Epithelial cells form monolayers of polarized cells with apical and basolateral surfaces. Madin-Darby canine kidney epithelial cells transiently lose their apico-basolateral polarity and become motile by treatment with hepatocyte growth factor (HGF), which causes the monolayer to remodel into tubules. HGF induces cells to produce basolateral extensions. Cells then migrate out of the monolayer to produce chains of cells, which go on to form tubules. Herein, we have analyzed the molecular mechanisms underlying the production of extensions and chains. We find that cells switch from an apico-basolateral polarization in the extension stage to a migratory cell polarization when in chains. Extension formation requires phosphatidyl-inositol 3-kinase activity, whereas Rho kinase controls their number and length. Microtubule dynamics and cell division are required for the formation of chains, but not for extension formation. Cells in the monolayer divide with their spindle axis parallel to the monolayer. HGF causes the spindle axis to undergo a variable “seesaw” motion, so that a daughter cells can apparently leave the monolayer to initiate a chain. Our results demonstrate the power of direct observation in investigating how individual cell behaviors, such as polarization, movement, and division are coordinated in the very complex process of producing multicellular structures.

INTRODUCTION

One of the most common types of cell polarity is found in polarized epithelial cells, which have an apical surface facing the lumen and a basolateral surface facing other cells and extracellular matrix (Drubin and Nelson, 1996; Mostov et al., 2000). Another category of cell polarity is found in migrating cells, which includes cell movements during development, chemotaxis, wound healing, and metastasis (Nabi, 1999; Franz et al., 2002; Webb et al., 2002). In these types of movement, the cells become polarized in the direction of movement. Migratory polarization seems very different from the seemingly more static polarization of epithelial cells. Nevertheless, epithelial cells can be induced to become motile and assume a morphology and polarization that resembles that of a migratory cell, such as a fibroblast. This occurs during epithelial-mesenchymal transitions that happen in development or oncogenesis (Arias, 2001; Savagner, 2001). Herein, we focus on how cells transition from epithelial polarity to migratory polarity.

One model to address this is to treat polarized epithelial cells, such as Madin-Darby canine kidney (MDCK) cells, with hepatocyte growth factor (HGF) (Birchmeier and Gherardi, 1998). HGF is also known as scatter factor, because it induces MDCK cells grown subconfluently on a solid support to “scatter,” i.e., become highly motile (Comoglio and Boccaccio, 2001). The transition from epithelial polarity to migratory polarity can also be studied in cells grown in three-dimensional gels of extracellular matrix, such as type I collagen. These culture conditions may more closely resemble the in vivo environment and have provided useful model systems for studying the formation of multicellular epithelial structures (Cukierman et al., 2001; O’Brien et al., 2002; Zegers et al., 2003).
When a single cell suspension of MDCK cells is embedded in such a gel, the cells divide and form spherical cysts, where a monolayer of cells surrounds a central lumen. Cells are polarized, with the apical surface facing the lumen (O'Brien et al., 2001). Treatment of these cysts with HGF causes them to elaborate branching tubules. This process involves a complex series of changes in cell polarity and movement (Montesano et al., 1991a,b; Metzger and Krasnow, 1999; Hogan and Kolodziej, 2002; O'Brien et al., 2002). HGF-induced tubulogenesis can be divided into four stages, termed extension, chains, cords, and tubules (Pollack et al., 1997, 1998). Approximately 6–12 h after addition of HGF some cells produce extensions, large pseudopods protruding from their basolateral surface. These cells retain a small apical surface and are thus part of the monolayer. At ~24 h, chains of one to three cells protrude from the cyst. At this stage, cells have lost apico-basal polarity. Subsequently, chains transform into cords two to three cells thick. Small lumens lined by newly established apical surfaces form between cells. These lumens expand to ultimately form a single lumen contiguous with the lumen of the original cyst, marking the completion of mature tubules.

Herein, we have studied the early stages in which the cells in the monolayer of the cyst wall form extensions and chains, because these may correspond to the transition from epithelial polarity to the polarity of a migrating cell. These stages also involve cell movement, because formation of extensions involves movement of a portion of the cell, and formation of chains involves movement of entire cells away from the cyst. To analyze the mechanisms underling extension and chain formation, we examined four classes of molecules and processes that are known to be involved in polarization and movement in other systems. 1) Phosphatidylinositol 3 kinase (PI3K). We found that the products of PI3K are concentrated at the leading edge in extensions and that PI3K is needed for extension formation. 2) Microtubules. Surprisingly, microtubule dynamics is not needed for extension formation. 3) The Rho effector, Rho kinase (ROCK). We found that ROCK determines the number and length of extensions. 4) Cell division. Cell division is not needed for extension formation but is required for making chains. As cells divide and migrate out to form chains, they change the orientation of their division from parallel to the plane of the monolayer to roughly perpendicular to this plane. This is an example of an oriented cell division in an experimentally tractable mammalian system, which should facilitate future analysis of this important process.

MATERIALS AND METHODS

Cell Culture, Cystogenesis, and Tubulogenesis

MDCK cells were maintained in minimal essential medium containing Earle’s balanced salt solution (Cellgro; Mediatech, Herndon, VA) supplemented with 5% fetal bovine serum (HyClone Laboratories, Logan, UT), 100 unit/ml penicillin, 100 μg/ml streptomycin (control medium below) in 5%CO2, 95% air. For growth of cells in three-dimensional collagen gels, cells were trypsinized and then triturated at room temperature into a single-cell suspension of 4 × 10^6 cells/ml in type I collagen solution containing 66% vitrogen (control medium below) in 5%CO2, 95% air to allow the collagen to gel and then control medium was added. Cells were fed every 3 d and grown for 7–10 d until cysts with lumen were formed. For tubulogenesis, 7-d-old cysts were stimulated with 1:1 conditioned medium (CM) containing HGF from MRC5 human lung fibroblasts: control medium.

Cell Transfection

Early passage MDCK cells grown to ~30% confluence on 10-cm plastic dishes were transfected with 20 μg of a cDNA expression vector encoding green fluorescent protein (GFP)-α-tubulin (BD Biosciences Clontech, Palo Alto, CA), GFP-pleckstrin homology (PH)-Akt (Servant et al., 2000), or GFP-myosin light chain (MLC), according to previously published protocols (Breitfeld et al., 1989; Lipschutz et al., 2001). Transfected cells were replated in medium containing 700 μg/ml G418 72 h after transfection. Medium was changed every 3–4 d. After 12 d of G418 selection, surviving cloned were picked and screened for GFP expression. Clones expressing GFP-tagged proteins were detected by observation with a confocal laser scanning microscope and Western blot with anti-GFP antibody diluted at 1:1000. One clone of each that formed cysts in collagen gel was chosen for use. All cell lines were kept under selection with G418.

Antibodies and Reagents

The primary antibodies used in this study were mouse anti-cis-Golgi enzyme GM130 (Transduction Laboratories, Lexington, KY), mouse anti-α-tubulin antibody (Sigma-Aldrich, St. Louis, MO), and goat anti-P-MLC (Thr18,Ser19) and rabbit-anti-MLC (Santa Cruz Biotechnology, Santa Cruz, CA). Secondary antibodies were anti-mouse Alexa 594 (Molecular Probes, Eugene, OR). Actin filaments were stained with Alexa Fluor 488 phalloidin (Molecular Probes). Nuclei were labeled with TOPRO-3 (Molecular Probes). Reagents used in this study were nocodazole and LY29004 (Sigma-Aldrich), mitomycin C and Y27632 (Calbiochem, San Diego, CA), and G418 (Invitrogen, Carlsbad, CA).

Western Blot Analysis of MLC Levels

MDCK cells were grown to confluence in 24-mm-diameter wells and then treated with CM, CM with 30 μM Y27632, and Y27632 for 16 h, respectively. Cells were lysed in 500 μl of Triton X-100 lysis buffer (10 mM triethanolamine, pH 7.4, 1.1% Triton X-100, 0.1% SDS, 100 mM NaCl, 1 mM EDTA, 1 mM EGTA, 1 mM NaF, 20 mM Na2PO4, and 2 mM NaVO3). Lysates were boiled for 5 min and cleared by centrifugation. Protein concentrations were determined by a bicinchoninic acid protein assay (Pierce Chemical, Rockford, IL), and equal amounts of proteins were electrophoresed on 4–15% SDS-PAGE gels (Bio-Rad, Hercules, CA), immunoblotted, and detected with an enhanced chemiluminescence system (PerkinElmer Life Sciences, Boston, MA).

Treatment of MDCK Cysts with Reagents

LY294002. Seven-day-old MDCK cysts were incubated with 20 μM LY294002 in CM at 37°C for 16 h. As a control, cysts were treated with CM only for 16 h. For chain formation, cysts were treated with CM for 16 h to produce extensions and then treated with 20 μM LY294002 in CM.

Nocodazole. Seven-day-old cysts were preincubated at 4°C for 1 h and then incubated in 200 ng/ml nocodazole in CM at 37°C for 16 or 40 h separately. As a control, cysts were treated with only CM for 16 or 40 h, without the 4°C incubation. To arrest cells at a mitotic stage during tubule formation, cysts were stimulated with CM for 72 h and then incubated in 200 ng/ml nocodazole for an additional 24 h.

www.current-biology.com
W. Yu et al.

Molecular Biology of the Cell

750
Mitomycin C. Seven-day-old cysts were preincubated with 0.5 ng/ml mitomycin C at 37°C overnight and then treated with mitomycin C in CM at 37°C for 16 or 40 h, respectively. As control, cysts were treated with CM only for 16 or 40 h.

**Immunofluorescence Staining**

The procedure for the immunofluorescence staining was described in detail in previous publications (Pollack et al., 1998; O’Brien et al., 2001). All samples were rinsed with phosphate-buffered saline + twice quickly and then treated with collagenase type VII (Sigma-Aldrich) at 37°C for 10 min. Cysts were fixed with 3% paraformaldehyde for 30 min and with 0.25% Triton X-100 for 10 min at room temperature, and then quenched with 50 mM NH4Cl for 10 min and permeabilized with 0.1% saponin and 0.7% gelatin for 30 min. Samples then were incubated in primary antibodies at 4°C for overnight. After sufficient washing, samples were treated with 0.1 mg/ml RNase at 37°C for 10 min and then incubated in Alexa 594-conjugated secondary antibody 1:200 dilution and Alexa Fluor 488-phalloidin at 1:50 dilution at 4°C overnight. After washing, cysts were stained for nuclei with TOPRO3 at 1:500 dilution at 37°C for 1 h. All samples were mounted with Prolong medium (Molecular Probes) and then dried at room temperature overnight.

**Time-Lapse Images**

Cells expressing GFP-e-tubulin were grown in three-dimensional collagen gels as described above. Coverglass-bottomed chambers (Nalge Nunc, Naperville, IL) coated with Sigmacote (Sigma-Aldrich) were used for growing cysts to obtain live cell images. After 7 d of culture, cysts were incubated in phenol red-free medium for 48 h and then fed with phenol red-free Leibovitz’s L-15 medium to permit incubation in room air (Invitrogen) with 1% antibiotic and 5% fetal bovine serum for 2 d. Therefore, we used recombinant human HGF (rhHGF; generous gift of R. Schwall, Genentech, South San Francisco, CA) in these experiments, because the buffer component, L-15 medium, CM). CM contains high concentrations of HGF, which had been conditioned by MRC5 fibroblasts (conditioned medium, CM). CM contains high concentrations of HGF, which is the tubulogenesis-promoting activity in CM (Figure 1A and B) (Montesano et al., 1991a,b). Indeed, when we added 2.5 μg/ml neutralizing anti-HGF antibodies to the CM, tubulogenesis was completely abolished (Figure 1C). We found that in control cysts GFP-PH-Akt was uniformly localized at the basolateral membrane but was excluded from the apical surface (Figure 1D). A similar localization of this probe was previously observed in MDCK cells grown as a monolayer on filters (Watton and Downward, 1999), (Montesano et al., 1991a,b). In cells that had been treated with CM for 16 h and formed extensions, GFP-PH-Akt strikingly accumulated in extensions, thus suggesting a possible involvement of PI3K in extension formation (Figure 1E).

Next, we treated cysts with a specific inhibitor of PI3K, LY294002. At 20 μM, LY294002 strongly inhibited the formation of extensions and resulted in cysts that formed only a very few short and thin extensions (compare Figure 1F and G). Similar results were obtained after treatment with 25 μM wortmannin (our unpublished data). Moreover, extension formation in LY294002 and wortmannin treated cells (our unpublished data) was fully restored upon washout of the drugs (Figure 1H), which argues against the possibility that the effects of the drugs were due to toxicity or induction of apoptosis. These findings suggest a requirement for PI3K during extension formation.

We also examined the localization of GFP-PH-Akt in cells that had been treated with CM for 40 h to allow the formation of chains. In contrast to unstimulated cells in cysts or CM-stimulated cells containing extensions, cells in chains had GFP-PH-Akt uniformly around the entire plasma membrane (Figure 1I). To test the role of PI3K in chain formation,
CM

A

B

CM + nocodazole

C

C'

D

E

W. Yu et al.
we first treated cysts with CM for 16 h to produce extensions and then incubated the cysts for an additional 24 h in the continued presence of CM and in the concurrent absence or presence of LY294002. As expected, in the absence of LY294002 the extensions went on to form chains (Figure 1J). However, in the presence of LY294002, no chains and indeed very few extensions were seen (Figure 1K). Presumably, the LY294002 caused the extensions that had formed to collapse.

**Golgi Orientation**

During migration of several cell types, such as fibroblasts, T cells, and astrocytes, the Golgi apparatus reorients toward the leading edge (Schliwa and Höner, 1993; Stowers et al., 1995; Nobes and Hall, 1999; Etienne-Manneville and Hall, 2001). This may facilitate the delivery of biosynthetic and perhaps recycling vesicles to the growing region of plasma membrane. It was therefore interesting to ask whether the Golgi similarly reoriented toward the leading edge of extensions and/or chains. As described previously, in control cells, and astrocytes, the Golgi apparatus reorients toward the leading edge (Schliwa and Höner, 1993; Stowers et al., 1995; Nobes and Hall, 1999; Etienne-Manneville and Hall, 2001). This may facilitate the delivery of biosynthetic and perhaps recycling vesicles to the growing region of plasma membrane. It was therefore interesting to ask whether the Golgi similarly reoriented toward the leading edge of extensions and/or chains. As described previously, in control
cysts the Golgi is located beneath the apical surface, i.e., between the nucleus and the apical surface (Figure 2A) (O’Brien et al., 2001). Surprisingly, for treatment with CM for 16 h in cells that remained in the cyst wall and had extensions, the Golgi did not reorient toward the leading edge of the extension. Rather, in 100% (n = 30) of such cells, the Golgi remained under the apical surface (Figure 2B). Some of these cells had only very small apical surfaces remaining and the nuclei were starting to move away from the cyst. In contrast, after 40 h of CM, for cells in chains (i.e., no detectable apical surface), the Golgi frequently reoriented. For cells at the tip of the chain, in 69% the Golgi was now oriented at the leading edge (example shown in Figure 2B), whereas in only 18% the Golgi was still facing toward cyst wall, and in 13% the Golgi was at the sides of the cell (n = 61). In cells located at the middle of the chains (i.e., neither in cyst wall nor at the tip of the chain), the Golgi was apparently randomly located (31% toward the tip, 36% at the sides, 33% located toward the cyst wall, n = 45).

**Role of Microtubule Dynamics**

The Golgi is often located near the centrosome or microtubule organizing center, and its localization can be used as a marker for the location of the microtubule organizing center. The centrosome reorients during migration of some types of cells (e.g., Chinese hamster ovary) but not others (e.g., PK) (Yeon et al., 2002). Given the changes in localization of the Golgi, we asked whether dynamic changes in microtubules (MTs) play a role in formation of extensions and chains (Wittmann and Waterman-Storer, 2001). Although the MTs in polarized epithelial cells are relatively stable and resistant to depolymerization by nocodazole, low concentrations of this drug (e.g., 200 ng/ml) can block MT dynamics (Vasquez et al., 1997). Cysts were preincubated at 4°C for 1 h and then incubated at 37°C with CM for 16 or 40 h, in the presence or absence of 200 ng/ml nocodazole. The number of cells with mitotic figures, most with normal-looking spindles, increased 13- and 41-fold after a 16- and 40-h nocodazole treatment, respectively, thus confirming that the nocodazole was inhibiting MT dynamics. Higher concentrations of nocodazole produced abnormal spindles. Inhibition of MT dynamics was furthermore apparent from a fragmented Golgi (Figure 2, C and C’). At 16 h, cysts treated with nocodazole had formed extensions, which were similar in number, size, and shape to those in the minus nocodazole control (Figure 2, C and D, compare with control in Figure 1, B and E). However, at 40 h no chain formation was observed. Some isolated, rounded cells were seen outside of the cyst, but these do not seem to be in contact with each other or with the cyst (Figure 2E). These cells seem to arrested in mitosis, due to the inhibition of MT dynamics by nocodazole. This is in contrast to the well-formed chains seen in controls treated with CM alone for 40 h (Figure 1, I and J).

**Role of the Actin Cytoskeleton and Rho Kinase**

We observed that the formation of both extensions and chains was blocked by the actin inhibitors latrunculin B and jasplakinolide (our unpublished data). The function of the actin cytoskeleton in cell polarity and movement is regulated by small GTPases of the Rho family, such as Rho, Rac, and Cdc42 (Bishop and Hall, 2000; Ridley, 2001).

We have previously reported that a dominant negative form of Rac1 causes an inversion of the polarity of cysts (O’Brien et al., 2001) and prevents tubulation in response to HGF (O’Brien and Mostov, unpublished data). We have found similar results with a dominant negative version of Cdc42 (Datta and Mostov, unpublished data). To investigate the role of RhoA in extension and chain formation, we used the drug Y27632, which blocks a major class of RhoA effectors, ROCK I and ROCK II (Uehata et al., 1997). ROCK is necessary for maintaining a cortical actin-myosin network and for formation and contractility of stress fibers (Nobes and Hall, 1999). We first treated cells with 30 μM Y27632 alone for 24 h. Although the treated cysts looked nearly normal by phase contrast microscopy (our unpublished data), confocal microscopy by using fluorescent phalloidin showed the presence of very short basolateral extensions (compare Figure 3A, control, with E, Y27632 treated). These extensions were visible as early as 3 h after treatment with Y27632 and were comparable to the extensions seen with a 3-h treatment of CM alone (our unpublished data). How-

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**Figure 2 (facing page).** Orientation of the Golgi in extensions and chains and effect of nocodazole on extension and chain formation. (A) Cyst treated with CM for 16 h to produce extension. Golgi marker (GM130) is red, fluorescent phalloidin is green, and nuclei are blue. One cell has formed an extension and has a concentration of actin at the tip of this extension, near the right edge of the panel. This cell still has a small apical surface, which is at the base of a deep indentation of the cyst lumen. Note that the Golgi in this cell is still underneath the apical surface (arrow). (B) Cyst treated with CM for 40 h to form a short chain. The cell in the chain has left the monolayer. Note that the nucleus is close to the nucleus of a cell that remains in the monolayer. Arrow indicates the Golgi in this chain, which is oriented toward the leading edge of the cell, away from the cyst. (C and C’) Disassembly of the Golgi in nocodazole-treated extension. Cyst was preincubated for 1 h at 4°C and then treated for 16 h with CM and 200 ng/ml nocodazole. (C) Staining as in A. C’ emphasizes fragmented Golgi by GM130 staining alone. (D) Cyst was preincubated for 1 h at 4°C and then treated for 16 h with CM and 200 ng/ml nocodazole. Green is fluorescent phalloidin, blue is nuclei, and red is microtubules. Fairly normal extensions are seen. (E) As in D, except treatment was for 40 h. Note many cells arrested in mitosis (red spindles) in and around the cyst. Bar, 10 μm.
ever, the CM-only extensions continued to grow, whereas the Y27632-only extensions apparently remained the same length at times >3 h (our unpublished data).

When cells were treated with CM and 30 μM Y27632 for 24 h, we noticed a dramatic increase in the number and the length of extensions (Fig. 3, compare B with F). The number of extensions at 24 h increased ~9.5-fold (n = 59 cysts for CM alone, n = 35 for CM + Y27632). The mean extension length with CM alone was 32.2 ± 5.0 (SD) μm, and 67.8 ± 8.4 μm with Y27632 + CM (n = 55 extensions in each group). Taken together, these data suggest that ROCK normally restricts the number and length of extensions. We also examined the effect of Y27632 on chain formation. In cysts treated for 40 or 72 h with CM and Y27632, cells migrated out of the cysts (Figure 3, compare C with G and D with H). However, the cells lost contact with their neighbors and formed isolated cells (Fig. 3, G and H). Chains were not seen.

Strikingly, in addition to migrating away from the cyst, some cells migrated into the center of the cyst after 24 h (Figure 3F), and by 40–72 h the lumen of the cyst was obliterated by cells that had migrated into it (Figure 3, G and H). This suggests that Y27632 disrupts the directional mechanism that normally causes the cells to migrate only away from the cyst and instead allows them to migrate in multiple directions.

ROCK has several known substrates. One of its major substrates is myosin phosphatase, which results in elevated phosphorylated myosin light chain (P-MLC) (Kimura et al., 1996). P-MLC is in turn correlated with assembly of myosin II filaments and activation of myosin ATPase activity (Chrzansowska-Wodnicka and Burridge, 1996). MLC may also be a substrate of ROCK (Amano et al., 1996). We used a goat antibody that is specific for P-MLC (phosphorylated on Thr18 and Ser19) to localize the P-MLC. In cells in the monolayer of the cyst wall, of untreated cyst most of the P-MLC signal is at the basal surface (Figure 4A), coincident with the actin stress fibers that underlie the basal surface (our unpublished data). Some of the P-MLC is concentrated apically, in the area of tight junctions (arrowheads).

We were unable to localize total endogenous MLC, because the available antibodies did not work for immunocytochemistry on cysts embedded in thick collagen gels. We therefore expressed a GFP-MLC fusion, which has been shown to localize similarly to endogenous MLC (Ward et al., 2002). This GFP-MLC localized mainly to the basal surface, although a small amount localized to the apical region, similar to P-MLC (Figure 4, A’ and A”, arrowheads). We were able to measure the total amount of MLC by using a Western blot, and this did not change with CM and/or Y27362 (Figure 4H).
In extensions, most of the P-MLC and GFP-MLC are at the elongated basal surface (Figure 4, B and C, solid arrows). Arrowheads indicate P-MLC in region of TJ. GFP-MLC (A’, green in merge) was mainly localized at basal plasma membrane, nuclei in blue. (B, D, and F) P-MLC in extension (B, 16 h CM), chain (D, 40 h CM), and extension after Y27632 treatment (F, 16 h CM + 30 μM Y27632), respectively. (B) Solid arrows indicate P-MLC associated with basolateral plasma membrane in extension. (D) Open arrowheads indicate P-MLC in cytoplasm of cell in chain. Left open arrow is in leading region of the cell, whereas right open arrow is in trailing region of the cell. (F) Asterisks indicate elongated, narrow extension. (C, E, and G) GFP-MLC in extension (C, 16 h CM), chain (D, 40 h CM), and extension after Y27632 treatment (G, 16 h CM + 30 μM Y27632), respectively. GFP-MLC is primarily located at the basolateral membrane, but some staining is observed apically, near tight junctions (arrowheads). (H) CM and Y27632 do not affect MLC expression levels. Confluent MDCK cells on 24-mm wells remained untreated or were treated for 16 h with CM in absence or presence of 30 μM Y27632, or with Y27632 alone. MLC in whole cell lysates was detected by Western blotting by using an anti-MLC antibody.

This relocalization of P-MLC is yet another difference between extensions and chains. When cysts were treated with both CM and Y27632, the P-MLC signal was weaker (Figure 4F), consistent with the idea that Y27632 causes an inhibition of phosphorylation of P-MLC. Some of the P-MLC was in the elongated extensions (Figure 4F, asterisks), which seemed narrower than those formed without Y27632 (e.g., solid arrows in Figure 4B). Due to the apparent narrowness...
of the extensions formed in the presence of CM + Y27632, it was difficult to be certain whether the P-MLC was cytoplasmic and/or associated with the inner surface of the plasma membrane of the extension. Some of the P-MLC remained associated with the TJ (Figure 4F, arrowhead). This TJ-associated P-MLC was not appreciably diminished in intensity compared with the TJ-associated P-MLC in control (Figure 4B), suggesting that the TJ-associated P-MLC may be relatively resistant to loss of phosphorylation caused by Y27632.

Taken together, these results suggest that ROCK normally causes contractility of actin-myosin bundles, which acts to inhibit production and elongation of extensions. Blockage of ROCK by Y27632 relieves this inhibition of extensions, resulting in more and longer extensions. Phosphorylation of MLC is likely to be a relevant result of ROCK activity, although other downstream effectors of ROCK may be involved.

Role of Cell Division

The development of some invertebrate tubules, such as the trachea in *Drosophila*, occurs postmitotically (Metzger and Krasnow, 1999; Hogan and Kolodziej, 2002). We investigated whether cell division was necessary for tubulogenesis in the HGF/MDCK system. Apart from enhancing cell motility and cell scattering, HGF is a potent mitogen in a variety of cell types, including filter-grown MDCK cells (Birchmeier and Gherardi, 1998). Indeed, we observed a sixfold increase in mitotic figures in cysts that were treated with CM for 16 h (47 per 100 cysts in CM-treated cells vs. 7 in untreated controls). As shown in Figure 2, nocodazole treatment leads to a block in both cell division and chain formation, which indicates a role for cell division in tubulogenesis. To further analyze the role of cell proliferation in HGF-induced tubulogenesis, we inhibited cell proliferation by preincubating cysts with the cell cycle inhibitor mitomycin C at 37°C overnight. Cysts were then treated with CM in the presence of mitomycin C for an additional 16 h. As was observed with nocodazole, cysts treated with mitomycin C did not form chains (Figure 5, compare A and B). Moreover, the cysts lost most of their extensions. Importantly, the general polarized structure of the cysts was largely unchanged. This suggests that the treatment with mitomycin C was not completely toxic to the
cells, because toxicity would have resulted in a collapse of cyst structure and cell death, which was also not observed. Similar results were obtained with another inhibitor of the cell cycle, t-mimosine (our unpublished data). These results show that cell division is not required for formation of extensions but is required for the formation of chains.

Location of Cell Division
Where do cells divide during tubulogenesis? In some mammalian systems, such as lung and endothelia, cell division takes place primarily at the tips of the growing tubules and branches (Mollard and Dziadek, 1998; Nogawa et al., 1998; Huang and Ingber, 1999). We analyzed the location of mitotic figures in CM-treated cells. We found that cell division did not occur at specific sites in the tubule. A typical result is shown in Figure 6, which displays three confocal sections (taken at intervals of 4 μm) of the same branching tubule, which contained several mitotic cells. Of these mitotic cells, one was at the tip of tubule (cell a), two at the base (cells d and e, both of which are late in mitosis), and several were located in the middle of the tubule (cells b, c, and f). This result demonstrates that the tubules can grow by cell proliferation along the whole tubule.

Orientation of Cell Division during Chain Formation
The orientation of cell division plays an important role in morphogenesis in several systems. This has mainly been studied in genetically tractable, such as yeast, Caenorhabditis elegans, and Drosophila (Jan and Jan, 2001; Knoblich, 2001; Wodarz, 2002). Oriented division also plays a role in neurogenesis in the mammalian brain and during wound healing (Chenn and McConnell, 1995; Heins et al., 2001; Estivill-Torres et al., 2002; Song et al., 2002). We investigated whether the orientation of cell division could play a role during tubulogenesis. We concentrated on the initial step in formation of chains, in which a cell first leaves the monolayer of the cyst wall.

The orientation of the spindle and the axis of division during mitosis of MDCK cells grown on filters has been reported previously (Reinsch and Karsenti, 1994). To summarize, the centriole in interphase MDCK cells is located just beneath the center of the apical surface. This centriole duplicates and in early prophase, one of the centrioles moves to the basal side of the nucleus. By early prophase, the early spindle forms, with its axis parallel to the apical-basal axis of the cell, i.e., perpendicular to the plane of the monolayer. In prometaphase, the spindle starts a rotation of 90°, so that by metaphase the spindle is now parallel to the plane of the monolayer. The spindle remains in this position until mitosis is completed, so that both daughter cells remain in the plane of the monolayer.

We first analyzed spindle orientation in dividing cells in cysts that were not treated with HGF. (To follow this in live cells, we established a stable MDCK cell line expressing GFP-α-tubulin and collected time-lapse images by confocal microscopy.)

In all cells we analyzed, the spindle rotates by 90° from parallel to the apical-basal axis (Figure 7A, 0°) to parallel to the plane of the monolayer (Figure 7A, 15°), much like cells on a filter. The spindle remains parallel to the monolayer until division is complete.

In contrast, HGF-treated cysts displayed a wide range of degree and type of rotation. Some cells behaved much like untreated cysts, i.e., they underwent a 90° rotation in spindle axis, so that both daughter cells remained in the plane of the monolayer. (Figure 7B).

However, in some HGF-treated dividing cells the spindle did not undergo a 90° rotation and one of the daughter cells apparently left the plane of the monolayer. For instance, in Figure 7C, the spindle failed to rotate from its apical-basal orientation but did undergo small excursions. In other cases (Figure 7D), the spindle underwent larger excursions, e.g., seesaw movement, oscillating back and forth around the apical-basal axis. At 0 min the spindle was parallel to the apical-basal axis, as indicated by the solid line in Figure 7D. The spindle first rotated away from the apical-basal axis in a counterclockwise direction, with a maximum excursion of ~70° at 7 min. However, the spindle then rotated in the opposite direction, reaching ~45° rotation in the opposite direction at 25 min. The spindle then apparently rotated partially out of the plane of the section. It should be kept in mind that the spindle may be rotating in three dimensions, and so only part of its rotation may be evident in the our time-lapse images, which are all in one plane of focus. Pronounced three-dimensional rotation has been seen also in the developing mammalian brain (Adams, 1996).

Taken together, these data are consistent with the hypothesis that in the absence of HGF, a spindle rotation mechanism normally ensures that both daughters remain in the monolayer. In contrast, HGF treatment causes an apparently variable loss of this mechanism.

We also examined the effect of Y27632 on spindle orientation in cysts without HGF. We found that the effects to be variable. In Figure 8A, a spindle is observed at 0 min oriented parallel to the monolayer, and this rotates to perpendicular to the monolayer by 25 min. This cell seems to be delayed in mitosis, which is not completed by 86 min. Figure 8B shows a very different pattern, where the spindle seesaws approximately perpendicular to the monolayer (16 and 36 min) and eventually divides, with the cell possibly leaving the monolayer (60 min). It is possible that these variable defects in spindle orientation could account for the effects we saw of Y27362 alone on cyst morphology, such as the spikes seen at 24 h. However, these spikes largely do not seem to involve cells that have divided, but rather seem more likely due to changes in the actin cytoskeleton. These spindle orientation defects may also account, at least in part, for some the effects seen with combined treatment with Y27632 and HGF, such as the isolated cells seen at long time points, and perhaps even the increased number of extensions (though these were more likely produced without mitosis).

DISCUSSION
Our basic result is that MDCK cells in collagen exhibit two types of polarization and movement, and that HGF induces the cells to switch from one type to the other as the cells make the transition from extensions to chains. Cells with extensions resemble the apparently nonmotile polarized ep-
The epithelial cells in the monolayer in that they still have apical-basolateral polarity. However, these cells have a type of movement, i.e., they protrude a pseudopodial extension. In contrast, MDCK cells in chains have lost almost all remnants of epithelial polarity and instead resemble other typical migrating cells, such as migrating fibroblasts.

PI3K is required for both the initiation and maintenance of extensions. As was observed in the leading edges of fibro-
Figure 7. Time-lapse images of orientation of mitotic spindle. All images are of cysts expressing GFP-tubulin, taken by time-lapse confocal microscopy. (A) Untreated cyst. The mitotic spindle is oriented in the apical-basal axis at 0 min (dotted line), rotates 90° to the plane of the monolayer by 15 min, and remains in this orientation through the completion of mitosis. (B) Cyst treated with CM. The mitotic spindle is oriented in the apical-basal axis at 0 min (dotted line) and rotates to the plane of the monolayer by 9 min. In this case, the spindle axis remains in this orientation through the completion of mitosis and the cell does not leave the monolayer. (C) Cyst treated with CM. This spindle exhibits little rotation and remains parallel to the apical-basal axis. (D) Cyst treated with CM. This cyst exhibits seesawing motion of spindle. The spindle is oriented in the apical-basal axis at 0 min (solid line). It first rotates counterclockwise to 70° at 7 min (dotted line). The spindle then rotates clockwise, passing through the apical-basal axis at 17 min and reaches 45° in the opposite direction at 25 min. Eventually, the spindle rotates out of the plane of focus.
blasts, neutrophils, and Dictyostelium, lipid products of PI3K accumulate in HGF-induced extensions. Moreover, inhibition of PI3K activity with LY294002 blocks extension formation. Presumably, PI3K plays a similar role to that in neutrophils and Dictyostelium, where it provides an intracellular signal for the direction of migration. However, not all cell migration depends on PI3K, e.g., the long pseudopodial extensions that astrocytes form in direction of migration during wound healing resemble MDCK extensions (both are long, large, and relatively slowly moving) but are not affected by PI3K inhibitors (Etienne-Manneville and Hall, 2001).

Cells that change direction quickly, e.g., neutrophils, do not rely on the relatively slow polarization of the Golgi or MTs (Schliwa and Hörner, 1993) but rather depend on rearrangements of the actin cytoskeleton. In contrast, cells that form pseudopodia slowly and move slowly, e.g., astrocytes, rely more on MTs for movement, and astrocyte polarization and pseudopod formation do not depend on actin polymerization (Etienne-Manneville and Hall, 2001). MDCK extension formation is, like pseudopod formation in astrocytes, much slower than neutrophil migration (time scale of hours rather than minutes). It is therefore surprising that extension formation does not rely on MT dynamics or Golgi reorientation but does depend on actin polymerization.

From this viewpoint, extensions somewhat resemble neutrophils, because they require PI3K and actin, but do not involve Golgi reorientation or MT. In contrast, chain formation involves Golgi reorientation, and thus resembles fibroblasts and astrocytes. It was difficult to assess the role of PI3K and polymerized actin in chains, because their inhibitors led to a collapse of extensions, which are the precursors of chains. It was also hard to assess the role of MT dynamics in chains, because the nocodazole inhibits mitosis, which is necessary for chain formation.

The differences between formation of extensions and chains have in vivo parallels. In the development of salivary glands and trachea in Drosophila, cells first send out pseudopodial extensions, and then the entire epithelial cell moves. Mutation of the ribbon gene does not prevent extension formation by these developing gland cells but does prevent movement of the cell bodies, which may be analogous to chain formation in the MDCK system (Bradley and Andrew, 2001; Shim et al., 2001).

ROCK Controls the Number and Length of Extensions

It is not known what controls the number or length of extensions or tubules. Even though cysts are uniformly exposed to HGF and all of the cells are theoretically capable of responding, only a minority of cells in the cyst produce extensions and then tubules. Y27632 greatly increased the number and length of HGF-induced extensions and produced very small extensions on its own. These effects are compatible with previous data that activation of Rho inhibits HGF-induced scattering of MDCK cells (Ridley et al., 1995). Inhibition of ROCK also produces extensions (neurites) in neuroblastoma cells and pseudopodia in fibroblasts (Hirose et al., 1998) and increases the rate of fibroblast movement in a wound-healing assay (Nobes and Hall, 1999). Our data suggest that normally ROCK (and presumably the contractility and stress fibers that it controls) inhibits movement and pseudopod formation in a number of systems, including MDCK cell extensions.

ROCK activity leads to the production of P-MLC, which in turn promotes actin-myosin contractility. We find that in cells in the monolayer of the cyst wall, most of the P-MLC underlies the basal surface, with some in the region of the tight junction. During extension formation the P-MLC remains underneath the elongated basal surface. However, a completely different localization of P-MLC is seen in chains, where most of the P-MLC is diffuse in the cytoplasm, especially the trailing region of the cytoplasm. This trailing location is similar to that seen in migrating neutrophils, where P-MLC is predominantly located in the tail or uropod of the migrating neutrophil (J. Xu, F. W. and H. B., unpublished). Indeed, ROCK is needed for detachment of the uropod in migrating leukocytes (Albl et al., 2001; Worthylake et al., 2001).

Ordinarily, in response to HGF, cells move centripetally away from the cysts, whereas with Y27632, some cells move toward the center. This may be due to disruption of the mechanism that normally causes cells to move in only one direction, toward the source of HGF. Intriguingly, a similar effect is seen with Y27632 treatment of neutrophils, which produces multiple pseudopods in random directions (Wang and Bourne, unpublished data). We cannot exclude other possibilities, e.g., that Y27632 makes tight junctions leaky, allowing cells to chemotax toward HGF that has leaked into the lumen (Walsh et al., 2001).
Role of Cell Division in Tubulogenesis

Most tubulogenesis in vertebrates involves cell proliferation (Hogan and Kolodziej, 2002). Consistent with that, we found that inhibition of MDCK cell division blocked chain formation, though extensions were not affected.

Orientated cell divisions play a crucial role in many steps in development. Cells in the wall of unstimulated cysts behave similarly to those grown on a filter support. The spindle forms parallel to the apical-basal axis and then consistently rotates 90° to be parallel to the plane of the monolayer. In striking contrast, with HGF treatment the rotation was highly variable. Some spindles underwent almost no rotation, some rotated normally, and some had large seesaw excursions. A similar type of seesaw motion has been observed in oriented cell division in Drosophila melanogaster (Lu et al., 2001; Roegiers et al., 2001). Our results suggest that the mechanism that normally rotates the spindle and produces coplanar division is variably lost with HGF treatment. This causes some daughter cells to leave the monolayer and perhaps initiate or contribute to a chain.

Superficially, at least, the reorientation of cell division in MDCK cells resembles the orientation of cell division in Drosophila neurogenesis. Neuroblasts arise from an epithelium, where some cells divide in the plane of the monolayer and thereby remain in this monolayer. Recent evidence indicates that the orientation of the spindle is controlled by a hierarchical mechanism (Lu et al., 2001; Roegiers et al., 2001). The default situation in the Drosophila system is for the spindle to orient in the apical-basal direction. A superimposed signal involving the adherens junction can orient the spindle parallel to the plane of the monolayer. When viewed in this context, our data suggest a model wherein normal division of MDCK in a monolayer, the spindle rotates 90° because it is captured into a position parallel to the monolayer. This capture mechanism may involve the interaction of astral microtubules with a protein associated with the adherens junctions. We suggest that during HGF-induced tubulogenesis, this normal capture mechanism may be abrogated in some (but not all) cells. The result could be that the spindle is not anchored in any particular orientation and in some cases can undergo a seesaw motion.

MDCK tubulogenesis may provide an accessible system to dissect the mechanisms of orientation of cell division in mammalian cells. Proteins that have been suggested to play roles in controlling orientation of the spindle and cell division in mammalian cells and are therefore possible targets for HGF-induced reorientation of division include LIS1 (Faulkner et al., 2000), PINS (partner of inscuteable), NuMA (Du et al., 2001), dynein, EB1, and APC (adenomatous polyposis coli) (Dujardin and Vallee, 2002; Gundersen, 2002). We also found that inhibition of ROCK leads to variable defects in spindle orientation and rotation, suggesting that ROCK may be involved in this process. Our results raise the possibility that orientation of cell division may play a role in tubulogenesis, and therefore is potentially important both during development and during tubulogenesis in adults, which occurs during normal physiology (e.g., cycling of mammmary andendometrial glands), regeneration (e.g., after renal tubular necrosis), and pathological proliferation of tubules (e.g., hypersecretory lung diseases).

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REFERENCES


