Cell shape and contractility regulate ciliogenesis in cell cycle–arrested cells

Amandine Pitaval, Qingzong Tseng, Michel Bornens, and Manuel Théry

In most lineages, cell cycle exit is correlated with the growth of a primary cilium. We analyzed cell cycle exit and ciliogenesis in human retinal cells and found that, contrary to the classical view, not all cells exiting the cell division cycle generate a primary cilium. Using adhesive micropatterns to control individual cell spreading, we demonstrate that cell spatial confinement is a major regulator of ciliogenesis. When spatially confined, cells assemble a contractile actin network along their ventral surface and a protrusive network along their dorsal surface. The nucleus–centrosome axis in confined cells is oriented toward the dorsal surface where the primary cilium is formed. In contrast, highly spread cells assemble mostly contractile actin bundles. The nucleus–centrosome axis of spread cells is oriented toward the ventral surface, where contractility prevents primary cilium growth. These results indicate that cell geometrical confinement affects cell polarity via the modulation of actin network architecture and thereby regulates basal body positioning and primary cilium growth.

Introduction

Tissue development and renewal rely on the spatial and temporal regulation of the balance between cell growth and quiescence. Most cells in tissues are quiescent. In vertebrates, these cells generally possess a primary cilium (Gerdes et al., 2009). It is widely considered that ciliogenesis is tightly coupled to the cell cycle (Wheatley, 1971; Santos and Reiter, 2008). In postmitotic cells entering the G0 phase, the mother centriole differentiates into a basal body, anchoring itself to the plasma membrane and growing the cilium (Alieva and Vorobjev, 2004). When quiescent cells resume cell cycle, the cilium is resorbed. According to this classical view, quiescent cells are ciliated, whereas nonciliated cells are engaged in the cell cycle. However, several examples of differentiated, but not ciliated, cells in situ suggest that cell cycle exit might not be sufficient to induce ciliogenesis (Seeley and Nachury, 2010). The parameters regulating ciliogenesis in growth-arrested cells remain to be elucidated.

It has been concluded from experiments on cultured cells that cell cycle arrest and cell confluence are required for ciliogenesis (Wheatley et al., 1994). Primary cells progressively stop cycling after a limited number of passages, and immortalized cell lines stop cycling upon serum starvation and deprivation of growth factors. In both cases, a high level of confluence seems necessary to grow the primary cilium (Alieva and Vorobjev, 2004). However, because contact inhibition in confluent cells is known to induce cell growth arrest, it is not clear whether confluence affects ciliogenesis by promoting cell quiescence or by playing a more direct role on ciliogenesis. We designed specific tools and methodology to control separately the degree of cell confluence and cell cycle exit. We then analyzed the parameters governing ciliogenesis in cell cycle–arrested cells.

Results and discussion

Human retinal pigment epithelial (RPE1) cells are diploid immortalized cells that maintain normal checkpoints on cell cycle progression as well as contact inhibition of growth. We confirmed the role of cell confluence in ciliogenesis in these cells stop cycling after a limited number of passages, and immortalized cell lines stop cycling upon serum starvation and deprivation of growth factors. In both cases, a high level of confluence seems necessary to grow the primary cilium (Alieva and Vorobjev, 2004). However, because contact inhibition in confluent cells is known to induce cell growth arrest, it is not clear whether confluence affects ciliogenesis by promoting cell quiescence or by playing a more direct role on ciliogenesis. We designed specific tools and methodology to control separately the degree of cell confluence and cell cycle exit. We then analyzed the parameters governing ciliogenesis in cell cycle–arrested cells.

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Supplemental material can be found at: http://doi.org/10.1083/jcb.201004003
confluent cells had a cilium, whereas only a small proportion of dispersed cells had one (Fig. 1A). We checked by video microscopy during 48 h that most cells had actually stopped dividing at all densities. This demonstrated that cells could exit the cycle by plating RPE1 cells at various densities. Coverslips were then fixed at various time points after serum starvation and stained for primary cilium (Alieva et al., 1999). Few cells had primary cilium soon after starvation. However, 48 h later, a majority of confluent cells had a cilium, whereas only a small proportion of dispersed cells had one (Fig. 1A). We checked by video microscopy during 48 h that most cells had actually stopped dividing at all densities. This demonstrated that cells could exit the cycle.
Figure 2. Cell cycle exit in G1. (A) Cells were plated on large, fibronectin-coated discoidal micropatterns to prevent them from escaping the observation field. They were monitored by time-lapse microscopy to measure cell division time with respect to the time of starvation and to follow the fate of daughter cells. The red cell illustrates starvation in early G1: the cell divided 2.5 h before starvation and stopped dividing after. The blue cell illustrates starvation in late G1: the cell divided 5 h before starvation and divided again after. See corresponding Video 1 and Fig. S1. Bar, 100 µm. (B) The histogram shows the proportion of dividing and nondividing cells, depending on the cell cycle stage when starvation occurred (S, n = 31; G2, n = 31; early G1 [<2 h after mitosis], n = 92; and late G1 (>5 h after mitosis), n = 130). (C) Experimental procedure used for the following experiments. Cells were synchronized in early G1, plated on a micropatterned coverslip with 10 different sizes of discs ranging from 500 to 3,500 µm², and serum starved. Bar, 50 µm.

(A) t=0, t=14h, t=56h

(B) S, G2, early G1 (<2h), late G1 (>5h)

(C) double thymidine block + trypsin

(synchronization in early G1) plating on micropatterns starvation

(D) t = 0, t = 40h

(E) without serum with serum

Ki67 F-actin, hoechst Ki67 F-actin, hoechst

3000 µm² 3000 µm²

750 µm² 750 µm²

on August 20, 2017 jcb.rupress.org Downloaded from
with or without growing a primary cilium (Fig. 1B) and that specific parameters associated with cell confluence seemed to regulate this fate.

We further investigated the role of cell spatial confinement and the presence of cell–cell contact on ciliogenesis by plating individual cells on micropatterned surfaces of defined sizes. To analyze cell cycle exit and ciliogenesis in these conditions, it was necessary to determine the exact time window in which cells could be plated on micropatterns and serum starved to obtain cell cycle exit before any further division. Therefore, we investigated the cell response to serum starvation all along the cell division cycle (Fig. 2A and Fig. S1). As expected from previous studies (Pardee, 1974; Zetterberg and Larsson, 1985), most cells starved in early G1 did not reenter the cell cycle (Fig. 2B). We thus proceeded as follows: cells were synchronized in early G1, plated on individual micropatterns, and starved. The role of cell confinement was tested by using an array of discoidal micropatterns of various sizes ranging from 500 to 3,500 µm² (Fig. 2C).

Because cell spreading has been shown to positively regulate cell cycle progression (Chen et al., 1997; Huang et al., 1998; Schwartz and Asoian, 2001), we checked whether RPE1 cells actually left the cell cycle on all micropattern sizes. Just after spreading, cells were starved and monitored by video microscopy. Cells on small or large micropatterns left the cell cycle in similar proportions (Fig. 2D and E). Control non-starved cells kept on dividing. We concluded that cell cycle exit was specifically induced by serum starvation independently of the extent of cell spreading.

Dramatic differences in the ciliation rate 48 h after serum starvation were observed, depending on the size of the micropatterns. Although cilia were present on most cells confined on small micropatterns, they became less frequent as cell spreading increased (Fig. 3A and B). Most of the highly spread cells had exited cell cycle without growing a cilium. Instead, these cells displayed a large network of acetylated microtubules in their cytoplasm. In addition, the mean primary cilium length decreased from 5 to 2 µm as cell spreading increased (Fig. 3C). Long cilium, >6 µm, could only be observed in spatially confined cells. These differences were already present 24 h after starvation and maintained for at least 72 h (Fig. 3D, E and D), suggesting they were not caused by a delay in basal body maturation (Anderson and Stearns, 2009). When two cells were found on the same micropattern, each cell adopted half of the area, and their ciliation rate matched that of individual cells on micropatterns twice as small (Fig. 3B). Therefore, cell–cell contact did not seem to be required for, nor regulate, ciliogenesis in these cells. Until now, primary cilium growth defects in cell cycle–arrested cells had only been seen in cells in which basal body integrity (Graser et al., 2007; Mikule et al., 2007), centrosome-associated cell cycle regulation (Spektor et al., 2007), or intraflagellar transport (Schneider et al., 2005) required for ciliogenesis was specifically impaired. Our results demonstrate a strong dependency of ciliogenesis on the degree of cell shape extension in wild-type cells.

The degree of cell spreading affects actin network architecture. As cell spreading increased, cells displayed larger transversal stress fibers and more circumferential bundles (Fig. 3A). Because actin dynamics have been shown to affect ciliogenesis (Kim et al., 2010), we tested whether these actin structures were involved in the cell shape–associated regulation of primary cilium growth by disrupting actin network integrity with 1 µM cytochalasin D. Several large cells managed to maintain their spreading, whereas their actin network was damaged. Interestingly, after a 24-h starvation in the presence of cytochalasin D, most cells, regardless of their size, were ciliated (Fig. 4A). In addition, cilium length was no longer reduced in large cells; instead, large cells had slightly longer cilia. The actin network, therefore, appeared fully responsible for the regulation of ciliogenesis in response to cell shape extension.

We then looked for more specific actin structures that could be implicated in this regulation. The location of myosin II strongly affects actin network architecture and thus plays a key role in the establishment of cell polarity at all stages of organism development (St Johnston and Ahringer, 2010). In epithelial cells, ezrin is restricted to the apical pole where, upon phosphorylation, it links the polymerizing actin meshwork in membrane protrusion to the plasma membrane (Dard et al., 2004; Fievet et al., 2004). After 24 h of serum starvation, RPE1 cells confined on small micropatterns displayed a polarized actin network with myosin II–positive actin bundles at their ventral surface and an ezrin-rich actin meshwork at their dorsal surface (Fig. 4B and Fig. S2). In extended cells, the actin network architecture appeared less polarized. Ezrin staining at the cell dorsal surface appeared fainter and more dispersed. Most actin structures were strongly decorated with myosin II (Fig. 4B and Fig. S2). Indeed, cell shape extension has been correlated to high levels of actomyosin contraction (McBeath et al., 2004; Polte et al., 2004). Because up-regulation of RhoA in response to the loss of meckelin, a transmembrane protein of the primary cilium, has been shown to induce the formation of stress fibers and prevent primary cilium growth (Dawe et al., 2009), high levels of contraction could be responsible for the impaired ciliogenesis in spread cells.

To test this hypothesis, cell contractility was inhibited with either 10 µM Y27632, to inactivate Rho kinase, or with 50 µM blebbistatin, to inactivate myosin II ATPase. Both treatments induced actomyosin relaxation of spread cells as revealed by the large reduction of phosphorymosin II staining (Fig. 4B). Interestingly, the proportion of ciliated cells in relaxed spread cells
Figure 3. Cell confinement and ciliogenesis. The experimental procedure for all panels is described in Fig. 2 C. (A) Cells on various micropattern sizes were serum starved for 48 h, fixed, and stained for F-actin (top) and acetylated tubulin (bottom with an inset magnification) to reveal the primary cilium. Two examples of cells on 500, 1,500, and 3,500 µm² are shown to illustrate that most cells on 500 µm² had a primary cilium, half had one on 1,500 µm², and most cells on 3,500 µm² had no primary cilium. Bars, 20 µm. Inset bar, 4 µm. (B) The proportion of individual ciliated cells on 10 different micropattern sizes (n > 100 for each) is shown with the black curve. The proportion of ciliated cells in cell doublets (two cells per pattern) on 10 different micropattern sizes (n > 100 for each) is shown with the gray curve with respect to individual cell size. (C) Quantification of primary cilium length depending on micropattern size. Primary cilium length was shorter in larger micropatterned cells. (D) Quantification of ciliated cell frequency in cells plated on small (750 µm²) and large (3,000 µm²) micropatterns over time (n > 170 for each). The rate slightly increased from 24 to 48 h and then reached a plateau. (E) Quantification of primary cilium length on small (750 µm²) and large (3,000 µm²) micropatterns 72 h after serum starvation. Primary cilium length was similar to the one measured 24 h after serum starvation (compare with C). *, P < 0.05 and ***, P < 0.001. Plotted bars represent standard deviations. Horizontal bars represent mean values.
Figure 4. Actin network assembly, contractility, and ciliogenesis. The experimental procedure for A and B is described in Fig. 2 C. RPE1 cells were plated on micropatterns (from 750 to 3,000 µm²) and serum starved for 24 h. (A) Micropatterned cells were treated with cytochalasin D during starvation. Cells were fixed and stained for acetylated tubulin (red), F-actin (green), and DNA (blue). Primary cilium occurrence (top, n = 82, 97, 82, and 41) and length (bottom, n = 30, 26, 27, and 25) were measured in cells whose shape still covered the entire micropattern after the treatment. Bar, 5 µm. (B) Micropatterned cells were treated with either Y27632 or blebbistatin during starvation. Cells were then fixed and immunolabeled for phospho-ezrin (top) and phosphomyosin II (bottom). Z acquisitions were performed and projected on a single image containing the maximal intensity of each pixel. Several images on each micropattern were averaged and color coded with the fire look-up table to highlight intensity variations. Bar, 5 µm. Primary cilium occurrence was measured in each condition (n > 400 for each). (C) RPE1 cells in early G1 were plated at low density on hard substrate (polystyrene-coated glass coverslips) or soft substrate (polyacrylamide gel grafted on glass coverslips). Cells were then serum starved for 48 h, fixed, and stained for F-actin (red), DNA (blue), and acetylated tubulin (green). Most cells on hard substrates had no primary cilium (left). A majority of cells on soft substrates were ciliated (right). Primary cilium occurrence was measured at a cell density of 200 cells/mm² on hard substrates (n = 300) and on soft substrates (n = 1,140). Primary cilium length was shorter on hard substrates than on soft ones. *, P < 0.05; **, P < 0.01; and ***, P < 0.001. Plotted bars represent standard deviations. Horizontal bars represent mean values. Bar, 20 µm.
was twice that of control spread cells in both cases (Fig. 4 B). It is noteworthy that no significant effect could be detected on mean primary cilium length (control: 2.9 ± 1.5 µm, n = 20; blebbistatin: 2.9 ± 1 µm, n = 30; and Y27632: 3.6 ± 1 µm, n = 30).

To further confirm the negative regulation of cell contraction on ciliogenesis, we plated cells on soft substrates to release cell contraction (Polte et al., 2004; Engler et al., 2006; Solon et al., 2007). Cells displayed large and straight stress fibers on hard substrates but thinner and looser bundles on soft substrates (Fig. S2 D). Strikingly, although only 25% of sparsely populated cells had a primary cilium on hard substrate (polystyrene), almost 80% were ciliated on soft substrates (polyacrylamide gel; Fig. 4 C). In addition, cilia were longer. Altogether, these results demonstrated that, in quiescent cells, the high level of contraction associated with cell spreading hindered ciliogenesis.

Rho kinase is also known to regulate centrosome positioning (Chevrier et al., 2002). Because early stages of ciliogenesis rely on basal body migration and docking to the plasma membrane, we wondered whether actin architecture and myosin II activity could play a role in basal body positioning. After serum starvation, the basal body was found to migrate above was twice that of control spread cells in both cases (Fig. 4 B). It is noteworthy that no significant effect could be detected on mean primary cilium length (control: 2.9 ± 1.5 µm, n = 20; blebbistatin: 2.9 ± 1 µm, n = 30; and Y27632: 3.6 ± 1 µm, n = 30).

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the nucleus, to the ezrin-rich cell dorsal surface of confined cells, where they could grow a primary cilium. However, in starved spread cells, basal bodies were found below the nucleus, in contact with the ventral surface where they rarely formed a primary cilium (Fig. 5, A–C and Fig. S3 B). Dorsal positioning of basal body and ciliogenesis appeared to be tightly coupled (Fig. S3 D). After cytochalasin D treatment, basal bodies adopted a dorsal position, and ciliogenesis was improved (Fig. S3, B and D). After Rho kinase inhibition in confined cells, basal bodies were positioned below the nucleus, and ciliogenesis was reduced (Fig. S3, B and D). Inhibition of cell contractility in confined cells did not affect cilium length (control: 5 ± 1.5 µm, n = 60; Y27632: 5.3 ± 2 µm, n = 37; and blebbistatin: 5.1 ± 2.5 µm, n = 34). As previously suggested from RhoA inhibition in multiciliated cells (Park et al., 2006; Pan et al., 2007), this showed that Rho kinase activity is necessary for the apical positioning of the basal body and the subsequent growth of the primary cilium. However, in extended cells, Rho kinase inhibition did not affect the ventral positioning of basal bodies but promoted the growth of ventral cilia. This showed that cell contractility acts separately on both cilium extension and basal body positioning.

In this study, we thus report a major role for physical parameters such as spatial confinement and substrate rigidity, which, through their effect on actin cytoskeleton architecture, regulate ciliogenesis at cell cycle exit (Fig. 5 D). When individual cells were sufficiently spatially confined on adhesive substrate, they could assemble both a contractile ventral domain and a protrusive, ezrin-rich dorsal domain mimicking the apical-basal polarity of epithelial cells. This polarization was further transmitted to the internal cell organization with the nucleus–centrosome axis oriented toward the dorsal surface in a Rho kinase–dependent manner. Under these conditions, most cells assemble a primary cilium. When individual cells were highly spread, they mostly formed a highly contracted actin network in contact with their adhesive substrate. In this case, the internal polarity was reversed: the nucleus–centrosome axis was oriented toward the ventral surface. In this position, below the nucleus, and in proximity with stress fibers, the centrosome could not induce the growth of the primary cilium. This was partially a result of the local contractility of the cell cortex. Overall, these results demonstrate that cell cycle exit is not sufficient to induce primary cilium growth. Depending on the global organization of actin cytoskeleton architecture and dynamics, cells can enter a quiescent state with or without assembling a primary cilium.

Whether such a regulation occurs during development could have far-reaching consequences given the sensory functions of primary cilia.

Materials and methods

Cell culture

Human telomerase-immortalized, retinal-pigmented epithelial cells (hTERT-RPE1; Takara Bio Inc.) were grown at 37°C and 5% CO2 in DME/F12 (Invitrogen) containing 10% fetal calf serum, 2 mM glutamine, 100 U/ml penicillin, and 100 U/ml streptomycin. We used a double thymidine block to synchronize cells at the G1/S transition. Cells in a culture flask were treated with 5 mM thymidine for 16 h, released for 8 h with normal culture medium, and then were treated again with thymidine for 16 h.

To detect cilium incidence of G1-enriched cells on micropatterns, cells were trypsinized just after mitosis (10 h after release of G1/S synchronization) and plated on the micropatterned coverslip (10,000 cells/cm²) to obtain a majority of patterns with individual cells. Medium was gently flushed over cells and replaced with fresh medium 0.5–1 h later to remove nonattached cells in suspension. Serum was then removed 1 h later, and cells were cultured for 24–72 h.

Cytoskeletal drugs were added to the medium when the serum was removed, i.e., 2 h after the plating of cells on micropatterns. For Rho kinase inhibition, Y27632 was added at 10 µM. For myosin II ATPase inhibition, blebbistatin was added at 50 µM. For actin network destabilization, cytochalasin D was added at 1 µM 2 h after serum removal to allow starved cells to fully spread on large micropatterns before being treated.

Micropatterning

Glass coverslips were first spin coated with an adhesion promoter (T1 Prime; MicroChemicals) and then with 0.5% polystyrene dissolved in toluene at 3,000 rpm for 30 s. The polystyrene layer was further oxidized with an oxygen plasma treatment (FEMTO; Diener Electronic) for 10 s at 30 W and incubated with 0.1 mg/ml polylysine polyethylene-glycol (JanKem Technology) in 10 mM Hepes, pH 7.4, at room temperature for 1 h. Coverslips were then dried by spontaneous dewetting. Poly-ethylene-glycol–coated slides were placed in contact with an optical mask containing the transparent micropatterns (Toppan Photomasks, Inc.) using a home-made vacuum chamber and exposed for 3 min to deep UV light (UV Cleaner; Jelget Company). Micropatterned slides were washed once in PBS and finally incubated for 30 min with a solution of 20 µg/ml bovine fibronectin solution (Sigma-Aldrich) and 5 µg/ml Alexa Fluor 646– or Alexa Fluor 542–labeled fibrinogen (Invitrogen). Before plating cells, patterned coverslips were washed three times with sterilized PBS. We also used commercially available micropatterned glass coverslips (CYTOOChips, Cytoq).

Soft substrate fabrication

Acrylamide monomer solution (Sigma-Aldrich) containing 5% acrylamide and 0.13% bis acrylamide was polymerized by ammonium persulfate (Sigma-Aldrich) and N,N,N,N-tetramethylethylenediamine (Sigma-Aldrich) over a glass coverslip previously silanized with 3-(trimethoxysilyl)propyl methacrylate (Sigma-Aldrich). 25 mM 4-benzoylbenzyl-trimethylammonium chloride (benzophenone; custom synthesis by Sigma-Aldrich outsourced to LaboTest) and fibronectin at 40 µg/ml in PBS were added on the surface of the polymerized gel and then exposed to UV light (Delolux G3S; Supratec) at 400 W for 45 s to graft the fibronectin using the photoactivation of benzophenone (Fink et al., 2007). The gel was washed with PBS before cell seeding.

Video microscopy and cell imaging

Time-lapse acquisitions were taken with an inverted microscope (Axiovert 200M; Carl Zeiss, Inc.). The temperature, CO2, and humidity control were performed using a Box and Brick system (Life Imaging Services). Multiple positions were recorded using an XY motorized stage (Marzhauser) with a 10-min time frame over 40–54 h with a dry 20x objective (NA = 0.4) or with a dry 40x objective (NA = 0.9). Fluorescent images shown are maximal projections of Z stacks acquired with oil immersion objectives at either 63x or 100x (NA = 1.4) mounted on a piezo ceramic (Physics Instruments). Both microscopes were controlled with Metamorph software (MDS Analytical Technologies). Side views were obtained using XZ optical sections (not projections) of either classical epifluorescence or Z stacks with and incubated with confocal Z stacks (1.5 planes separated by 500 nm) acquired with a confocal microscope (TCS-SP2; Leica) through a 63x objective (NA = 1.4).

Immunostaining

For immunostaining experiments, cells were plated on polystyrene-coated glass coverslips to promote cell attachment and keep a good optical quality. Coating was performed by spin coating 0.5% polystyrene solution to tolune at 3,000 rpm for 30 s. The polystyrene layer was further oxidized with an oxygen plasma treatment. Alternatively, cells were plated on micro-patterned or soft substrates.

For actin-associated protein and primary cilium immunostaining, RPE1 cells were fixed at 4% paraformaldehyde in cytoskeleton buffer, pH 6.1, for 1.5 min at room temperature without any prior PBS wash. They were then rinsed twice with PBS and incubated in 0.1 M ammonium chloride in PBS for 10 min. Cells were then permeabilized in 0.1% Triton X-100 in PBS.
for 3 min and blocked with 3% BSA in PBS for 30 min. For phosphorysomycin immunostaining, cells were prepermeabilized for 15 s with 0.1% Triton X-100 in cytoskeleton buffer before paraformaldehyde fixation. For centrosome detection, cells were fixed for 5 min in methanol at -20°C. For Ki67 immunostaining, cells were fixed with 4% paraformaldehyde in PBS, pH 7.4, for 15 min at room temperature.

Quantification on images and statistical analysis

Fluorescence densities were measured on individual cells. Z stacks containing 20 images separated by 500 nm were projected along the Z axis using the maximal intensity of each pixel. We then quantified the ratio between the total fluorescence intensity and the cell area. The mean value of the maximal intensity of each pixel. We then quantified the ratio between the maximal intensity of each pixel. We then quantified the ratio between the total fluorescence intensity and the cell area. The mean value of the maximal intensity of each pixel. We then quantified the ratio between the total fluorescence intensity and the cell area. The mean value

Image processing

All of the images were processed using ImageJ (National Institutes of Health). For 3D reconstruction, images were first deconvolved using the plugin Extended Depth of Focus was applied to visualize all the cilium images at low magnification (Fig. 1), maximum projection of Z stacks or the plugin

Online supplemental material

Fig. S1 shows the analysis of cell cycle exit. Fig. S2 shows the quantification of ezrin and myosin II densities in confined and extended cells. Fig. S3 shows the labeling and quantification of centrosome position. Video 1 shows the monitoring of cell fate upon serum starvation. Online supplemental material is available at http://www.jcb.org/cgi/content/full/jcb.201004003/DC1.

We would like to thank Annie Delouvée for providing advice about RPE1 cultured through the primary cilium in fibroblasts.

Submitted: 1 April 2010
Accepted: 13 September 2010

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