveolae. Bars (A and B), 100 nm; [inset] 50 nm.

the previous MSK studies, including those cited above, investigated the ultrastructural features of the structure of interest, but within a very limited view field. By observing these large membrane surfaces, the spatial variations of the MSK mesh size and of the number density of CCPs and caveolae can be reliably examined.

Second, the 3D reconstruction of the undercoat structure within 100 nm from the cytoplasmic surface of the plasma membrane was performed using electron tomography for the platinum-replicated samples: 97–141 images for a specimen tilted at different angles (every 1°) with respect to the incident electron beam in the range of ± 48 – 70° were obtained and then converted to 100–121 sliced images of every 0.85–1.34 nm for the 3D reconstructed images (Perkins et al., 1997; Medalia et al., 2002; Lucic et al., 2005; McIntosh et al., 2005; Zeuschner et al., 2006).

Third, using the 3D reconstructed images of the MSK structure within 13.6 nm (16 slices of 0.85-nm thickness) from the cytoplasmic surface, the MSK mesh size distribution on the

cytoplasmic surface of the plasma membrane was determined. This part of the MSK, which is closely associated with the cytoplasmic surface of the plasma membrane, might form the compartment boundaries for partitioning of the plasma membrane for the diffusion of membrane molecules, thus determining the compartment size. Therefore, it is interesting to compare the distribution of the MSK mesh size on the membrane determined this way and that of the compartment size sensed by membrane molecules. Because the compartment size distributions for membrane molecules are very different between normal rat kidney fibroblast (NRK; median = 230 nm) and fetal rat skin keratinocyte (FRSK; median = 41 nm) cell lines (Fujiwara et al., 2002; Murase et al., 2004), the distribution of the MSK mesh size on the membrane surface was examined using these two cell lines. Although the compartment size is very different between these cell lines, within each cell type, the histogram for the MSK mesh size on the membrane surface is very similar to that for the diffusion compartment size. This strongly supports the MSK fence and MSK-anchored transmembrane protein picket models.

Results

Bird's-eye view of the undercoat structure of the upper cell membrane

Glass coverslips precatenized by a treatment with Alcian blue were placed on top of the cells cultured in 35-mm plastic dishes and were allowed to settle and attach to the upper cell membrane at 4°C for 15 min. The buffer containing 1% PFA and 0.25% glutaraldehyde was then added to the space between the coverslip and the plastic bottom of the culture dish. As the coverslip was floated apart, the cells were ruptured and the upper cell membrane was retained, still adhering to the overlaying Alcian blue-coated coverslip. The upper membrane was rapidly frozen by pressing its exposed cytoplasmic surface onto a pure copper block precooled by liquid helium. The frozen sample was deep etched, coated with platinum-carbon, and observed under an electron microscope. We have made extensive efforts to reproducibly prepare and observe large cell membrane fragments $>10 \mu\text{m}$ in diameter.

Fig. 1 is a typical electron micrograph providing a bird's-eye view of the cytoplasmic surface of a large area of the upper cell membrane of a cultured NRK cell. Many such EM images showing the cytoplasmic surfaces of large cell membrane fragments were obtained for NRK and FRSK cells, suggesting that the entire (upper) plasma membrane, except for the places where CCPs and caveolae exist, is coated with the filamentous netlike structure.

Fig. 2 (A and B), which was obtained for an NRK cell (Fig. 2 A) and an FRSK cell (Fig. 2 B), shows the magnified images of the cytoplasmic surface of the plasma membrane, exhibiting extensive filamentous netlike structures, which are the MSK. The presence of clathrin-coated structures shows that this is indeed the cytoplasmic surface. The striped banding patterns with a 5.5-nm periodicity on individual filaments are characteristic of actin filaments and, thus, indicate that these are actin filaments (Heuser and Kirschner, 1980; Heuser, 1983; Katayama,

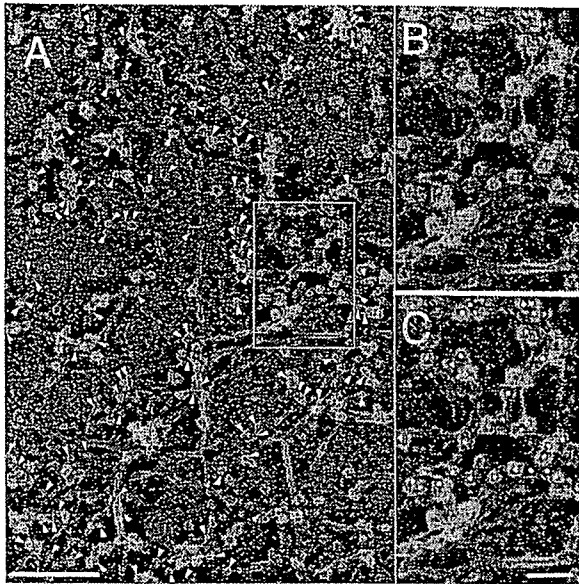


Figure 3. Immunogold labeling also indicates that the major component of MSK is actin filaments (NRK cell). Actin filaments were indirectly immunolabeled with 5-nm colloidal gold particles coated with secondary antibodies. Each gold particle can be identified as a clear white spot (colloidal gold particle) surrounded by a fuzzy gray ring, which is caused by the platinum rotary shadowing around the secondary antibody coating of the gold particle. Representative probe images are indicated by arrowheads. (A) Most of the filamentous structures are labeled by colloidal gold probes. (B and C) The boxed area in A is shown at a higher magnification. In C, the gold particles are marked by yellow dots. The filaments with the 5.5-nm striped banding pattern, which is characteristic of the actin filament, are labeled with these gold probes. Bars (A), 200 nm; (B and C) 50 nm.

1998; Schoenenberger et al., 1999). Because almost all of these filaments contain this striped pattern, it is concluded that the MSK is predominantly composed of actin filaments. This was also confirmed by immunogold staining (see Fig. 3 and related text).

The electron micrograph shown in the inset in Fig. 2 B indicates the spatial resolution: because each band in the striped pattern with a 5.5-nm periodicity is visibly separated, the effective resolution is thought to be ~ 2 nm (both the thickness of the platinum coating and the platinum granule size are ≤ 2 nm; Heuser and Kirschner, 1980; Heuser, 1983). The MSK structure observed here on the upper cell membrane is similar to that on the bottom cell membrane (the part of the cell membrane facing the coverslip) observed previously (Heuser and Anderson, 1989).

These results suggest that the cytoplasmic surface of a portion of the upper cell membrane $>10 \mu\text{m}$ in diameter was visualized with a spatial resolution of ~ 2 nm, which is much smaller than the width of a single actin filament or the repeat distance of the stripes. As shown in Figs. 1 and 2 (A and B), the MSK is likely to cover the entire cytoplasmic surface of the upper cell membrane except for the places where CCPs and caveolae are present in both NRK and FRSK cells. Such a notion of the complete coverage of the cytoplasmic surface of the plasma membrane by actin filaments might have existed for >30 yr in a part of the EM community (Byers and Porter, 1977; for review

see Sheetz et al., 2006), but the data specifically indicating that the actin filaments of the MSK may cover the entire cell membrane has not been presented in the literature, as done here, nor shared in the cell biology community. The EM observations shown in this study are consistent with the MSK fence and anchored transmembrane protein picket models, in which the entire plasma membrane except for the specific membrane domains is partitioned into many small compartments with regard to lateral diffusion of the molecules incorporated in the plasma membrane.

The MSK predominantly consists of actin filaments: immunogold labeling of actin and actin-binding proteins

To further examine whether the MSK is predominantly composed of actin filaments (and partly because the 5.5-nm periodicity of the banding pattern is somewhat difficult to discern in some of the filaments), we examined it using an indirect immunolabeling method with 5-nm-diameter colloidal gold particles (see Materials and methods; Fig. 3). On the filaments with striped patterns, the enlarged images (Fig. 3, B and C) show the presence of many colloidal gold actin probes, which appear as distinct white spots surrounded by somewhat blurred white halos, reflecting the platinum shadow over the antibody molecules attached to the gold particle. The electron micrographs in Fig. 3 revealed that almost all of the colloidal gold probes are bound to the filaments located on the cytoplasmic surface (yellow dots). Therefore, it is concluded that actin is the main constituent molecule of the MSK.

Electron tomography of the undercoat structure on the cytoplasmic surface of the plasma membrane

The 3D structure of the undercoat within 100–134 nm from the cytoplasmic surface of the plasma membrane, which includes CCPs, caveolae, and the actin-based MSK, was reconstructed using electron tomography for the platinum-replicated samples. Based on the 97–141 tilt images acquired in the range of ± 48 – 70° every 1° step for a single EM view field, 100–121 sliced images of every 0.85–1.34 nm perpendicular to the z axis (parallel to the image obtained at 0° of the tilt angle) were calculated by a computer (long wavelength [$\geq \sim 500$ nm] undulations of the cell membrane were corrected by the 3D reconstruction software IMOD). The 3D image was reconstructed based on these serial thin slices. Representative images obtained for an EM view field are shown in Video 1 (131 tilt images; an anaglyph produced from images taken at $\pm 12^\circ$ is shown in Fig. 4 A) and Video 2 (showing the 3D image by rotating the 3D reconstructed undercoat structure; a typical view is shown in Fig. 4 B; videos are available at <http://www.jcb.org/cgi/content/full/jcb.200606007/DC1>). Throughout the present research, this protocol was used to obtain 3D images.

In these images, because of their 3D representation, it is especially clear that the MSK, which is mostly composed of actin filaments, generally spreads along the membrane, covering almost the entire cytoplasmic surface of the upper membrane except for the places with caveolae and CCPs. In addition,

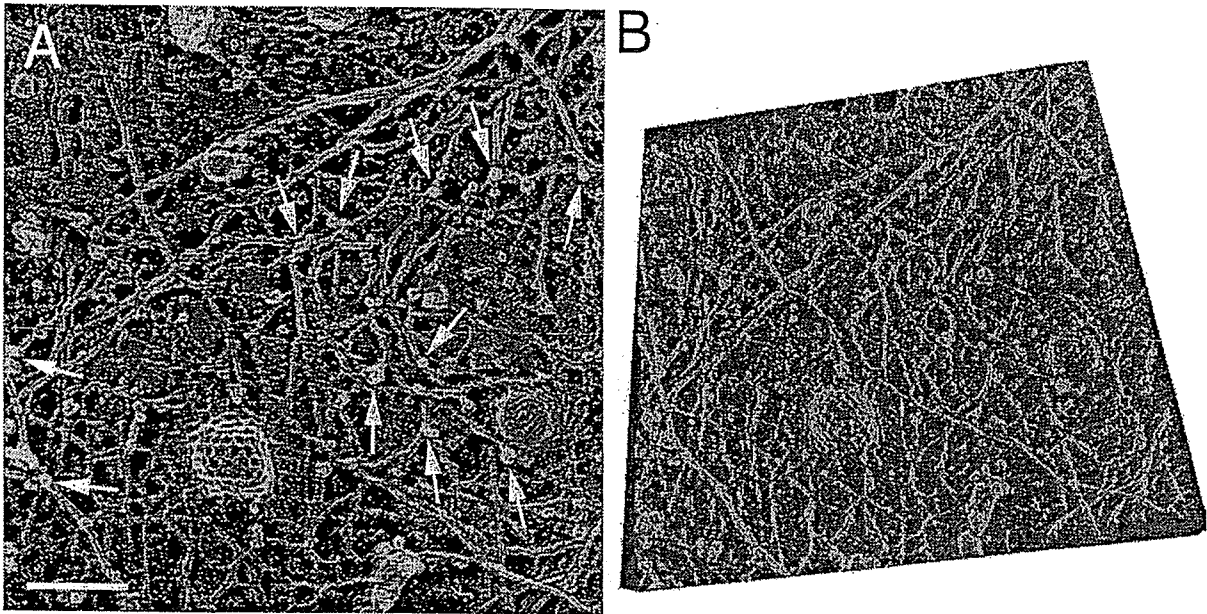


Figure 4. Stereo electron micrographs and 3D reconstructed images of the undercoat structure, CCPs, and caveolae in NRK cells. (A) An EM anaglyph of the undercoat structure generated at $\pm 12^\circ$ of the tilt angle among the 131 tilt images (acquired in the range of $\pm 65^\circ$ with 1° steps). Use view glasses for the 3D structure (left = red). See Video 1 for all 131 of the tilt images. Arrows indicate actin filaments protruding from the membrane cytoplasmic surface toward the cytoplasm. The arrows point to the places where the protruding actin filaments intersect with the MSK meshwork located close to the membrane. (B) The 3D undercoat structure reconstructed from the tilt images shown in Video 1. See Video 2, where the 3D structure is rotated (available at <http://www.jcb.org/cgi/content/full/jcb.200606007/DC1>). Bar, 100 nm.

CCPs and caveolae are very closely associated with the actin filaments in the MSK, as seen in these images and also in Figs. 2 (A and B) and 3. These results are consistent with Rothberg et al. (1992), Fujimoto et al. (2000), and Parton (2003), but in NRK cells studied here, many more actin filaments were found to be associated with each CCP or caveola. Furthermore, 92 and 93% of CCPs and caveolae ($n = 200$) were bound by the actin filaments. These results are consistent with the requirement of filamentous actin for CCP internalization (Qualmann et al., 2000; Merrifield et al., 2002).

Many short, thin filaments protrude toward the cytoplasm, mostly perpendicularly, from the membrane surface (they were short probably because they were broken at the time of the membrane rip off; Fig. 4 A, arrows). Note that these perpendicular filaments are almost always connected to the MSK network lying on the cytoplasmic surface (see the tips of the arrows; Fig. 4 A). Thus, the part of the MSK that is located on the cytoplasmic surface is connected three dimensionally to the cytoskeleton. Together, they will provide mechanical support for the membrane and the force for deforming the membrane.

3D reconstruction of the MSK structure

The part of the actin-based MSK that is in contact with the cytoplasmic surface of the cell membrane has been proposed to partition the cell membrane into 30–230-nm compartments by the fence and picket effect (Edidin et al., 1991; Kusumi and Sako, 1996; Kusumi et al., 2005). If these fence and picket models are correct, the distribution of the mesh size of the MSK on the cytoplasmic surface of the plasma membrane would be

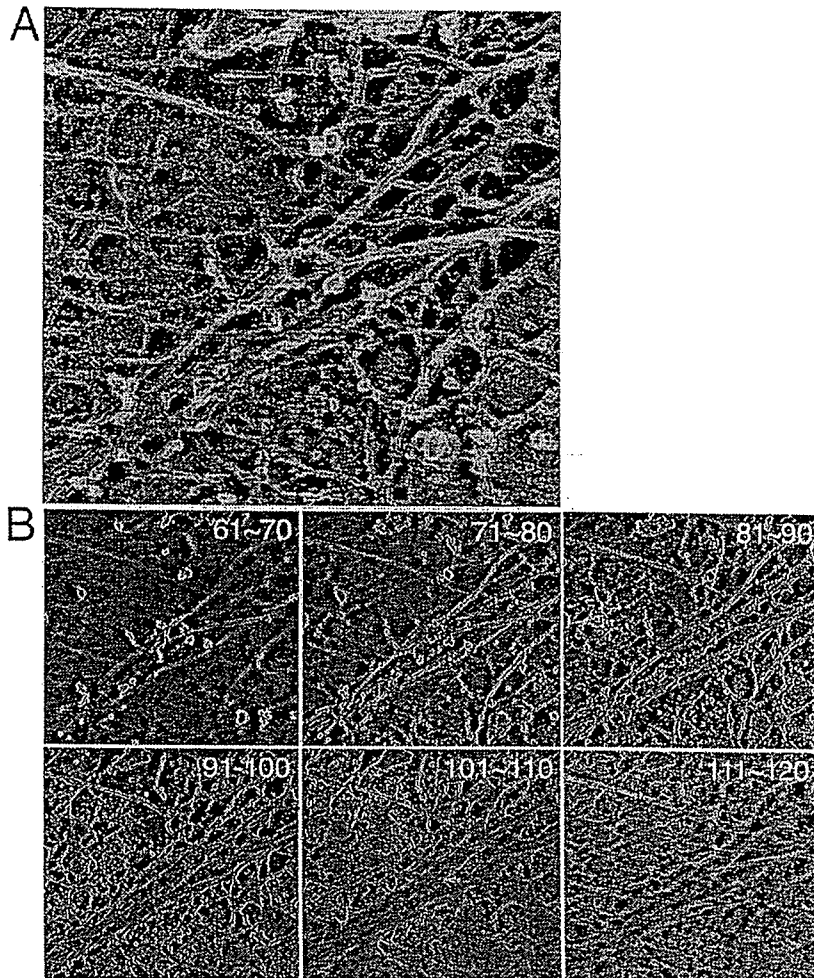
practically the same as that of the compartment size determined by diffusion measurements of membrane molecules. To carry out this examination, the 3D reconstruction of MSK by electron tomography provides a unique opportunity because the obtained images provide quantitative data on how far the individual filaments are located from the membrane surface.

In Fig. 5 A, a typical MSK structure quantitatively analyzed in this study is shown in an anaglyph, and its 8.5-nm-thick sections (created by superimposing 10 0.85-nm sections) of the MSK of an NRK cell, starting from the cytoplasmic side toward the membrane, are shown (Fig. 5 B; a series of the original tilt images is shown in Video 3, and a series of sliced images of every 0.85 nm is shown in Video 4, available at <http://www.jcb.org/cgi/content/full/jcb.200606007/DC1>). The actin-based MSK is visible on image sections 81–110. Individual actin filaments, forming a network as well as bundles, can be identified. Given the high density of the actin filament meshwork, which is much smaller than the optical resolution, conventional fluorescence microscopy will be unable to visualize the MSK meshwork and can visualize only the bundles of actin filaments.

Interface structure of MSK on the cytoplasmic surface of the plasma membrane

The filaments of the MSK that are directly associated with the cytoplasmic surface of the plasma membrane and may be involved in partitioning the plasma membrane were systematically determined. Out of the stack of 121 image slices taken every 0.85 nm from the cytoplasmic surface (~ 100 -nm thick altogether),

Figure 5. A series of sliced images of the actin MSK on the plasma membrane cytoplasmic surface of an NRK cell. (A) A typical actin MSK structure used for analysis of the mesh size on the cytoplasmic surface of the plasma membrane using computed tomography. An anaglyph of the undercoat structure generated at $\pm 12^\circ$ of the tilt angle among the 97 tilt images (acquired in the range of $\pm 48^\circ$ with 1° steps). See Video 3 for all 97 of the tilt images. (B) 10 consecutive sections, each 0.85-nm thick, are superimposed, and six of these superimposed images, which represent 60 image sections out of 121 image sections, are shown from the cytoplasmic side toward the plasma membrane side. The numbers here indicate the number of slices counted from the cytoplasmic side. The actin-based MSK near the cytoplasmic surface of the plasma membrane is visible on images 81–110. All 121 of the sliced images of every 0.85 nm are shown in Video 4 (available at <http://www.jcb.org/cgi/content/full/jcb.200606007/DC1>).



16 consecutive image slices from the membrane surface (~ 13.6 -nm thick altogether) were used for this analysis (Fig. 6, A and B).

In Fig. 6 A (four images on the right) and Fig. 6 B (the second to fourth images), the boxed areas in the left-most images were expanded, and the sections of every 1.7 nm (superposition of two 0.85-nm-thick slices; 330×330 nm) are displayed between 0 and 11.9 nm. Using these sections, the filaments that are closely associated with the cytoplasmic surface of the cell membrane were determined. Because the thickness (width in the image) of the actin filament after platinum shadowing is between 9 and 11 nm (consistent with Heuser, 1983) and the thickness of the platinum replica is ≤ 2 nm (consistent with Heuser, 1983 and Moritz et al., 2000), the height of the actin filament that is associated with the membrane will be 7–9 nm (because the height is given by the actin thickness and one replica thickness, whereas the width in the image is determined by the actin thickness plus two replica thicknesses), with 8 nm being a reasonable estimate. In the series of electron tomography sections shown in Fig. 6 (A and B), the existence of three major classes of filaments with regard to the distance from

the membrane surface can be discerned (for details of this analysis, see Materials and methods).

The first class of filaments is distinct in computer-reconstructed sections close to the cytoplasmic surface of the plasma membrane, even in the first ~ 0 –1.7-nm section (because the contrast is reversed in these micrographs, they look more lucent or white), but fade out of the reconstructions 8–10 nm away from the membrane surface (for details, see Materials and methods). These filaments are drawn in green in Fig. 6 C. We interpret that these filaments are in true contact with the plasma membrane (the gap between the filament and the inner membrane surface is < 0.85 nm) because they can be seen clearly even in the first 0.85-nm section. These filaments are likely to be the significant ones for generating membrane corrals.

The second class of filaments also looks clear in sections very close to the membrane surface but does not fade out until ~ 14 nm away from it. We interpret that these may be the actin filaments that had platinum coatings all around their surfaces because they stood off the surface somewhat, which slightly exaggerated their thickness and made them appear as though they were in contact with the plasma membrane when

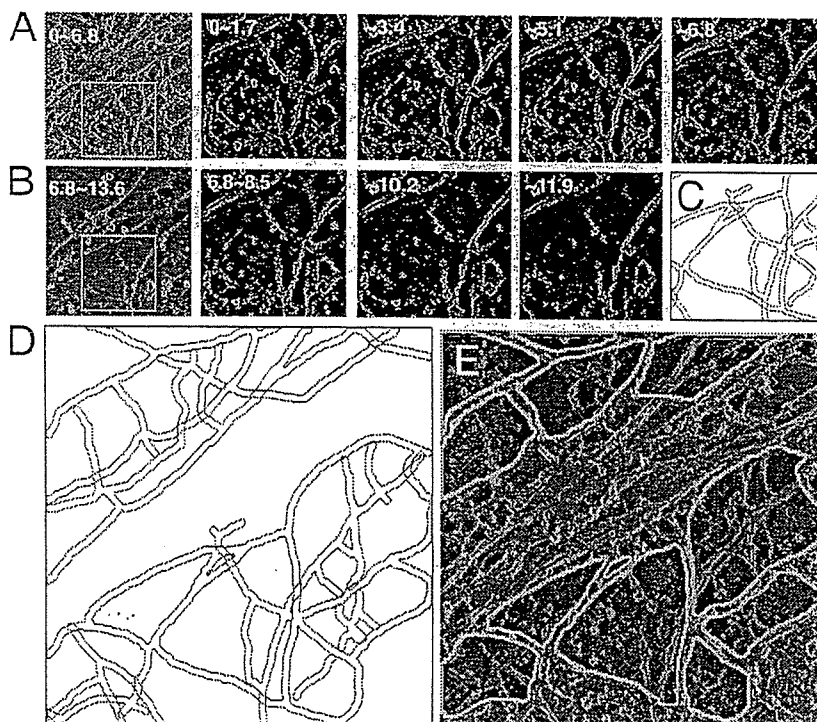


Figure 6. The method for determining the MSK mesh on the cytoplasmic surface of the plasma membrane, which possibly delimits the compartments of the plasma membrane, using the 3D reconstructed images of the MSK (an NRK cell). (A and B) The images on the far left are the ~ 0 –6.8- or ~ 6.8 –13.6-nm sections, each of which is a stack of eight 0.85-nm sections of 670×670 nm. These are from a series of 121 image sections (0.85-nm thick) from the cytoplasmic surface after the tilt and long wavelength undulation of the cell surface were corrected. The boxed areas in A and B (330×330 nm) are expanded on the right of these image stacks, with a section thickness of 1.7 nm (two 0.85-nm sections are superimposed; 330×330 nm for each image). (C) The outline of each actin filament adjacent to the membrane surface (green, which could not be observed above 10.2 nm) and that of each actin filament that could be observed above 10.2 nm (red). The same view field and magnification as those for the thinner sections shown in A and B (330×330 nm). See Materials and methods for details. (D) The outline of actin filaments in a greater view field, which is the same as those in the thick sections (~ 0 –6.8 and ~ 6.8 –13.6 nm) in A and B (670×670 nm, expanded here). (E) The image of the ~ 0 –6.8-nm sections (670×670 nm) superimposed by the image of areas surrounded by the filaments outlined in green in D (green areas with yellow outlines). According to the fence and picket models, these areas are likely to be the compartments where membrane molecules are temporarily confined.

in fact they probably were not quite in direct contact. We did not consider these filaments to be close enough to generate membrane corrals.

The third class of filaments is not apparent in sections closest to the plasma membrane but becomes clear some distance away from it (>2 –4 nm) and also does not fade out until ~ 14 nm. We interpret these as being filaments that definitely do not contact the plasma membrane directly and, thus, should not contribute to forming corrals. The second and third classes of filaments are drawn in red in Fig. 6 C.

Therefore, we considered that only the first class of filaments (those drawn in green in Fig. 6, C and D) forms the MSK fences and pickets, and the area surrounded by these filaments is colored green in the 0–6.8-nm section shown in Fig. 6 E. Note that areas are excluded from this analysis in which bundles of actin filaments are present (e.g., the structure crossing diagonally from the bottom left to the top right in Fig. 5), actin filaments are too crowded to be individually discerned, an actin filament is terminated in the middle of a domain (domains that contain a loose end of an actin filament), or CCPs, caveolae, and the smooth surface membrane invaginations are present (the white regions in Fig. 7 C).

Comparison of the MSK mesh size on the plasma membrane determined by electron tomography with the compartment size for the diffusion of membrane molecules
Similar determination of the MSK meshwork was performed for FRSK cells. Representative meshes of the MSK are shown

in Fig. 7 (for an FRSK cell, colored to aid in visualization). We performed such analyses for 10 representative stacks of image sections ($1,290 \times 1,290$ -nm plane) each for NRK cells and FRSK cells (eight different cell membrane sheets for each cell type) and identified 76 and 1,300 areas bounded by the MSK meshwork, respectively (excluding the regions occupied by stress fibers and other membrane undercoat structures such as CCPs and caveolae; about the same total membrane areas were examined for each cell type, and, thus, the difference in the number of identified areas represents the difference in the area size between these two cell lines). The 2D area size for each domain was measured by Amira software. The distributions of the square root of the area size (the side length, assuming a square shape for the area) for NRK (Fig. 8, pink open bars) and FRSK (blue open bars) cells are shown in Fig. 8. The median values of the area and its square root are 3.9×10^4 nm² and 200 nm, respectively, for NRK cells and 2.7×10^3 nm² and 52 nm, respectively, for FRSK cells.

The size distributions of the compartments for the diffusion of membrane molecules were obtained for an unsaturated phospholipid, L- α -dioleoylphosphatidylethanolamine, by Fujiwara et al. (2002) and Murase et al. (2004) for NRK and FRSK cells, respectively. The distributions of the side lengths for NRK (Fig. 8, pink closed bars) and FRSK (blue closed bars) cells are shown in the histograms in Fig. 8. The median values of the compartment area and the side length are 4.3×10^4 nm² and 230 nm, respectively, for NRK cells and 2.1×10^3 nm² and 41 nm, respectively, for FRSK cells (Murase et al., 2004).

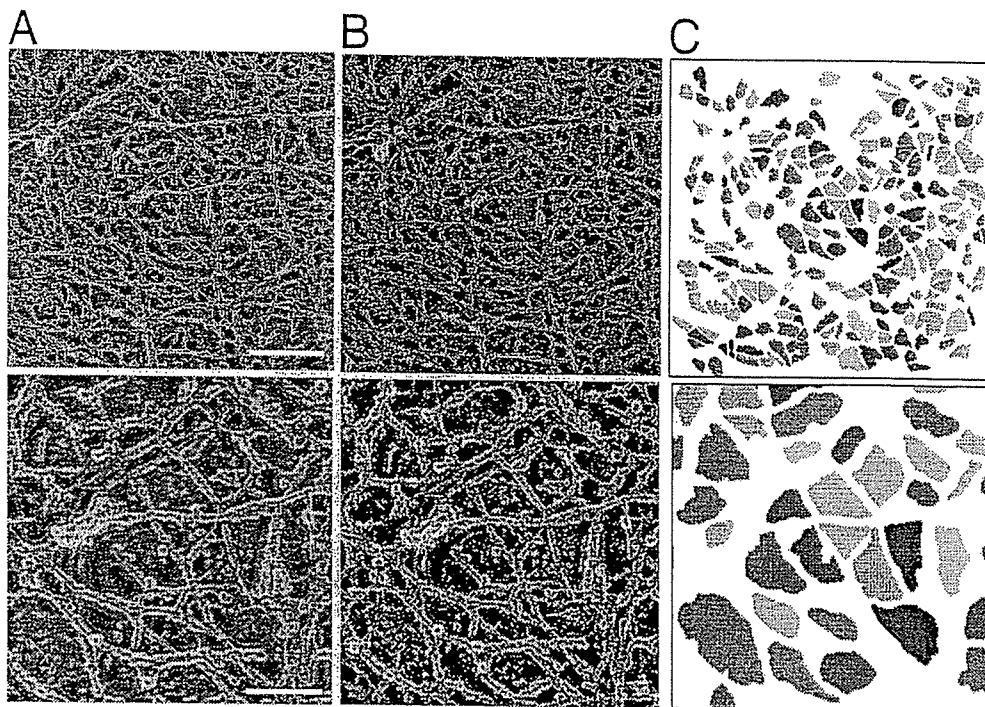


Figure 7. The MSK meshwork directly on the cytoplasmic surface of the plasma membrane. The central parts of the figures in the top row (bar, 300 nm) are magnified by a factor of three and are shown in the bottom row (bar, 100 nm). (A) Typical stereo views of the plasma membrane specimen (anaglyph; left = red). (B) Normal electron micrographs of the plasma membrane samples. The same view fields as those in A. (C) The areas delimited by the actin filaments closely apposed to the cytoplasmic surface of the cell membrane are shown. Different colors are shown to help the eye.

These results indicate that in the same cell line (for both the NRK and FRSK cases), the MSK mesh size determined by electron tomography and the diffusion compartment size determined by the high speed single-particle tracking of a phospholipid are similar to each other. However, between these two cell lines, both the MSK mesh and the diffusion compartment sizes differ greatly. The similarities between the MSK mesh sizes and the diffusion compartment sizes in cell lines that exhibit quite different distributions strongly support the MSK fence and picket models.

Discussion

We performed quantitative analyses of the undercoat structure of the cytoplasmic surface of the plasma membrane using electron tomography for samples prepared by a rapid-freeze, deep-etch, platinum replication technique. One of the most important limitations of this technique is that the cell has to be placed in a hypotonic medium at 4°C for 5–15 min to remove the upper cell membrane. However, with this method, large membrane fragments that were covered by the dense MSK meshwork could be obtained, which was important for the purpose of the present research.

We obtained the results by specifically addressing the following three questions. (1) Does the dense meshwork of the MSK exist everywhere on the cytoplasmic surface of the cell membrane, and, if so, how is it linked to the bulk cytoskeleton?

(2) If so, what is its relationship with other structures of the plasma membrane, such as CCPs and caveolae? (3) How is the distribution of the MSK mesh size right on the cytoplasmic surface of the plasma membrane?

The final point is important because this part of the MSK might form the corrals of the plasma membrane for the diffusion of membrane molecules. Therefore, it is interesting to compare the distribution of the mesh size of the MSK directly attached to the cytoplasmic surface of the plasma membrane, as determined by an EM method, with that of the compartment size for the diffusion of membrane molecules. NRK (median size = 230 nm) and FRSK (41 nm) cell lines were selected for such a comparison because their compartment sizes are very different (Fujiwara et al., 2002; Murase et al., 2004). This will be an interesting test for the MSK fence and MSK-anchored transmembrane protein picket models and became possible by obtaining the 3D reconstructed images of the MSK structure on the cytoplasmic surface of the plasma membrane.

Does the dense meshwork of the MSK exist everywhere on the cytoplasmic surface of the cell membrane, and, if so, how is it linked to the bulk cytoskeleton?

The cytoplasmic surface of the plasma membrane has been observed by EM for >30 yr (Byers and Porter, 1977). Stunning high resolution EM images of the structures near the cytoplasmic surface have been published previously (Heuser and Kirschner,

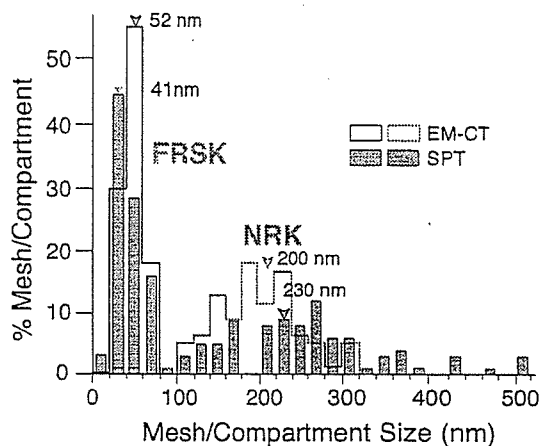


Figure 8. Comparison of the distributions of the MSK mesh size on the cytoplasmic surface of the plasma membrane estimated by electron tomography with that of the compartment size determined from the phospholipid diffusion data for NRK and FRSK cells. Electron tomography, open bars; phospholipid diffusion data, closed bars (adapted from Fujiwara et al., 2002 and Murase et al., 2004). NRK, pink; FRSK, blue. Within the same cell type, the MSK mesh size and diffusion compartment size exhibited similar distributions (compare the open and closed bars with the same color). The actual sizes are quite different between NRK and FRSK cells. EM-CT, EM-based computer tomograph; SPT, single-particle tracking.

1980; Hirokawa and Heuser, 1981; Heuser and Anderson, 1989; Hartwig and DeSisto, 1991; Rothberg et al., 1992), suggesting that the plasma membranes of animal cells are shaped by cytoskeletal interactions. However, despite these interesting and important studies, we felt that more extensive studies addressing the questions posed in the title of this subsection are necessary in particular for cells in culture.

We found that almost the entire cytoplasmic surface of the upper cell membrane is covered with the actin-based MSK except for the places where CCPs, caveolae, and noncoated membrane invaginations exist. In addition, this extensive pseudo-2D-type MSK network is linked to many actin filaments that extend from the membrane cytoplasmic surface toward the cytoplasm (Fig. 4 A) and is probably connected to the bulk cytoskeleton. This is consistent with the quantitative analysis of bleb formation, in which the density of the MSK filaments must be higher than one every 0.5–1 μm to avoid blebbing (Sheetz et al., 2006).

How closely is the MSK associated with CCPs and caveolae?

Almost all of the CCPs and caveolae are extensively linked to the MSK meshwork. As found in Figs. 2–4, many actin filaments that come out of the MSK meshwork are connected to these structures at their polygonal and striated patterns, respectively, and, in the cell types examined here, this occurs much more extensively than found previously (Rothberg et al., 1992; Fujimoto et al., 2000; Parton, 2003). Merrifield et al. (2002) found that actin rapidly becomes more concentrated at the CCPs right before they are internalized, but from the structures found in the present research, we could not tell what kind of actin-CCP interactions may be involved in such concentrations.

Is the distribution of the MSK mesh size right on the cytoplasmic surface of the plasma membrane consistent with that for the compartment size determined by a phospholipid diffusion probe?

3D reconstruction of the MSK using electron tomography allowed the determination of the MSK meshwork directly situated on the cytoplasmic surface of the plasma membrane. This meshwork is likely to partition the plasma membrane into many small compartments with regard to the lateral diffusion of membrane molecules. The size distribution of the areas bounded by the MSK meshwork agreed well with that determined for an unsaturated phospholipid undergoing hop diffusion in high speed single-particle tracking experiments in both NRK and FRSK cells, which have quite different mesh sizes (Fig. 8). These results support the MSK fence (corralling) model and the anchored transmembrane protein picket model.

How the MSK is attached to the cytoplasmic surface of the plasma membrane is unknown. Specific proteins that link the membrane and actin filaments at their barbed ends, such as gelsolin and villin (Pollard and Cooper, 1986; Hartwig et al., 1989), and at their sides, such as ponticulin (Wuesthube and Luna, 1987) and ezrin/radixin/moesin family proteins (Tsukita et al., 1997), exist. However, the weak nonspecific binding of actin filaments to membrane lipids and proteins may greatly contribute to the interactions of the actin filaments with the membrane. Although transmembrane proteins are temporarily trapped within a compartment, they hop to adjacent compartments in longer terms, suggesting that the link between the membrane and actin filaments may break at a rate comparable with the hop rate, which is once every several to several hundred milliseconds (depending on the molecule and cell type).

In addition to actin and actin-associated proteins, some other proteins may contribute to forming the MSK and membrane corrals. For example, septin, which is $\sim 7\text{-nm}$ thick in the negatively stained images (Kinoshita et al., 2002), and agorin (Apgar and Mescher, 1986) may play some roles. Further characterization of the proteins involved in the MSK and its interaction with the membrane components and further analysis of the dynamics of membrane-MSK interactions will be important for understanding the roles played by the MSK-plasma membrane interactions in signal transduction, domain formation in the plasma membrane, and reorganization of the cytoskeleton.

Materials and methods

Antibodies and other reagents

Rabbit anti-actin IgG was obtained from Biomedical Technologies, and colloidal gold probes (5-nm diameter) coated with secondary antibodies (produced in goat) were purchased from GE Healthcare.

Rapid-freeze, deep-etch, immunoreplica EM of the cytoplasmic cell surface
NRK and FRSK cells were maintained in HAM-F12 or DME mediums, respectively, supplemented with 10% FBS (Sigma-Aldrich) under a 5% CO_2 atmosphere at 37°C. The cells used for the experiments were grown in 35-mm plastic dishes to $\sim 70\%$ confluency, usually 2 d after inoculation. The cells were washed three times with ice-cold Pipes buffer (10 mM Pipes, 100 mM KCl, 5 mM MgCl_2 , and 3 mM EGTA, pH 6.8, which mimics the environment in the cytoplasm somewhat but is slightly hypotonic) and were exposed for 15–30 s to an ice-cold 70% Pipes buffer (the Pipes buffer diluted by a factor of 1.43 with water, making this solution more hypotonic;

Rutter et al., 1988). After the buffer on the coverslip was drained, the remaining excess water was removed by filter paper. To expose the cytoplasmic surface of the upper cell membrane, the upper cell membrane was removed from the rest of the cell after it was adhered to a coverslip placed on top of the cell layer (Rutter et al., 1988; Sanan and Anderson, 1991). 5 × 5-mm coverslips (Matsunami) coated with positively charged Alcian blue 8GX (Wako; Alcian blue-coated coverslips were prepared by first immersing them in 1% Alcian blue in distilled water at room temperature for 10 min, washing them with distilled water, and drying them in the air) were placed on top of the cells (upper surface facing the medium rather than the coverslip) and incubated at 4°C for 5–15 min. During this period, good contact between the cell surface and the coverslip was developed. Then, the coverslips were gently floated off from the cell using the surface tension of the buffer by slowly adding ice-cold Pipes buffer containing 1% PFA/0.25% glutaraldehyde into the space between the culture dish and the coverslip. When the coverslip floated off, the cells were cleaved, and the upper cell membrane came off with the coverslip. Then, the cells were further fixed by incubation in fresh ice-cold 1% PFA/0.25% glutaraldehyde in Pipes buffer for 10 min. After fixation, the coverslips were washed three times, for 10 min each time, with PBS [8.10 mM Na₂HPO₄, 1.47 mM KH₂PO₄, 137 mM NaCl, and 2.68 mM KCl, pH 7.4].

To identify the actin filaments on the cytoplasmic surface of the cell membrane, immunogold labeling was performed after fixation. The fixed upper cell membranes on the coverslips with their cytoplasmic surfaces exposed toward the buffer were incubated at 4°C for 2 h in PBS containing 10 µg/ml rabbit anti-actin IgG and 1% BSA (Sigma-Aldrich), and, after three washes for 10 min each in 25 mM Tris-buffer, pH 8.0, the cell membranes were incubated in Tris-buffer containing 1% BSA and 5-nm-diameter colloidal gold conjugated with anti-rabbit IgG goat antibodies (GE Healthcare) at 4°C for 2 h. After three washes in PBS, these labeled membranes were further fixed in 2% glutaraldehyde in PBS on ice for 5 min. Finally, the coverslips were washed in distilled water for 1 min before rapid freezing.

Each coverslip was placed on the plunger tip of the rapid-freezing device (Eiko; Usukura and Yamada, 1987) with the cytoplasmic surface of the membrane facing down. The specimen was slammed down (free fall) onto a polished pure copper block, which was precooled by direct immersion in liquid helium. The frozen coverslip was placed in liquid nitrogen and was transferred into the freeze-etching shadowing chamber (FR7000-S; Hitachi). The excess ice covering the cytoplasmic surface of the membrane was shaved off with a prechilled glass knife using a microtome placed in the chamber at -140°C or below. The cytoplasmic surface was then etched for ~10 min after the specimen temperature was raised to -90°C. The etched specimen surfaces were then rotary shadowed with platinum at an angle of 22.5° from the surface and with carbon from the top. The molecules as well as the gold probes localized on the cytoplasmic surface of the cell membrane were immobilized to the deposited platinum (Fujimoto, 1995; Fujimoto et al., 1996).

Collodion was applied immediately after the platinum-carbon replicas were removed from the cold chamber to fortify them. The platinum-carbon replica was removed from the glass coverslip in 1% hydrofluoric acid in distilled water. After the replicas were successfully removed from the glass surface and mounted on the grid, the collodion coat was dissolved away in n-pentyl acetate. In this protocol, the sodium hypochlorite solution, which is generally used to remove the replicas from the coverslip and also to clear the membrane and the undercoat structure of the replicas, was replaced with 1% hydrofluoric acid to keep the cell membrane, the undercoat structure, and the immunogold probes that had been attached to these structures on the platinum replicas [1% hydrofluoric acid is likely to only dissolve the glass, leaving the cell membrane molecules bound to the platinum replica; Fujimoto, 1995; Fujimoto et al., 1996]. An additional advantage of using 1% hydrofluoric acid is that the platinum replicas break less often, probably because it does not remove the membrane components and, thus, leaves the replicas rather intact. In addition, to keep as many colloidal gold particles and membrane molecules attached to the platinum replicas as possible, we included Photo-Flo 200 (Kodak), a detergent used to prevent water-drop stains on photographic film in all of the solutions used here (advice given by J. Heuser). After the replicas were washed with distilled water, they were mounted on 100–200 mesh copper grids (Ted Pella) coated with polyvinyl formvar (Nissin EM) and observed at magnifications of ~10,000–70,000 with a transmission electron microscope (I200EX; JEOL).

The following methodological precautions and improvements were made to reproducibly produce large cell membranes and replicas without excessive fragmentation. (1) An Alcian blue coat rather than poly-L-lysine coat was used (Rutter et al., 1988; Sanan and Anderson, 1991). (2) Before

overlaying the coverslips, excess water was removed from the specimen, leaving just enough buffer to cover the cell. (3) To cleave off the upper membrane attached to the overlaid coverslip, the coverslip was floated off very gently by adding cleavage medium (using the surface tension of the buffer to float the coverslip). If this was not performed gently enough, the membrane was fragmented. (4) The frozen sample was shaved with a glass knife, with the angle between the knife and the coverglass adjusted to a shallow angle (<6°) so that most of the excess water and the cytoplasm were removed and the cytoplasmic surface of the cell membrane could be exposed after light etching. Because replicas with too many variations in height tend to break when they are removed from the coverslip and placed on the water surface, removal of the excess cytoplasm helps to avoid replica breakage. (5) Collodion was applied immediately after the replicas were removed from the cold chamber (before the replicas were removed from the coverslip on the water surface) to fortify the replica (a technique learned from T. Baba and S. Ohno). This step also helped to prevent replica breakage when the replicas were removed from the coverslip. After the large replicas were removed from the glass surface, the collodion coat was dissolved away in n-pentyl acetate. (6) A solution of 1% hydrofluoric acid was used to slightly dissolve the glass surface to facilitate the removal of replicas from the coverslip. (7) A detergent, Photo-Flo 200 (Kodak), was included in all of the solutions that contacted platinum replicas.

Electron microscope tomography

For 3D reconstruction, the replica was imaged at tilt angles of every 1.0° in the range between ±48 and 70° (total of 97–141 images) for a single field by a transmission electron microscope (Tecnai Sphera F20; FEI) equipped with a CCD camera (1,024 × 1,024 pixels). The pixel size at the specimen was 0.85 nm. The image acquisition was fully automated as previously described (Medalia et al., 2002). The 100–121 image sections of every 0.85–1.34 nm were obtained by a calculation based on the set of 97–141 tilt images using an IMOD software package running on Linux (University of Colorado; Kremer et al., 1996). Corrections for the tilt of the specimen and the long wavelength undulations of the membrane were also achieved with IMOD software. 3D rendering (displaying 3D images in different ways) was performed using the Amira DEV software package (Mercury Computer Systems) operating on a Linux system.

Analysis of the 3D reconstructed images of the MSK

In the series of electron tomography sections shown in Fig. 6 (A and B), the existence of three major classes of filaments with regard to the distance from the membrane surface was found in the following way. The first class is the filaments that are highly electron dense in the first ~0–1.7-nm section (because the contrast is reversed in these micrographs, they look more lucent or white) and are continuously seen in the image sections up to the ~6.8–8.5-nm section, which then dim rapidly in the ~8.5–10.2- and ~10.2–11.9-nm sections. To quantitatively evaluate such signal intensity changes within individual filaments, we selected points that are clearly seen in the image of the first ~0–1.7-nm section every 100–250 nm on each filament, measured the signal intensity on each spot (five pixels), and looked for the section where the signal intensity on the spot decreases by >25% from that for the adjacent section closer to the membrane (the signal intensity tends to drop very rapidly around the threshold sections). If the 25% decrease in the signal intensity occurred between the sections of ~6.8–8.5 and ~8.5–10.2 nm or between the sections of ~8.5–10.2 and ~10.2–11.9 nm, these filaments were categorized into the first class (i.e., those closely associated with the cytoplasmic surface of the cell membrane). These filaments are drawn in green in Fig. 6 C (different regions within a single filament might become dim in either of these two sections).

The second class is similar to the first class but can be seen clearly even in the ~10.2–11.9-nm section (and also in the ~11.9–13.6-nm section, which usually looked similar to the ~10.2–11.9-nm section). This may be the result of the actin filaments that are located several nanometers away from the cytoplasmic surface. Because the platinum rotary shadowing was performed at a low angle (22° from the surface), the platinum may have been deposited in the space between the filament and the membrane. Another possibility is that two filaments are stacked together for a long distance, but we do not think that this happens very often. Therefore, we categorized these filaments into those that stay near the membrane but do not closely associate with the membrane surface. These filaments were not considered to contribute to delimiting the membrane compartments for molecular diffusion in the plasma membrane.

The third class is the filaments that exhibit dim signals in the first ~0–1.7-nm section and show higher electron densities in farther sections, at least up to the section of ~10.2–11.9 nm, before fading out in the ~11.9–13.6- and ~13.6–15.3-nm sections (the latter two sections are not depicted).