Activation of ROCK by RhoA is regulated by cell adhesion, shape, and cytoskeletal tension

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ARTICLE INFORMATION

Article Chronology:
- Received 15 March 2007
- Revised version received 1 July 2007
- Accepted 2 July 2007
- Available online 10 July 2007

Keywords:
- Rho
- ROCK
- Kinase assay
- Myosin phosphorylation
- Cell shape
- Focal adhesions
- FAK
- Hypertension

ABSTRACT

Adhesion to the extracellular matrix regulates numerous changes in the actin cytoskeleton by regulating the activity of the Rho family of small GTPases. Here, we report that adhesion and the associated changes in cell shape and cytoskeletal tension are all required for GTP-bound RhoA to activate its downstream effector, ROCK. Using an in vitro kinase assay for endogenous ROCK, we found that cells in suspension, attached on substrates coated with low density fibronectin, or on spreading-restrictive micropatterned islands all exhibited low ROCK activity and correspondingly low myosin light chain phosphorylation, in the face of high levels of GTP-bound RhoA. In contrast, allowing cells to spread against substrates rescued ROCK and myosin activity. Interestingly, inhibition of tension with cytochalasin D or blebbistatin also inhibited ROCK activity within 20 min. The abrogation of ROCK activity by cell detachment or inhibition of tension could not be rescued by constitutively active RhoA-V14. These results suggest the existence of a feedback loop between cytoskeletal tension, adhesion maturation, and ROCK signaling that likely contributes to numerous mechanochemical processes.

Introduction

Contractile tension generated within the actin cytoskeleton by myosin II is emerging as a key player in many cellular processes, from the stabilization of cell-matrix adhesions to the modulation of gene expression, cell proliferation, and differentiation [1–5]. A principal mediator of cytoskeletal tension is the small GTPase RhoA and its downstream effector Rho-associated kinase (ROCK). GTP binding and hydrolysis switches RhoA between a GTP-bound active and a GDP-bound inactive state [6]. The conformationally active RhoA propagates downstream signals in turn by binding to effector proteins such as ROCK. ROCK, through the phosphorylation and deactivation of the myosin binding subunit of myosin phosphatase (MYPT1), and direct phosphorylation of myosin light chain (MLC), leads to contractile force generation [7].

Integrin-mediated adhesion to the extracellular matrix (ECM) has been shown to regulate RhoA signaling [8,9]. It has been shown that placing cells in suspension leads to increased GTP-RhoA levels, [10] and replating cells on ECM leads to a transient down regulation of RhoA activity followed by a slow recovery [11]. While it has largely been thought that adhesion regulates the RhoA–ROCK-tension pathway through its effects on RhoA [8,9], evidence suggests the possibility that the coupling of RhoA to ROCK activity also may be an important control point. For example, despite high levels of GTP-RhoA...
[11], suspended cells exhibit low MLC phosphorylation [2,10]. Similarly, while cytochalasin D treatment has long been known to decrease MLC phosphorylation [12], it has also been shown to increase cellular GTP-RhoA [11]. Thus while high RhoA activity is often used to indirectly implicate high contractility, RhoA activity may in certain circumstances actually be decoupled from myosin-based contractility. Decoupling between RhoA and another effector, mDia, has recently been reported [13]. Elimination of signaling by FAK, a principal mediator of integrin signaling, abrogates mDia-mediated stabilization of microtubules at the leading edge despite the presence of active RhoA [13]. Furthermore, the related Rho family GTPase, Rac, requires adhesion to couple to its effector PAK. By recruiting cholesterol rich lipid rafts to the plasma membrane, integrin ligation localizes active Rac to the membrane, permitting Rac to activate PAK [14]. Together, these findings suggest that RhoA-mediated ROCK activity may also be regulated by adhesion.

Interestingly, RhoA signaling not only is modulated by cell-ECM adhesion, but also acts as an important regulator of adhesion. Adhesion to the ECM involves numerous interrelated processes including integrin binding, cell spreading and flattening against the substrate, and condensation of integrins to form large structures known as focal adhesions. RhoA–ROCK-mediated cytoskeletal tension appears to limit the degree to which cells spread and is critical for focal adhesion maturation and signaling [15–17], and ROCK-generated tension may feedback to regulate RhoA at least in cells on collagen gels [3]. These data highlight the possible existence of feedback and feed forward regulatory loops that, if present, would play a critical role in how cells adaptively alter adhesion, morphology, and mechanics in response to their ECM environment.

In this study, we investigated the possibility that both adhesion and tension are involved in regulating RhoA, ROCK and myosin activity. We examined how different aspects of adhesion (integrin ligation, cell spreading, focal adhesion formation, and cytoskeletal tension) modulated RhoA-ROCK signaling. Our findings demonstrate that cell adhesion and spreading are required for RhoA to activate ROCK, and that this direct modulation of ROCK activity by adhesion requires traction forces mediated by cytoskeletal tension. Thus, full myosin activation by RhoA requires adhesion to establish a positive feedback loop that links RhoA-myosin signaling and force-mediated adhesion maturation. This control system provides an explanation for how adhesion, cell mechanics, and RhoA GTPase signaling are so closely intertwined in many mechanotransduction processes.

Materials and methods

Cell culture and reagents

Primary bovine pulmonary artery endothelial cells (VEC Technologies, Rensselaer, NY) were grown in DMEM supplemented with 5% bovine serum, 100 U/ml penicillin, and 100 μg/ml streptomycin (all from Invitrogen). Cells were used between passages 5 and 8. Human umbilical vein endothelial cells (gift from Dr. Guillermo Garcia-Cardena, Harvard Medical School) were grown in M199 medium (Cambrex) supplemented with 20% FBS (Invitrogen), 50 μg/ml endothelial mitogen (Biomedical Technologies), 100 μg/ml heparin (Sigma), penicillin, and streptomycin. Cells were used between passages 5 and 7. Other reagents used were: fibronectin (BD Biosciences); BSA (Serologicals Corporation); anti-phospho-myosin light chain 2 (Thr18/Ser19) (Cell Signaling Technologies); antimonocline myosin II heavy chain (Biomedical Technologies); recombinant MYPT1 fragment, anti-ROCK-2, anti-phospho MYPT1 Thr 696, Rhotekin agarose beads (all from Upstate); anti-RhoA (Santa Cruz); anti-vinculin, protease, and phosphatase inhibitor cocktails (all from Sigma); protein-G sepharose beads (Amersham); ATP (Sigma); Blebbistatin (25 μM; Tocris); cytochalasin D (2 μg/ml; Sigma); TRITC-phalloidin (Molecular Probes); NaCl, EDTA, MgCl2, glycerol (all from JT Baker); HEPES (Gibco); NP-40 (Igepal CA-630); leupeptin, pepstatin, aprotinin (all from Sigma). The RhoA-V14 adenovirus was prepared and used as previously described [4].

Manipulation of cell adhesion

For experiments using cells in suspension, cells were plated on Petri dishes coated with 25 μg/ml of fibronectin (FN) and allowed to spread for 3 h at which time they were trypsinized and maintained in suspension with periodic gentle shaking in polystyrene tubes in the incubator for 1 h. ECM density was varied for some experiments by varying the concentration of the FN coating solution, as indicated in the text. Micropatterned substrates were generated on PDMS (polydimethylsiloxane)-coated Petri dishes by using a protein stamping technique described before [18].

Measurement of cytoskeletal tension

Traction forces were measured by using microfabricated substrates previously described [19]. Briefly, cells were plated on microfabricated elastomeric posts on which they attach and spread, applying traction forces to the posts. The resulting bending of the posts was measured and converted to a traction force. Fifteen cells were quantified for each condition, averaged over 3 different samples.

Immunofluorescence microscopy

Cells were washed twice with PBS and fixed with 3.7% paraformaldehyde for 30 min at room temperature. F-actin was visualized by staining with TRITC-phalloidin and epifluorescence images were captured on a Nikon TE2000 microscope using an Orca digital camera (Hamamatsu). Focal adhesions were quantified by vinculin staining and digital image analysis as previously described, including only adhesions larger than 0.25 μm2 [17].

ROCK activity assay

Endogenous ROCK kinase activity in immunoprecipitated ROCK was measured by an in vitro kinase assay using a recombinant fragment (aa 654–880) of human MYPT1 (rMYPT1) as substrate. Cells were washed twice with ice cold PBS and lysed into IP buffer: 50 mM HEPES pH 7.4, 150 mM NaCl, 1 mM MgCl2, 10 mM NaF, 1 mM Na3VO4, 5% glycerol, 1% Nonidet P-40,
1 mM dithiothreitol, 1 mM EGTA, 5 μl/ml each of protease inhibitor and phosphatase inhibitor cocktails. All manipulations after cell lysis were done at ice-cold temperature. Cell lysates were centrifuged at 14,000 rpm for 4 min to remove insoluble debris. Supernatants were transferred to fresh tubes, precleared with protein-G sepharose beads for 15 min, and incubated with anti-ROCK antibody for 40 min. The antibody–ROCK complexes were captured by incubating with protein G sepharose beads for 20 min. Beads were washed 3× in IP buffer and used for the kinase reaction. The lysis buffer quantitatively solubilized cellular ROCK and the immunoprecipitation process depleted greater than 85% of ROCK from the lysates (data...
not shown). We also verified that cellular ROCK levels did not change by the experimental treatments prior to lysis (data not shown). The kinase reaction mixture included, in a final volume of 100 μl, 0.5 μg rMYPT1, 50 μl of beads, and 100 μM ATP. rMYPT1 phosphorylation was measured by Western blotting using rabbit anti-phospho MYPT1 Thr 696. Results were normalized for total protein measured by BCA assay.

**RhoA activity assay**

RhoA activity was measured using a previously described method [11]. Briefly, cells were lysed in Rho buffer (25 mM HEPES, pH 7.5, 150 mM NaCl, 1% NP-40 (Igepal CA-630), 10 mM MgCl₂, 1 mM EDTA and 10% glycerol, 10 μg/ml leupeptin, 10 μg/ml pepstatin, and 10 μg/ml aprotinin) and GTP-Rho was affinity precipitated using Rhotekin-agarose beads. Affinity precipitated RhoA was quantified in parallel with total cellular RhoA from cell lysates by Western blot analysis with an antibody against RhoA.

**Western blot analysis**

Samples were resolved on 4–20% Ready Gels by SDS-PAGE, transferred to PVDF membranes, blocked using 5% nonfat milk powder, and incubated in primary antibody overnight, washed twice with 0.2% Tween in Tris-buffered saline (TBST), incubated with the secondary antibody for 1 h at room temperature, and washed thrice with TBST. Bands were detected using Pierce West Dura chemiluminescence system and quantified using unmodified, raw data obtained from images taken on a Versadoc imager and Quantity One software (Biorad). For measuring MLC phosphorylation, samples were collected by washing cells 2× in ice-cold PBS, lysed into 1× Laemmli sample buffer, boiled, and analyzed by Western blotting. Diphospho-MLC (Ser19/Thr18) signal was normalized to that of nonmuscle myosin II heavy chain.

**Results**

*Adhesion is required for the functional coupling of RhoA activity and ROCK activity*

For the first set of experiments, we examined how complete loss of adhesion to fibronectin (FN) affected myosin phosphorylation, RhoA activity and ROCK activity. Endothelial cells were plated on a high density coating (25 μg/ml) of fibronectin, allowed to attach and spread, and then were either released

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**Fig. 3** - The effect of tension inhibiting drugs on stress fibers, focal adhesions, and contractility. (A) Immunofluorescence micrographs of stress fibers labeled with TRITC-phalloidin in control (left), blebbistatin-treated (25 μM, middle), and cytochalasin D-treated (2 μg/ml, right) cells. (B) Immunofluorescence micrographs of focal adhesions labeled with anti-vinculin antibody in control (left), blebbistatin-treated (middle), and cytochalasin D-treated (right) cells. (C) Graphs of quantified number of focal adhesions per cell in drug-treated cells compared to controls. (D) Differential interference contrast micrographs of control (left), blebbistatin-treated (middle), and cytochalasin D-treated (right) cells on the microfabricated force-measuring device, overlaid with contractile force vectors. The top graph to the right the average number of focal adhesions per cell in control, blebbistatin-treated, and cytochalasin D-treated cells. The bottom graphs show histograms of contractile forces, each histograms representing pooled data from 15 cells under the respective conditions from 3 experiments: red—control cells, green—blebbistatin-treated cells, blue—cytochalasin D-treated cells. Error bars indicate standard deviations of measurement of about 50 cells from a representative experiment.
from the substrate or left attached for one additional hour. As an additional comparison to the suspension condition, some of the suspended cells were incubated with fibronectin-coated polystyrene beads to promote integrin engagement in the absence of cell spreading. Cells were lysed and assayed for myosin light chain (MLC) dual phosphorylation at Ser19/Thr18. MLC phosphorylation was low in suspended compared to attached cells even in the presence of integrin engagement with fibronectin-coated beads (Fig. 1A). Interestingly, loss of adhesion did not change GTP-loading of RhoA in suspended cells as compared to adherent cells, though, integrin engagement through fibronectin-coated beads decreased RhoA activity (Fig. 1B). To determine whether the disconnect between RhoA and myosin signaling occurred upstream or downstream of ROCK, the major regulator of myosin in these cells (Supplementary Fig 1), we examined the activity of endogenous ROCK by kinase assay. Cells in suspension exhibited dramatically lower ROCK activity compared to adherent cells (Fig. 1C), mimicking the loss of myosin activity. These initial findings suggested that, in the absence of adhesion, RhoA fails to activate ROCK, thereby disconnecting RhoA activity from myosin phosphorylation.

**Graded ECM ligand density and cell spreading modulate RhoA–ROCK coupling**

It has previously been shown that Rac–PAK coupling depends only on the presence or absence of adhesion; integrin ligation through fibronectin-coated beads in suspended cells is sufficient for Rac to activate PAK [14]. We examined whether Rho–ROCK signaling was also unaffected by varying the degree of substrate adhesion. Endothelial cells were plated on either high (25 μg/ml) or low (0.1 μg/ml) density of FN coated on polystyrene dishes and allowed to attach and spread (Fig. 2A). Cells on low FN exhibited threefold lower MLC phosphorylation relative to those on high FN (Fig. 2B) while RhoA activity was not different (Fig. 2C). ROCK activity again correlated with myosin phosphorylation, being substantially lower in the cells on low FN (Fig. 2D). Thus, ROCK activity was uncoupled from RhoA even under adherent conditions, albeit of low integrin ligation.

Cell adhesion involves not only changes in integrin ligation and clustering, but also changes in cell spreading and flattening on the substrate, both of which have distinct effects on cell signaling and function. To directly modulate the degree of cell spreading against a substrate without altering ECM density, we used a micropatterning approach to generate patches of ECM the size of individual cells. Cells were plated on 25×25 μm size patterns of FN that restricted them from spreading (‘unspread’) or allowed to freely spread on unpatterned FN (‘spread’) (Fig. 2E) and assayed for RhoA, ROCK and myosin activity. We found that myosin light chain phosphorylation was lower in the spreading-restricted cells on patterns (Fig. 2F) much like the low vs. high density FN condition (Fig. 2B). Interestingly, we found that RhoA activity was higher in the spreading-restricted cells (Fig. 2G). Despite the increase in RhoA signaling, ROCK activity was lower in unspread cells compared to spread cells (Fig. 2H). Together, these data suggest that cell spreading regulates both RhoA activity and the degree of coupling between RhoA and ROCK activity. That is, in unspread cells, RhoA activity is high but it fails to couple and activate ROCK and myosin. Cell spreading moderately decreases RhoA activity, but perhaps by allowing RhoA to act on its effector(s), ROCK and myosin activity dramatically increase.

**Tension is required for the coupling of RhoA and ROCK activity**

Cell shape could mediate RhoA–ROCK coupling through several potential mechanisms. Previous work from our lab and others has shown that unspread cells exert lower cytoskeletal tension than spread cells [19,20], and that this tension in turn alters the number of focal adhesions that form in cells that are spread to different extents [21]. Hence, we examined the possibility that cytoskeletal tension may mediate the coupling of RhoA activity to ROCK. We used two mechanistically different drugs to disrupt cytoskeletal tension in cells—blebbistatin, which is a direct inhibitor of myosin II ATPase activity, and cytochalasin D, which disrupts the actin cytoskeleton. To validate the effects of these inhibitors, endothelial cells were allowed to attach and spread on FN for 4 h and then treated with each inhibitor. Within 20 min, both of the treatments disrupted actin stress fibers, visualized by TRITC-phalloidin staining (Fig. 3A), and decreased the size and number of focal adhesions measured by vinculin staining (Fig. 3B). While these observations suggested that these inhibitors disrupted cytoskeletal tension, they were indirect measures of tension. Using a previously developed microfabricated system, we confirmed directly that cytoskeletal tension was inhibited by drug treatment within 20 min (Fig. 3C). Using these inhibitors, we examined the activity of

![Fig. 4](image-url)
RhoA and ROCK following the 20-min exposure. RhoA activity was unchanged upon blebbistatin treatment and increased upon cytochalasin D treatment (Fig. 4A). In contrast, ROCK activity was inhibited upon the disruption of tension in both cases (Fig. 4B). Thus, disrupting cytoskeletal tension decouples RhoA activity from ROCK.

These findings together demonstrated that both the degree of adhesion and cytoskeletal tension regulated ROCK activity. While our studies demonstrated a loss of correlation between RhoA and ROCK activity when cell spreading or cytoskeletal tension was disrupted, they did not provide a direct demonstration of the functional decoupling between these signals. That is, the effects of these manipulations on RhoA and on ROCK could be unrelated, as ROCK activity can be modulated by several other signals [22–24]. To examine if either adhesion or cytoskeletal tension was required for ROCK activation by RhoA, we infected cells with an adenovirus that expressed the constitutively active RhoA-V14 (Fig. 5 A) and measured ROCK activity under different conditions. As expected, RhoA-V14 expression increased myosin light chain phosphorylation as compared to GFP controls and maintained MLC phosphorylation even upon serum starvation (Fig. 5 B). ROCK activity also increased in RhoA-V14 cells as compared to GFP controls (Fig. 5C). Importantly, RhoA-mediated ROCK activity in RhoAV14 expressing cells was abrogated upon treatment with blebbistatin or cytochalasin D (Fig. 5C). Similarly, suspending RhoA-V14 expressing cells also eliminated RhoA-induced ROCK activity (Fig. 5D). Together, these data indicate that cell adhesion, spreading, and cytoskeletal tension are collectively necessary for RhoA to effectively activate ROCK.

Discussion

Previous studies have suggested that RhoA-ROCK signaling is intricately linked to adhesion signaling. Here, by separating the effects of integrin ligation, cell spreading, and cytoskeletal tension, and directly measuring endogenous RhoA and ROCK activity, we have uncovered several important regulatory mechanisms. First, RhoA activity appears to be regulated by cell shape. Previous studies have shown that integrin-mediated adhesion can antagonize RhoA activity [11], and are consistent with our observation of decreased RhoA activity upon addition of fibronectin-coated beads to suspended cells. This same pathway may be responsible for the further suppression of RhoA activity as cells are allowed to spread on larger micropatterned islands of fibronectin. Although we have not examined the basis for this decrease, it is plausible that the mechanism involves the increased actin polymerization observed with cell adhesion and spreading [25]. It has been suggested that F-actin may sequester a Rho guanine-nucleotide exchange factor (Rho GEF) [26], such that increased actin polymerization would suppress RhoA activity. Consistent with this model, we observed increased RhoA-GTP loading upon cytochalasin D treatment, a phenomenon that has been previously noted by other researchers [11]. Additional studies will be required to further delineate this mechanism.

Second and perhaps most interestingly, the ability of RhoA to modulate ROCK kinase activity is dependent on cell adhesion, cell spreading, and cytoskeletal tension. This decoupling
of RhoA and ROCK is evident in suspended cells, which exhibited high levels of Rho-GTP but low MLC phosphorylation and ROCK activity. Remarkably, allowing cells to adhere on a high density of FN was not sufficient to rescue ROCK activity; instead, cell spreading and cytoskeletal tension were also required (Figs. 2 and 4). Previous studies have shown for other small GTPase effectors PAK and mDia, that integrin ligation is sufficient to mediate their activation by Rac and Rho, respectively [13,14]. This decoupling of Rho and ROCK by changes in cell shape may be one important specific mechanism by which cell shape modulates cell signaling, cytoskeletal tension, and adhesion maturation. The requirement of cytoskeletal tension to couple Rho-GTP to ROCK may be mediated through the effects of mechanical tension on the maturation of focal adhesions. That is, tension-mediated changes in focal adhesion maturation could affect adhesion signaling and thereby the coupling of RhoA and ROCK activity. Nonetheless, these data importantly demonstrate that GTP-RhoA activity by itself is not an accurate measure of either ROCK or contractile activity in cells (Figs. 1 and 4).

How does RhoA couple to ROCK in an adhesion dependent manner? Given that PAK is a major tyrosine kinase in focal adhesions and is important in transducing mechanical forces into biochemical signals, it may be involved in this control mechanism. PAK is required for microtubule stabilization by another RhoA effector, mDia [13]. Tyrosine kinase activity has been shown to be required for Rho induced stress fiber formation [27] and Rho can directly induce the phosphorylation of PAK in the absence of stress fiber formation [28], suggesting the possibility of PAK activation being upstream of ROCK. The colocalization of RhoA and ROCK may also play a role in the regulation of ROCK activity by adhesion and tension, perhaps mediated by membrane lipid rafts, as has been shown in two other cases. Adhesion-mediated activation of both PAK by GTP-Rac and mDia by GTP-Rho has been shown to be mediated by GM1-ganglioside containing lipid rafts [13,14]. The dependence of lipid raft internalization or fusion to the plasma membrane by adhesion may represent a general mechanism by which Rho GTPases and their effectors are regulated. Similar mechanisms may underlie the adhesion and cytoskeletal regulation of ROCK that we observed in this study.

The bi-directional coupling between tension and ROCK activity has numerous implications. Once activated, this positive feedback loop may be an important mechanism to amplify and sustain ROCK activity, where ROCK activates tension, which in turn strengthens the RhoA–ROCK coupling. Conversely, this architecture also explains the observation that disruption of ROCK, tension, or adhesion, functionally disrupts the entire mechano-adhesive system and results often in similar phenotypes such as decreased proliferation [2,5,29]. It has recently been suggested that change in substrate stiffness may also affect adhesions and RhoA–ROCK coupling [3]. Here our findings would suggest that the decreased substrate stiffness alters the mechanical stress at adhesions generated by cytoskeletal tension, in turn decreasing ROCK activity, cytoskeletal tension, and adhesion maturation, and, thus providing a general control mechanism whereby cell adhesion, substrate mechanics, cytoskeletal tension, and cell signaling are all intricately related (Fig. 6). Since ROCK is required for generating tension, this raises the question of what generates the tension required to activate ROCK in the first place. There are other well-characterized kinases for myosin such as MLCK and PAK, that may temporally precede ROCK activation [30]. There also exist as yet uncharacterized mechanisms for generating tension independent of MLC phosphorylation [31] that may precede and contribute to ROCK activation.

RhoA–ROCK signaling has long been known as an important regulator of cytoskeletal tension and adhesion maturation. The numerous feedback loops between these adhesives, cytoskeletal, and mechanical functions of the cell and this signaling pathway highlight the complex control system that cells have constructed, perhaps in order to navigate the complex relationship between ECM mechanics, structure, and composition. Understanding this control system ultimately will be central to our understanding of how cells coordinate the chemical and mechanical worlds.

Acknowledgments

We thank D. Cohen and N. Sniadecki for a critical reading of the manuscript, and Lixin Qi for expert technical assistance. This work was funded in part by grants from the NIH (HL073305, EB00262, GM74048), the Department of Defense Multidisciplinary University Research Initiative, and the University of Pennsylvania MRSEC.

Appendix A. Supplementary data

Supplementary data associated with this article can be found, in the online version, at doi:10.1016/j.yexcr.2007.07.002.

REFERENCES


