Direct binding of RalA to PKCζ and its crucial role in morphological change during keratinocyte differentiation

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ABSTRACT During differentiation, keratinocytes undergo a dramatic shape change from small and round to large and flat, in addition to production of proteins necessary for the formation of epidermis. It has been shown that protein kinase C (PKC) ζ is crucial for keratinocyte differentiation. However, its role in this process has yet to be fully elucidated. Here, we show that catalytic activity is not necessary for enlarged and flattened morphology of human keratinocytes induced by overexpression of PKCζ, although it is important for gene expression of the marker proteins. In addition, we identify the small G protein RalA as a binding partner of PKCζ, which binds to the C1 domain, an indispensable region for the morphological change. The binding led activation of RalA and actin depolymerization associated with keratinocyte differentiation. siRNA techniques proved that RalA is involved in not only the keratinocyte differentiation induced by PKCζ overexpression but also normal keratinocyte differentiation induced by calcium and cholesterol sulfate. These results provide a new insight into the molecular mechanism of cytoskeletal regulation leading to drastic change of cell shape.

INTRODUCTION

For unicellular organisms, cell shape change associated with movement, cell division, and mating is fundamental for survival. In multicellular organisms, cell shape varies considerably in response to different, cell division, and mating is fundamental for survival. In multicellular organisms, cell shape varies considerably in response to different cues. For example, neurons develop long neurites in response to several factors and hormones, including nerve growth factor (Wiesmann and de Vos, 2001; Tojima and Ito, 2004; Arimura and Kaibuchi, 2008), and lymphocytes chemotax to chemokines (Stephens et al., 2008). In most cases, dynamic morphological changes are triggered by environmental signals and involve cytoskeletal reorganization mediated by the small-molecular-weight G proteins. It is well known that RhoA, Rac1, and cdc42 induce stress fibers, membrane ruffles, and filopodia, respectively (Narumiya, 1996; Takai et al., 2001). Although how cells generate and maintain their shape and size has been well studied, the precise molecular mechanisms leading to dramatic morphological changes are not fully understood. The terminal differentiation of epidermal keratinocytes involves drastic and dynamic morphological changes, making them a model cell for studying drastic cytoskeletal rearrangement.

The epidermis consists of four layers: basal, spinous, granular, and cornified layers (Fuchs 1990; Koster and Roop, 2007). Once committed to differentiation, the basal cells lose their proliferative
potentia and are differentiated toward cornified cells becoming larger and flatter. Keratinoctyes differentiation involves not only these morphological changes but also a shift in protein expression (Fuchs, 1990; Koster and Roop, 2007). When the basal cells are differentiated to the spinous cells, keratin expression shifts from K5/K14 to K1/K10. Differentiation from spinous to granular is marked by expression of keratinocyte-specific transglutaminase 1 (TGap1), and its substrates, including involucrin, filaggrin, and small proline rich protein (SPR), are expressed. TGap1 covalently cross-links the substrates, resulting in forming the cornified layer. Together the morphological and biochemical changes produce the epidermis, an effective function as a barrier to external surroundings.

Keratinocyte differeniation is partly regulated by protein kinase C (PKC), particularly the transition from spinous to granular cells (Denning, 2004; Dlugosz and Yaspa, 1994; Koster and Roop, 2007). Ten subtypes of PKC have been cloned and classified into three groups based on the structure of their regulatory domain (Nishizuka, 1992; Newton, 2006). Conventional PKCs (cPKC: α, βI, βII, and γ) have two common regions, dioacylglycerol (DAG)-binding C1 domain and Ca⁺⁺-binding C2 domain, in the regulatory domain. Calcium, phosphatidylserine, and DAG are required for their activation. The novel PKCs (nPKC: δ, ε, η, and θ) are activated by DAG, but not by Ca⁺⁺, and atypical PKCs (aPKC: 1 and ζ) are insensitive to both DAG and Ca⁺⁺. Of the 10 subtypes, keratinocytes express at least 5 PKC subtypes: α, δ, ε, η, and ζ (Denning, 2004). Of these, the role of PKCη in keratinocyte differentiation has been well studied (Osada et al., 1990, 1993; Ohba et al., 1998). Overexpression of PKCη induces the expression of TGap1 necessary for differentiation. Cholesterol sulfate (CS), an abundant lipid in the granular layer (Kagehara et al., 1994), specifically activates PKCη (Ikuta et al., 1994) and induces keratinocyte differentiation (Denning et al., 1995). However, the molecular mechanism by which this occurs is still unknown.

Therefore we performed a series of experiments to determine the molecular mechanism of PKCη-induced keratinocyte differentiation. Here, we show that the kinase activity of PKCη is not necessary for the morphological change, although it is very important for gene expression of the marker proteins. The small-molecular-weight G protein RaIA binds to the C1 domain of PKCη and contributes to the morphological change during keratinocyte differentiation induced by not only PKCη but also calcium and CS. This is the first report to show RaIA binds to C1 domain of PKCη and that this interaction results in RaIA activation and actin depolymerization, necessary for keratinocyte differentiation. These findings provide information critical to our understanding of the mechanism regulating cell morphology.

RESULTS
PKCη is important for keratinocyte differentiation, but its kinase activity is not required for the morphological change
To elucidate the molecular mechanisms underlying the PKCη-associated keratinocyte differentiation, we first studied the effect of PKCη overexpression on cell height and area as morphological changes are a hallmark of it. Normal human epidermal keratinocytes (NHEK) expressing green fluorescent protein (GFP)-PKCη were flatter and more spread than cells expressing untagged GFP (Figure 1). Morphological changes were evident by 36 h and more pronounced by 48 h after adenoviral infection (Figure 1A). Before the infection with GFP-PKCη, the average diameter of NHEK was ~46 μm; the diameter increased to 69.3 and 83.2 μm at 36 and 48 h postinfection, respectively (Figure 1B). In contrast, NHEK cells expressing GFP alone were significantly less spread at both time points (55.9 and 64.1 μm at 36 and 48 h, respectively). To quantify enlargement, the number of the cells larger than 90 μm in diameter, which is almost twice as large as untreated cells, was counted and this percentage reported in Figure 1C. For cells expressing PKCη, the percentage of the enlarged cells was 15% at 36 h and 35% at 48 h, whereas that of the control cells was <10%, even at 48 h (Figure 1C). Interestingly, the distribution of PKCη-GFP changed from homogeneous at early time points to punctuate in the flattened cells (Figures 1A and 2A, arrows). This distribution of PKCη-GFP was also seen in CS-treated cells at 24 and 36 h (Figure 1D). Additionally, CS treatment induced the translocation of PKCη-GFP to the plasma membrane at 0.5 min, 12 h, and 24 h (Figure 1D, arrows). Finally, CS stimulation enhanced the effects, increasing the average size of PKCη-overexpressing cells from 69.3 μm to 79.7 μm at 36 h (Supplemental Figure S1), and the percentage of cells larger than 90 μm was ~25% and 33% at 24 and 36 h, respectively (Figure 1C), more than double that of non-CS-treated PKCη-overexpressing cells. These results indicate that CS enhanced the effect of PKCη on the morphology, confirming that PKCη is involved in keratinocyte differentiation.

To determine whether PKCη activity is required for the observed morphological changes, we tested the effects of kinase-negative (KN)-PKCη and the PKCη regulatory domain (RD) on spreading. Compared with cells expressing wild-type PKCη, those expressing KN-PKCη and RD-PKCη were significantly larger at 24 h (Figure 2A). More than 40% of the cells expressing KN-PKCη or RD were larger than 90 μm by 24 h (Figure 2B), compared with <10% for wild-type PKCη (Figure 1, B and C). KN- and RD-PKCη showed punctuate localization in the cytoplasm in addition to the membrane localization at 24 h (Figure 2A). Similarly, overexpression of RD-PKCη enlarged and flattened HaCaT cells (see Figure 7, B and C, later in this article), suggesting that this phenomenon is not unique to NHEK cells. These results demonstrate that PKCη activity is not necessary for the enlargement and flattening of keratinocytes; RD is sufficient.

However, keratinocyte differentiation involves both morphological and biochemical changes. We therefore used quantitative RT-PCR to determine the mRNA levels of marker proteins for keratinocyte differentiation. As reported previously, message levels for keratin 5 decreased in keratinocytes differentiated by calcium and CS, whereas those of keratin 1, involucrin, and TGap1 increased (Figure 2C). In contrast, the levels of these maker proteins did not increase upon expression of RD-PKCη. Notably, keratin 1 and 5 decreased in RD-PKCη-expressing cells relative to control. These results indicate that overexpression of RD-PKCη induces the morphological, but not biochemical changes associated with differentiation. In other words, the two major processes in the differentiation, namely morphological changes and transcriptional regulation, can be distinguished by their sensitivity to PKCη. We therefore focused our study on defining the role of PKCη in the morphological changes associated with keratinocyte differentiation.

PS and C1 domains are important for RD-induced morphological change
The RD of PKCη consists of several domains, including variable region 1 (V1), pseudosubstrate region (PS), conserved region 1 (C1), and variable region 3 (V3). To identify the domain(s) involved in the RD-induced morphological changes, RD mutants lacking each domain were produced and their effects on cell spreading determined. As shown in Figure 3, deletion of V1 domain did not significantly affect the RD-induced morphological changes; more than 30% of the cells overexpressing ∆V1 were flattened and enlarged. Further deletion of PS from ∆V1 (∆V1PS), as well as deletion of C1 and V3 (∆C1V3), abolished the shape change. Deletion of V3 alone (∆V3), or in combination with V1 (∆V1V3), decreased cell enlargement ~50%.
interaction (Colon-Gonzalez and Kazanietz, 2006), we made glutathione S-transferase (GST) fusion proteins of RD or ΔC1 and immobilized them on glutathione–sepharose. Lysates of HaCaT cells were applied to the columns, and the bound proteins were eluted with glutathione, separated by SDS–PAGE, and compared by silver staining. A band of ∼23 kDa was present in the RD but not the ΔC1 lane (Figure 4A, red arrow). Mass spectrometric analysis revealed that RalA was present in this band, and immunoblotting for RalA confirmed its presence only in the RD lane (Figure 4B). RalA is a small GTP protein expressed in many tissues, but its expression in skin has not been reported. Therefore we determined the expression of RalA message and protein in keratinocytes. The results of RT-PCR and Western blotting revealed the presence of RalA in the cultured keratinocytes (Figure 4, C and D). Additionally, immunofluorescent staining revealed RalA expression throughout the epidermis.

Notably, when either PS or C1 was removed (ΔPS and ΔC1), no morphological changes were seen. These results clearly show that PS, C1, and V3 are involved in the RD-induced morphological change; PS and C1 are particularly important. Interestingly, the mutants that did induce the morphological change (e.g., ΔV1 and ΔV3) localized to the plasma membrane like RD and KN-PKCη, but those mutants that did not induce cell enlargement tended to associate with nuclear and Golgi membranes or remain in the cytosol (Figure 3 and Supplemental Figure S2), demonstrating that plasma membrane localization is important to induce the morphological change.

**RalA binds to PKCη through C1 domain**

To identify candidate proteins that bind to the PKCη C1 domain and are involved in the RD-induced morphological change, because some C1 domains are known to be involved in the protein–protein interaction (Colon-Gonzalez and Kazanietz, 2006), we made glutathione S-transferase (GST) fusion proteins of RD or ΔC1 and immobilized them on glutathione–sepharose. Lysates of HaCaT cells were applied to the columns, and the bound proteins were eluted with glutathione, separated by SDS–PAGE, and compared by silver staining. A band of ∼23 kDa was present in the RD but not the ΔC1 lane (Figure 4A, red arrow). Mass spectrometric analysis revealed that RalA was present in this band, and immunoblotting for RalA confirmed its presence only in the RD lane (Figure 4B). RalA is a small GTP protein expressed in many tissues, but its expression in skin has not been reported. Therefore we determined the expression of RalA message and protein in keratinocytes. The results of RT-PCR and Western blotting revealed the presence of RalA in the cultured keratinocytes (Figure 4, C and D). Additionally, immunofluorescent staining revealed RalA expression throughout the epidermis.
bound to GST-RD but not to GST-ΔC1 or unconjugated GST (Figure 5A). Coincident with this, FLAG-RalA was colocalized with RD on the plasma membrane of unenlarged cells (Figure 5B, third row) and at dotty accumulation in the cytoplasm of enlarged cells (Figure 5B, second row). In contrast, there was no evidence of colocalization of RalA and ΔC1 (Figure 5B, bottom rows). Together these results indicate that RalA binds to PKCη via its C1 domain.

RD-PKCη specifically binds to RalA among several small-molecular-weight GTPases, and the binding is independent of activation state of RalA

Ral is a member of the Ras family of small GTPases, and Ral proteins consist of two proteins, RalA and RalB, which are 85% identical. We therefore determined the specificity of the binding. Of the two, RD-PKCη preferentially associated with RalA (Figure 6A). In addition, RD-PKCη showed no binding to Rho family small GTPases RhoA, Rac1, and cdc42. These results suggest that binding of RD-PKCη to RalA is relatively specific.

Like other small GTPases, RalA has two states: activated and inactivated. The former binds to GTP, whereas the later binds to GDP. To determine which state prefers to bind to RD-PKCη, effects of GTP-γ-S or GDP-β-S on the binding were

Specifically, its dotty localization on the plasma membrane was evident (Figure 4E, magnified image). Pull-down assays using purified FLAG-RalA and the GST fusion proteins confirmed that RalA directly

FIGURE 2: Effect of PKCη activity on keratinocyte morphology and mRNA level of marker proteins. (A) Confocal images of NHEK cells expressing KN- or RD-PKCη. The images were taken 24 h after infection. Arrows indicate the enlarged and flattened cell having typical punctuate localization of GFP-PKCη. Bars are 50 μm. (B) Static analysis of the effect of KN- or RD-PKCη on cell size. NHEK cells expressing KN-GFP, RD-GFP, PKCη-GFP, or GFP alone were cultured for 24 or 48 h without any stimulation, followed by the same procedure described in Figure 1C. (C) Effect of overexpression of RD on mRNA level of the marker proteins. The mRNA was extracted from NHEK cells 48 h after the infection with RD or treated with or without 50 μM C5 and 0.12 mM CaCl2 for 48 h. Data points represent mean ± SE of three independent experiments.

FIGURE 3: Localization of the PKCη mutant lacking domain(s) and their effects on the morphological changes. Twenty-four hours after the infection with adenoavir encoding PKCη mutants or GFP alone, NHEK cells were fixed following the same procedure described in Figure 1C. *, P < 0.05; **, P < 0.01 compared with RD. + or – represents localization of the mutants based on the results in Supplemental Figure 2. ++++, strong localization; +, weaker localization; PM, plasma membrane; nm, nuclear membrane.
Importance of the C1 domain for the RalA binding. (A) Direct binding of FLAG-RalA to RD-PKC\(\eta\) was independent of GTP-\(\gamma\)-S or GDP-\(\beta\)-S (Figure 6C). In addition, both mutants of RalA, G23V and S28N, mimicking active and inactive forms, bound to RD-PKC\(\eta\) (Figure 6D). These results demonstrate that the binding of RD-PKC\(\eta\) to RalA is independent of the nucleotide exchange.

RalA is involved not only in the RD-induced morphological change but also in the normal differentiation of keratinocytes induced by calcium and CS

To determine whether RalA is really necessary for the RD-induced morphological changes, we used siRNA to down-regulate RalA in HaCaT cells. siRNA in the amount of 100 nM significantly decreased endogenous RalA expression in HaCaT cells and reduced the RD-dependent cell enlargement (Figure 7, A and B). The average diameter of the cells expressing RD was 101.32 \(\mu\)m, whereas 10 and 100 nM siRNA reduced the diameter to 92.14 and 81.88 \(\mu\)m, respectively (Figure 7B). The percentage of cells larger than 90 \(\mu\)m was reduced from 70–50\% (10 nM) and 30\% (100 nM), respectively (Figure 7C). siRNA treatment, however, did not affect the size of cells expressing GFP alone (Figure 7, B and C). Importantly, control siRNA, which has only four unmatched nucleotides, did not reduce the protein level of RalA, nor did it inhibit the RD-PKC\(\gamma\)-induced morphological changes (Figure 7, A–C). Similar inhibitory effects of siRNA were obtained when NHEK cells were used (Figure 7D). These results demonstrate that RalA is involved in the RD-induced morphological changes. Consistent with this, treatment with GDP-\(\beta\)-S or overexpression of inactive form of RalA inhibited the RD-induced morphological change (Figure 8, A and B), suggesting that RalA is activated during the RD-induced morphological change. Indeed, activation of RalA was detected in the keratinocyte morphological change induced by RD but not \(\Delta\)C1 (Figure 8C).

Furthermore, to determine whether RalA acts in morphological changes under more physiological conditions, we repeated the siRNA down-regulation in NHEK treated with calcium and CS. Down-regulation of RalA expression blocked the morphological changes induced by CS and calcium (Figure 7E). Similarly, siRNA of PKC\(\eta\) inhibited the morphological change induced by CS and calcium (Supplemental Figure S3). These results implicated that binding of endogenous RalA to endogenous PKC\(\eta\) is necessary for the normal keratinocyte differentiation. Interestingly, FLAG-RalA and full-length PKC\(\eta\)-GFP did not colocalize 24 h after infection (Figure 9A, top), when 90\% of the cells are still smaller than 90 \(\mu\)m (Figure 1). Similarly to endogenous RalA, FLAG-RalA was localized on the plasma membrane, whereas full-length PKC\(\eta\) was detected throughout the cytoplasm (Figure 9A, top). In contrast, CS stimulation resulted in extensive colocalization of PKC\(\eta\) and RalA in puncta of enlarged and flattened cells (Figure 9A, bottom). This result is consistent with a model in which CS treatment enables full-length PKC\(\eta\) to bind to RalA, possibly by changing its conformation and targeting PKC\(\eta\) to the plasma membrane where RalA is localized, with following redistribution of RalA from the plasma membrane to the punctuate structure. Indeed, binding of endogenous RalA to PKC\(\eta\) was detected only in CS-treated cells (Figure 9B), under conditions in which endogenous RalA moved from the membrane to punctuate intracellular structure (Figure 9C). More importantly, activation of RalA was confirmed in the CS-induced keratinocyte differentiation, and the activation was inhibited by siRNA of PKC\(\eta\) (Figure 8D). These results support that binding of RalA to PKC\(\eta\) and following activation of RalA

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**FIGURE 4:** RalA as binding protein of PKC\(\eta\) and its localization in the skin. (A) Detection of binding protein specific for RD-PKC\(\eta\) via its C1 domain. Silver staining of proteins in the HaCaT lysate eluted from the GST-RD or GST-\(\Delta\)C1 column. (B) Presence of RalA in the eluate from GST-RD column. The same sample described in (A) was blotted onto PVDF membrane followed by immunoblotting using anti-RalA. (C) Detection of RalA mRNA in the keratinocytes by RT-PCR. mRNA was extracted from normally cultured NHEK cells. mRNA from mouse muscle was used for positive control. (D) Detection of endogenous RalA by mouse anti–RalA antibody in the cultured HNEK. For negative control, mouse anti–FLAG antibody was used. (E) Fluorescent immunostaining of endogenous RalA in the human epidermis. Bars are 20 \(\mu\)m.

**FIGURE 5:** Importance of the C1 domain for the RalA binding. (A) Direct binding of FLAG-RalA to GST-RD, but not to GST-\(\Delta\)C1. Pull-down assay using FLAG-RalA and GST-RD or GST-\(\Delta\)C1 was performed as described in Materials and Methods. (B) Colocalization of FLAG-RalA with RD-GFP. NHEK cells expressing FLAG-RalA were subsequently infected with viruses coding RD-GFP (RD), \(\Delta\)C1-GFP (\(\Delta\)C1), or GFP alone. The cells were fixed 24 h after the infection, followed by immunocytochemistry using anti–FLAG antibody and Alexa 594–conjugated secondary antibody. Both red and green fluorescences were observed under confocal microscopy.


**DISCUSSION**

As previously reported (Ohba et al., 1998), overexpression of PKCη in human keratinocytes induced the enlarged and flattened morphology associated with differentiation. Interestingly, catalytic activity of PKCη was not necessary, and RD is sufficient for the induction of morphological change. Instead, binding of PKCη-RD to RalA was involved in the process. In other words, RD of PKCη not only prevents the kinase activity in the resting state but also functions as a scaffold region in the active state. A similar additional function of RD of PKC was reported previously; RD of PKCε induces neurite outgrowth (Zeidman et al., 1999) and involvements of binding protein and/or lipids in this process are suggested (Zeidman et al., 2002; Ling et al., 2005; Shirai et al., 2007). Thus it may be common that regulatory domain of PKC has important roles in addition to "rid" of kinase domain. On the other hand, the kinase activity was necessary for the gene expression of the marker proteins, such as involucrin and TGase, necessary for differentiation. These results are very consistent with previous reports (Ueda et al., 1996; Ohba et al., 1998).

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**PKCη and RalA are colocalized with actin depolymerized during the differentiation of keratinocytes**

The morphological changes associated with keratinocyte differentiation require actin rearrangement. Thus the localization of actin and PKCη or RalA in response to CS or expression of RD was investigated. The cortical actin cytoskeleton was apparent in the untreated NHEK cells. In response to CS, F-actin reorganized from cortical bands to more fibrous and punctate structures as the cell spread (Figure 10A); PKCη colocalized with actin in these areas (Figure 10B, third row). In the absence of CS, neither unconjugated GFP nor PKCη-GFP colocalized with actin (Figure 10B, first and second rows). Similar F-actin reorganization and its colocalization with RD were confirmed in the case of RD expression; fibular actin was evident in the flattened cells by RD overexpression, whereas cortical actin was still observed in the small cells expressing no RD (Figure 10B, bottom row). These results suggest that actin depolymerization occurs during keratinocyte morphological changes induced by CS treatment or RD expression, and PKCη catalytic activity was not necessary for association with reorganized actin. Consistent with a requirement for actin depolymerization in the spreading, stabilization of actin with jasplakinolide inhibited the RD-induced morphological changes in a dose-dependent manner (Figure 10C).

Interestingly, in contrast to PKCη, RalA colocalized with actin in both resting and CS/Ca²⁺-treated cells; endogenous RalA was localized on the plasma membrane before the stimulation (Figure 10D). The presence of actin in RalA immunoprecipitates from control and CS/Ca²⁺-treated NHEK cells (Figure 10E) confirmed the association of endogenous RalA and actin in both the resting and differentiated cells, with more recovered following CS/Ca²⁺ treatments (Figure 10E). Notably, RalA immunoprecipitation from CS/Ca²⁺-treated cells also brought down PKCη (Figure 9B). These results are consistent with a model in which CS/Ca²⁺ increases RalA association with actin and PKCη, resulting in actin depolymerization with redistribution of the actin/RalA/PKCη complex, leading to cell spreading.

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**FIGURE 6:** Characteristics of the binding between RalA and RD-PKCη. (A) Weaker binding of RD-PKCη to RalB than RalA. HA-tagged RalA or RalB was expressed in Cos-7 cells. The lysates were incubated with purified GST-RD or GST and pulled down by glutathione–sepharose 4B. The precipitates were applied to 10% SDS–PAGE followed by immunoblotting using HA or GST antibody. (B) No binding of RD-PKCη to Rho family GTPases. Pull-down assay using HA-tagged cdc42, RhoA, or RalA was performed as described in (A). (C) Effect of GDP-β-S or GTP-γ-S on the binding of GST-RD to FLAG-RalA. The binding assay using purified proteins was performed in the presence of 1 mM GDP-β-S or GTP-γ-S. (D) Binding of GST-RD to active form (G28V) and inactive form (G23V) of Ral A. *+, degraded products of GST-RD and GST-ΔC1.
RalA is thought to be multifunctional in mammalian cells, such as regulating membrane transport, apoptosis, transcription cell migration, cell proliferation, and oncogenesis (Feig, 2003; Cammonis and White, 2005; van Dam and Robinson, 2006). However, before this study, there was no report to show its involvement in the morphological change during keratinocyte differentiation. Our novel findings provide a new idea, as shown in Figure 11, in the scenario that PKCη plays a crucial role in normal keratinocyte differentiation and CS is one of the triggers of the activation (Kashiwagi et al., 2002). In undifferentiated keratinocytes, PKCη is localized in the cytoplasm, whereas RalA and actin colocalized at the plasma membrane. Production of CS during the differentiation activates PKCη, resulting in up-regulation of marker proteins for differentiation. Coincidently, CS activation induces the membrane translocation of PKCη and enables its association with RalA. Alternatively, membrane localization of PKCη might require binding to RalA. In either case, the binding of PKCη to RalA appears to be a trigger of activation of RalA and actin depolymerization, leading to the morphological change. Finally, PKCη, RalA, and depolymerized actin are associated and colocalized in the cytoplasm of enlarged and flattened cells. In short, PKCη may function as scaffold proteins of RalA and actin, leading to actin depolymerization. The hypothesis is supported by the following findings. First, actin depolymerization occurs after PKCη translocation; actin depolymerization was evident at 36 and 48 h, whereas PKCη translocation was observed at 12 and 24 h (Figures 1 and 10). Simultaneous observation of PKCη and actin in the living keratinocytes proved the order more directly (data not shown).
Second, overexpression of RD, which does not need any stimulator or time to induce both membrane translocation and the RalA binding, enhanced the rate of the morphological change and the actin depolymerization (Figures 2, 5, and 10B). Third, only mutants that retained both plasma membrane localization and RalA binding induced the morphological change and RalA activation (Figures 3, 5, and 8C). In addition, full-length PKCα cannot bind and colocalize with RalA without CS stimulation, but RD can (Figures 5B and 9). Furthermore, jasplakinolide inhibited the RD-induced morphological change (Figure 10C), whereas latrunculin A enhanced the morphological change (data not shown).

In addition to PKCα, many binding proteins have been identified, and the functions of RalA tightly depend on the protein–protein interactions. One of the first identified RalA binding partners was Ral binding protein 1, RalBP1 (Cantor et al., 1995), and the other binding proteins include PLD1, Sec5, ZONAB, and PLCδ (van Dam and Robinson, 2006). Some of the interactions with binding proteins depend on RalA being activated, whereas others are constitutive. For example, RalBP1 binds to the active form of RalA, whereas PLD1 interacts with both forms. The former type of binding protein tends to bind to the switch region of RalA (van Dam and Robinson, 2006). The binding of RalA to PKCα was independent of the nucleotide exchange (Figure 6), suggesting that PKCα does not bind to the switch region. Although PKCα can bind to both forms, the active form of RalA seems to function in the morphological change because GDP-β-S and coexpression of inactive mutant (S28N) inhibited the RD-induced morphological change (Figure 8).

How RalA is activated during the differentiation is still unknown. Ral proteins can be activated by specific guanine nucleotide exchange factors (GEFs), including RalGDS (Feig, 2003; Cammonis and White, 2005; van Dam and Robinson, 2006). However, we failed to detect RalGDS associated with RalA or PKCα after the CS and calcium stimulation (data not shown), although PKCα is necessary for RalA activation (Figure 8D). RD-PKCα showed no GEF activity (data not shown). It has been recently reported that overexpression of novel class of GEF, RalGPS, causes morphological changes in HEK cells (Rebhum, 2000; Ceriani et al., 2007). Thus RalGPS is a potential candidate for activation of RalA during keratinocyte differentiation. Alternatively, additional production of RalA may be one of the activation mechanisms because the RD overexpression or CS treatment tended to increase mRNA of RalA (data not shown).

In addition, how the binding of RalA to PKCα induces morphological changes is still unclear. Overexpression of wild-type or active RalA alone did not induce the morphological change (data not shown), suggesting a requirement for other factors, possibly binding partner(s). One of the candidates is RalBP1, which interacts with Cdc42 and Rac1 (Cantor et al., 1995; Feig et al., 1996). Indeed, overexpression of inactive Rac1 inhibited the RD-induced morphological change (data not shown). However, we cannot exclude the possibility that inactive Rac1 prevented the adhesion of cells itself because Rac1 has been reported to be necessary for keratinocyte adhesion (Benitah et al., 2005; Tscharntke et al., 2007). The involvement of Rac1 in the morphological changes associated with keratinocyte differentiation is unlikely as it is highly expressed in the basal cells but not in their differentiated counterparts (Benitah et al., 2005), and reduction of Rac1 induces the keratinocyte differentiation (Nikolove et al., 2008). Rather, actin binding proteins are more likely candidates based on the fact that RalA targets filamin to induce filopodia (Ohta et al., 1999; Sugihira et al., 2001). An alternative possibility is the involvement of GTPase-activating proteins (GAP) because immunoprecipitates with RD-PKCα from differentiated keratinocytes, but not RD-PKCα itself, showed slight GAP activity (data not shown). These results suggest that both activation and inactivation of RalA may be involved in the morphological change, contributing to membrane trafficking to provide the additional plasma membrane needed for cell spreading. In fact, RalA is present on secretory vesicles in many cell types (Feig, 2003; van Dam and Robinson, 2006), and PLD1, one of the RalA binding proteins, is involved in vesicle budding and transport (Jenkins and Frohman, 2005). More interestingly, PLD is also reported to regulate the differentiation of keratinocytes (Jhung et al., 1998, 1999). The speculation is very attractive, but further experiments are necessary.
FIGURE 9: CS-dependent interaction between RalA and full-length PKCζ. (A) CS-dependent colocalization of full-length PKCζ with FLAG-RalA. NHEK cells expressing FLAG-RalA (RaLa) and PKCζ-GFP (PKCζ) were cultured for 24 h with or without 50 μM CS. After fixation, FLAG-RalA and PKCζ-GFP were detected as described in (A). (B) Binding of endogenous RalA to endogenous PKCζ. Immunoprecipitation using RalA antibody was performed as described in Materials and Methods. (C) CS-induced relocation of endogenous RalA. After 24 h incubation with or without 50 μM CS, NHEK cells were fixed followed by immnostaining with RalA antibody. Bars are 20 μm.

In addition to filopodia formation and vesicle transportation described above, a link between RaLa and regulation of cell morphology has been reported. Specifically, RaLa promotes neurite branching (Lalli and Hall, 2005), and its activation has been implicated in cytoskeletal changes associated with cell stimulation by chemoattractant peptides (Bhattacharya et al., 2002). However, compared with Rho family small GTPases, the molecular mechanism by which RaLa modulates the actin cytoskeleton is still enigmatic, and the biological function of RaLa has not been fully elucidated. Our findings provide one possible model for how RaLa and PKCζ regulate the actin cytoskeleton and cellular morphology during keratinocyte differentiation.

MATERIALS AND METHODS

Materials

CS, GDP-β-S, GDP-γ-S were purchased from Sigma (St. Louis, MO). HuMedia-KG2 medium and DMEM were obtained from Kurabo Industries (Osaka, Japan) and Wako (Osaka, Japan). Both penicillin and streptomycin were products of Life Technologies (Carlsbad, CA), and fetal bovine serum (FBS) was from ICN Biomedicals (Irvine, CA). Anti-FLAG and RaLa monoclonal antibodies were purchased from Sigma and BD Transduction Laboratories (Lexington, KY), respectively. Anti-GST was from Santa Cruz Biotechnology (Santa Cruz, CA). cDNA encoding FLAG or GST-fused RaLa and hemagglutinin (HA)-tagged RaB were provided by T. Kataoka (Kobe University) and T. Sato (Kobe University), respectively. The plasmids encoding HA-tagged Rho family GTPases, Rac1, RhoA, and cdc42, were given by K. Kaibuchi (Nagoya University).

Construction of adenovirus vector

The cDNA encoding PKCζ and mutants were produced by a PCR with cDNA for mouse PKCζ as the template. The sense and anti-sense primers used were as follows: for full-length PKCζ, 5′-TTAGATCTCGGATCCATGCCTCGGCGAAGGACATG3′ and 5′-GGGAATTCATGACCTGAGGAGGATTACGAG3′; for RD, 5′-TTAGATCTCGGATCCATGCCTCGGCGAAGGACATG3′ and 5′-GGGAATTCATGACCTGAGGAGGATTACGAG3′; for CS-Dependent interaction between RalA and full-length PKCζ. The cDNA encoding PKCζ and mutants were produced by a PCR with cDNA for mouse PKCζ as the template. The sense and anti-sense primers used were as follows: for full-length PKCζ, 5′-TTAGATCTCGGATCCATGCCTCGGCGAAGGACATG3′ and 5′-GGGAATTCATGACCTGAGGAGGATTACGAG3′; for RD, 5′-TTAGATCTCGGATCCATGCCTCGGCGAAGGACATG3′ and 5′-GGGAATTCATGACCTGAGGAGGATTACGAG3′. Further, the respective cDNA fragments were amplified by ExSite PCR-based site-directed mutagenesis kit (Stratagene, La Jolla, CA). In the case of ΔV1V3, ΔPS, and ΔC1, site-directed mutagenesis was performed according to manufacturer’s recommended protocol with ExSite PCR-based site-directed mutagenesis kit (Stratagene, La Jolla, CA). RD in pUC118 was used as a template, and the primers used were as follows: for ΔV1V3, 5′-GGGAATTCATGACCTGAGGAGGATTACGAG3′ and 5′-ATTAGCTATTGGCAATGGACCGCTGACGAC-3′; for V1 and P1 deletion (ΔVP1), 5′-GGGAATTCATGACCTGAGGAGGATTACGAG3′ and 5′-ATTAGCTATTGGCAATGGACCGCTGACGAC-3′; for V1 and V3 deletion (ΔV1V3), 5′-GGGAATTCATGACCTGAGGAGGATTACGAG3′ and 5′-ATTAGCTATTGGCAATGGACCGCTGACGAC-3′. Further, the respective cDNA products were first subcloned into pUC118 Invitrogen (San Diego, CA).

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After being digested with BglII, the respective cDNA fragments and a cDNA encoding GFP, which was obtained from BS340 (Shirai et al., 1998) with HindIII digestion, were subcloned into the BglII and HindIII sites of pShuttle–CMV vector. After linearization by Pmel, the pShuttle vector having cDNA encoding fusion protein of GFP with PKCζ or its mutant was coelectroporated with pAdEasy backbone vector into BJ5183 bacterial cells. The recombination was checked by Pad, cut, and the plasmids were purified by CsCl banding. After Pad digestion, 10 μg of the purified plasmids was lyophoected into 50–70% confluent HEK293 cells plated on a 6-cm dish by Fugene II, cut, and the plasmids were purified by CsCl banding and titrated.

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using a soft wear of LSM501 for each experiment; three independent experiments were performed. Finally, the data were plotted as a percentage of the infected cells.

Construction of FLAG-tagged RalA mutants and HA-tagged RalA
The cDNA encoding FLAG-tagged RalA mutants were produced by site-directed mutagenesis according to manufacturer’s recommended protocol with QuickChange II XL site-directed mutagenesis kit (Stratagene). A cDNA for FLAG-tagged human RalA was used as the template. The sense and anti-sense primers used were as follows: for inactive form, S28N RalA, 5′-GGCAAGTGGTG-3′ and 5′-CATGAACTGTAGAGTCAACGCGTTCTTGCCCACGCCACCACTGCC-3′; for active form, G23V RalA, 5′-GGGTGCTGGGCAAGAGTGCAATGACTCTACAGTTCATG-3′ and 5′-GAACTGTAGAGTCAGTGCACTCTTGCCCACGCCAACACTGCCCACCACATGATG-3′.

To obtain a plasmid encoding HA-tagged RalA, a cDNA for human RalA obtained by BamHI cut from the plasmid encoding FLAG-tagged RalA was cloned into pEF-BOS-HA vector donated by K. Kaibuchi.

Cell culture
NHEK cells were kindly provided by Masahiro Oka (Kobe University). Keratinocytes were cultured in HuMedia-KG2 medium supplemented with insulin (10 μg/ml), epidermal growth factor (0.1 mg/ml), hydrocortizone (0.5 μg/ml), and gentamicin (50 μg/ml). HaCaT cells were cultured in DMEM supplemented with penicillin (100 U/ml), streptomycin (100 μg/ml), and 10% FBS. COS7 cells were purchased from Riken Cell Bank (Tsukuba, Japan) and cultured in DMEM supplemented with penicillin (100 U/ml), streptomycin (100 μg/ml), and 10% FBS. All cells were kept at 37°C in a humidified atmosphere containing 5% CO₂.

Adenoviral infection and measurement of cell size
A total of 1 × 10⁶ of the cells were seeded in a 3.5-cm, glass-bottom dish with 2 ml of the medium a day before the infection. High-titer virus solution (>MOI = 5) was added to the dish, and medium was changed to normal medium after 1 h. After culturing with or without reagents including CS, the cells were fixed at appropriate time points by 4% paraformaldehyde and 0.2% picric acid in 0.1 M phosphate buffer overnight at 4°C. The fixed cells were observed under confocal microscopy. The longest axis of 100 cells was measured using a soft wear of LSM501 for each experiment; three independent experiments were performed. Finally, the data were plotted as a percentage of the infected cells.

Construction of FLAG-tagged RalA mutants and HA-tagged RalA
The cDNA encoding FLAG-tagged RalA mutants were produced by site-directed mutagenesis according to manufacturer’s recommended protocol with QuickChange II XL site-directed mutagenesis kit (Stratagene). A cDNA for FLAG-tagged human RalA was used as the template. The sense and anti-sense primers used were as follows: for inactive form, S28N RalA, 5′-GGCAAGTGGTG-3′ and 5′-CATGAACTGTAGAGTCAACGCGTTCTTGCCCACGCCACCACTGCC-3′; for active form, G23V RalA, 5′-GGGTGCTGGGCAAGAGTGCAATGACTCTACAGTTCATG-3′ and 5′-GAACTGTAGAGTCAGTGCACTCTTGCCCACGCCAACACTGCCCACCACATGATG-3′.

To obtain a plasmid encoding HA-tagged RalA, a cDNA for human RalA obtained by BamHI cut from the plasmid encoding FLAG-tagged RalA was cloned into pEF-BOS-HA vector donated by K. Kaibuchi.
FIGURE 11: Hypothetical roles of binding of PKCζ and RalA during keratinocyte differentiation. (A) In basal cells, PKCζ is localized in the cytoplasm, whereas RalA and actin present at the membrane. N, nucleus. (B) Production of CS during the differentiation activates PKCζ, resulting in translocation to the membrane. The activated PKCζ up-regulates marker proteins for differentiation (dotted lines) and binds to RalA. (C) Binding of PKCζ to RalA somehow induces actin depolymerization, resulting in dotty colocalization of PKCζ, RalA, and actin in the cytoplasm of enlarged cells. (D) Finally, the enlarged cells differentiated to cornified cells.

Immunocytochemistry
To compare localization of PKCζ and its mutants with FLAG-RalA, NHEK cells were lipofected with FLAG-RalA by Fugene (Roche) as described previously (Shirai et al., 2007). Simultaneously, the cells were infected with adenoviruses and fixed as described. The fixed cells were permeabilized with 0.3% Triton X-100/PBS (PBS-T) for 30 min at room temperature and blocked for 1 h with 10% bovine serum albumin (BSA) in PBS. One-hour incubations with anti-FLAG or RalA monoclonal antibody (mAb) were performed at a 1:800 or 1:1000 dilution with 0.03% PBS-T, respectively. Alexa Fluor 594-conjugated anti–mouse immunoglobulin (IgG) (Molecular Probes, Eugene, OR) was used as a secondary antibody.

To visualize F-actin, the fixed cells were sequentially treated with 0.3% PBS-T, 10% NGS, and diluted rhodamin–phalloidin (Molecular Probes) with PBS-T (1:250) for 1 h.

Human skin voluntarily provided was fixed with 4% paraformaldehyde in 0.1 M phosphate buffer. After dehydration, the fixed skin was blocked in paraffin wax. The blocked skin sample was sliced and transferred to slides coated with poly-L-lysine. The wax was removed with Xylene, and the tissue sections were rehydrated through a graded series of ethanol before being placed in 10 mM PBS. The sections were preincubated with 0.3% H2O2 for 20 min to inactivate endogenous peroxidase activity, with 10% NGS for 1 h to block nonspecific binding sites, and then with 0.1% phenylhydrazine for 20 min to inactivate residual endogenous peroxidase activity. The sections were incubated with RalA antibody (1:2000) in PBS-T overnight at 4°C. After washing with PBS-T, the sections were incubated with Alexa 488–conjugated anti–mouse IgG secondary antibody (1:2000) for an additional 24 h. Finally, the fluorescence was observed under confocal microscopy after washing well.

Confocal microscopic analysis
The fluorescence of the GFP or Alexa 488 was monitored with a confocal laser scanning fluorescence microscope (LSM 510 invert; Carl Zeiss, Jena, Germany) at 488-nm argon excitation using a 515- to 535-nm band-pass barrier filter, whereas red fluorescence was monitored at 543-nm HeNe excitation using a 560-nm long-pass barrier filter.

Recombinant protein expression and purification
cDNA encoding RD or ΔC1 was digested with BglII from the pUC118-based plasmids described previously and subcloned into the BglII site of pGEX 6P-1 (GE Healthcare Biosciences, Uppsala, Sweden). For bacterial expression of GST-fusion proteins, BL21 (DE3) pLys cells were transformed with the expression plasmids. Expression of the recombinant proteins was induced by 0.1 mM isopropyl-1-thio-β-D-galactoside at 25°C for 16 h. The cells were harvested and lysed in the lysis buffer containing 20 mM Tris-HCl (pH 8.0), 1 mM EDTA, 1 mM dithiothreitol (DTT), 5 mM MgCl2, 250 mM sucrose, 1% Triton X-100, 20 μg/ml leupeptin, 1 mM phenylmethylsulfonyl fluoride (PMSF) with handy sonic (Tomy Seiko, Tokyo, Japan). After ultracentrifugation at 10,000 × g for 30 min, fusion proteins were purified using Glutathione-Sepharose 4B (GE Healthcare Biosciences) as previously described (Yamaguchi et al., 2006).

Detection of protein(s) binding to C1 domain of PKCζ
HaCaT cells harvested from 20 plates of 10-cm dishes were homogenized in 5 ml of the lysis buffer described above and stored on ice for 30 min. After ultracentrifugation at 10,000 × g for 30 min, the supernatant was applied onto the glutathione–sepharose 4B column immobilizing purified GST-RD or GST-ΔC1. After washing with TEDM buffer (20 mM Tris-HCl [pH 8.0], 1 mM EDTA, 1 mM dithiothreitol (DTT), 5 mM MgCl2, 250 mM sucrose), proteins bound to GST-RD or GST-ΔC1 were eluted with 10 mM reduced glutathione and 0.1% SDS in 50 mM Tris-HCl buffer (pH 8.0). The eluted proteins were applied to SDS–PAGE followed by silver staining.

Mass spectrometric analysis
Proteins in SDS polyacrylamide gels were visualized by the reverse-staining method. The bands corresponding to proteins were excised, and then proteins in gels were reduced by incubating with 10 mM EDTA/10 mM DTT/100 mM ammonium bicarbonate for 1 h at 50°C and alkylated by treatment with 10 mM EDTA/40 mM iodoacetamide/100 mM ammonium bicarbonate for 30 min at room temperature. They were digested in gel with lysyl endopeptidase (LEP) from Achromobacter lyticus (Wako Pure Chemical Industries) in 100 mM Tris/HCl (pH 8.9) for 15 h at 37°C. Resulting peptide fragments were extracted from gels and then concentrated in vacuo. After desalting with ZipTipC18 (Millipore, Billerica, MA), peptide fragments were subjected to mass spectrometric analysis. Positive ion mass spectra were acquired in a Micromass
Q-ToF2 mass spectrometer equipped with a nano electrospray ionization source. Tandem mass spectrometry (MS/MS) was performed by collision-induced dissociation using argon as the collision gas.

**Pull-down assay**

cDNA encoding FLAG-RalA or its mutant was transfected into COS7 cells by electroporation (Yamaguchi et al., 2006). After 48 h, cells were harvested and lysed in 200 μl of Tris-buffered saline (TBS)-T buffer (50 mM Tris-HCl [pH7.4], 150 mM NaCl, 1% Triton X-100, 1 mM PMSF, 20 μg/ml leupeptin). Total lysate of the COS7 cells expressing FLAG-RalA or its mutants was mixed with 2 μg of purified GST, GST-RD, or GST-AC1 in 1 ml of PBS(-) containing 1% bovine serum albumin and 50 μl of glutathione–sepharose 4B for 2 h at 4°C. To determine the effect of GTP or GDP on the binding, GDP-β-S or GTP-γ-S was added into the mixture. After centrifugation and washing with PBS(-) three times, the bound proteins were eluted with 50 μl of SDS buffer for 10% SDS–PAGE followed by immunoblotting using anti–GST, FLAG, or RalA antibody.

**Immunoprecipitation**

To detect interaction between endogenous RalA and PKCη, immunoprecipitation was performed. NHEK cells treated with or without calcium and CS were homogenized in the lysis buffer with sonication on ice and then centrifuged at 10,000 × g for 5 min. Cleared lysates were incubated with RalA mAb for 2 h at 4°C, and then 50 μl of protein G sepharose (GE Healthcare) were added to the mixture. After centrifugation and washing with PBS(-) three times, the bound proteins were eluted with 50 μl of SDS buffer for 10% SDS–PAGE followed by immunoblotting using anti-PKCη (1:1000 dilution) and RalA antibody (1:10,000 dilution).

**Qantitative PCR**

Total cellular RNA was extracted from NHEK using SV Total RNA Isolation System (Promega, Madison, WI) according to manufacturer’s recommended protocol and quantified spectrophotometrically. A total of 200 ng of total RNA was reverse-transcribed into cDNA using ThermoScript RT-PCR System (Invitrogen) following standard protocol and was then applied to real-time PCR using CYBER green and an ABI Prism 7000 (Applied Biosystems, West Grove, PA) for 1 h at room temperature. After extensively washing with PBS-T, the membrane was immunostained with appropriate primary antibodies for 1 h at room temperature. After three rinses with PBS-T, the membrane was incubated with peroxidase-labeled anti-rabbit IgG or anti–mouse IgG (Jackson Immunoresearch, West Grove, PA) for 1 h at room temperature. After extensively washing with PBS-T, the immunoreactive bands were visualized using an enhanced chemiluminescence detection kit (GE Healthcare Bioscience).

**siRNA experiments**

Target sequence for Ral A, CGCGGTCAGATCTTCTTA (HsRALA), and control sequence, CGGAGCGGAUAUCGUAATTT, were obtained from QiAGEN (Valencia, CA) and Invitrogen, respectively. siRNA for human PKCη (sc-44020) was obtained from Santa Cruz Biotechnology. Cells were transfected with dsRNA using lipofectamine 2000 (Invitrogen) and cultured for 48 h. After 24 h transfection of siRNA, the cells were infected with adenovirus for RD or GFP and cultured for an additional 24 h with or without CS and calcium.

**RalA activation assay**

The assay was essentially performed as described previously (Kawato et al., 2008). Briefly, treated or nontreated HaCaT cells were lysed in lysis buffer as described. After removal of insoluble materials, the lysate was incubated with glutathione–sepharose beads coated with 10 μg GST-Sec5-RBD at 4°C for 1 h. After washing, active RalA, which was pulled down with beads, was analyzed by immunoblotting using the anti–RalA mAb.

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**Tandem mass spectrometry**

Q-ToF2 mass spectrometer equipped with a nano electrospray ionization source. Tandem mass spectrometry (MS/MS) was performed by collision-induced dissociation using argon as the collision gas.
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