From autoinhibition to inhibition in trans: the Raf-1 regulatory domain inhibits Rok-α kinase activity

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Introduction

The GTPases Rho, Rac, and Cdc42 control fundamental processes including cell shape, polarity, and migration but also gene expression and cell cycle progression. Thus, Rho GTPases and their effectors are promising therapeutic targets for several diseases, including cancer (Heasman and Ridley, 2008; Olson, 2008).

The Rho effectors Rok-α and -β (Riento and Ridley, 2003; Zhao and Manser, 2005) are serine/threonine kinases with a modular structure comprising an N-terminal catalytic domain, a coiled-coil region containing the Ras/Rho-binding domain (RBD), and a C-terminal regulatory region with an unusual pleckstrin homology (PH) domain interrupted by a cysteine-rich domain (CRD; Riento and Ridley, 2003). Roks are regulated by autoinhibition; their C-terminal regulatory region, particularly the PH/CRD domain, binds to the kinase domain and inhibits its activity (Amano et al., 1999; Chen et al., 2002). Interaction of two RhoA molecules with the RBD domains arranged in a parallel coiled-coil dimer relieves autoinhibition (Amano et al., 1999; Shimizu et al., 2003; Dvorsky et al., 2004) and leads to kinase domain dimerization, transautophosphorylation, and activation (Riento and Ridley, 2003; Zhao and Manser, 2005).

Raf-1, a serine/threonine kinase member of the Ras/extracellular signal-regulated kinase (ERK) signaling pathway, interacts with Rok-α (Ehrenreiter et al., 2005; Piazzolla et al., 2005). In Raf-1 knockout (KO) cells, hyperactive Rok-α causes cytoskeletal changes, leading to inhibition of cell migration domain (RBD), and a C-terminal regulatory region with an unusual pleckstrin homology (PH) domain interrupted by a cysteine-rich domain (CRD; Riento and Ridley, 2003), which is similar to Rok-α’s own autoinhibitory region. Thus, Raf-1 mediates Rok-α inhibition in trans, which is a new concept in kinase regulation. This mechanism is physiologically relevant because Raf-1reg is sufficient to rescue all Rok-α-dependent defects of Raf-1–deficient cells. Downstream of Ras and Rho, the Raf-1–Rok-α interaction represents a novel paradigm of pathway cross talk that contributes to tumorigenesis and cell motility.
Rok–Raf-1 interaction of full-length (FL) Raf-1 with a series of intramolecular autoinhibition. To test these possibilities, we alternatively, Raf-1 could interact with the negative regulatory domain (Rok–α). Raf-1 could conceivably prevent binding of Rok–α to the cell membrane and activated by a small GTPase, in this case, Ras. Raf-1 and stabilize intramolecular autoinhibition; their N-terminal regulatory domain, particularly the CRD, binds to the kinase domain, suppressing its catalytic activity (Cutler et al., 1998). Raf activation requires Ras binding, membrane recruitment, and phosphorylation of S/T sites in the activation loop of the CR3 region (Wellbrock et al., 2004).

All Raf kinases can activate the MAPK/ERK kinase (MEK)–ERK module, yet the main in vivo roles of Raf-1 in migration, survival, and Ras-induced tumorigenesis are MEK–ERK independent and rely on Raf-1’s ability to interact with and inhibit other kinases such as Rok–α (Ehrenreiter et al., 2005; Piazzolla et al., 2005; Ehrenreiter et al., 2009), MST2 (O’Neill et al., 2004), and ASK-1 (Yamaguchi et al., 2004). Until now, the mechanisms underlying this inhibition were unknown.

Negative regulation of the activity of a kinase by other kinases can occur in the context of a negative feedback loop, as does the inhibition of MEK1 by ERK (Eblen et al., 2004; Catalanotti et al., 2002), or in the context of pathway cross talk, as exemplified by the down-regulation of Raf-1 by Akt or PKA (Wellbrock et al., 2004). In these and other cases, negative regulation is achieved by direct phosphorylation of one kinase by the other. In this study, we report a novel form of kinase regulation and pathway cross talk mediated by protein–protein interaction instead of phosphorylation. Upon growth factor stimulation, GTPase binding to Raf-1 and Rok–α relieves autoinhibition, engendering a change from a closed, inactive state to an open, active conformation essential for Raf-1–Rok–α interaction. In the open state, the Raf-1 regulatory domain (Raf-1reg) binds to the kinase domain of Rok–α and inhibits its enzymatic activity directly. This kinase-independent inhibition in trans represents a new paradigm in pathway cross talk and regulation of kinase activity.

Results and discussion

Activation increases Raf-1–Rok–α interaction

In mouse embryonic fibroblasts (MEFs), Raf-1 binds to Rok–α, limiting its activation and cell membrane localization (Ehrenreiter et al., 2005). Raf-1 can conceivably prevent binding of Rok–α to RhoA by competing for or masking the Rho-binding site. Alternatively, Raf-1 could interact with the negative regulatory PH/CRD domain or the kinase domain of Rok–α and stabilize intramolecular autoinhibition. To test these possibilities, we examined the interaction of full-length (FL) Raf-1 with a series of Rok–α deletion mutants (Fig. 1 A). A mutant lacking the PH/CRD domain (ΔPH/CRD) and a truncated protein containing the kinase domain (Rok–α–K) bound to Raf-1 more efficiently than FL Raf–α (Fig. 1, B and C; and Fig. S1 A). In contrast, Raf-1 hardly interacted with the Rok–α regulatory domain (Rok–αreg; Fig. 1 B). Next, we used multiphoton fluorescence resonance energy transfer (FRET)/fluorescent lifetime imaging microscopy (FLIM) to directly monitor protein–protein interactions in cells. The fraction of FL Raf-1 bound to FL Rok–α in asynchronously growing cells was under the detection limit. However, robust interaction was detected upon coexpression of active RhoA with the FL proteins or using Rok–α–K as an acceptor (Fig. 1, D and E). FRET efficiency was much higher in cell protrusions (28%; Fig. 1 D, inset), suggesting protein accumulation and increased interaction in these locations. In line with the coinmunoprecipitation experiments, these results show that Raf-1 preferentially binds to the kinase domain of Rok–α. They rule out the possibility that Raf-1 inhibits Rok–α activation by competing with RhoA and suggest instead that RhoA favors intramolecular interaction between Rok–α and Raf-1 by disrupting the intramolecular interaction between the kinase and Rok–αreg.

Ras binding similarly disrupts the interaction between the regulatory and kinase domains of Raf-1, rendering both more accessible for intermolecular interactions (Terai and Matsuda, 2005). Indeed, EGF stimulation increased complex formation between endogenous (Fig. 2 A) or ectopically expressed proteins, as shown by both FRET/FLIM and coinmunoprecipitation experiments (Fig. 2, B and C; and Fig. S1 B). Constitutively active Ras or activation of endogenous Ras by a membrane-tethered form of the Ras guanine nucleotide exchange factor SOS (Sibilia et al., 2000) also stimulated Raf-1–Rok–α interaction (Fig. 2 D). Conversely, mutating the Raf-1 RBD (R89L) or CRD (CC165/168SS; CC/SS) significantly reduced complex formation (Fig. 2, E–G). Thus, activation by Ras is both necessary and sufficient to promote Raf-1–Rok–α interaction.

Ras binding induces a conformational change in Raf-1 and recruits it to the membrane to be phosphorylated by activating kinases (Bondeva et al., 2002). Tethering Raf-1 to the membrane by fusing it to the Ki-Ras membrane-targeting signal (Raf-1 CAAX) activates the MEK–ERK pathway (Leevers et al., 1994), but it abolished binding to Rok–α (Fig. 2 E). Thus, the change from a closed to an open conformation mediated by Ras binding is essential both for MEK–ERK activation and Raf-1–Rok–α interaction, but these two functions of Raf-1 take place in distinct subcellular compartments. Indeed, single fluorophore video tracking of Raf proteins has shown that Raf-1 binds to Ras-GTP and activates MEK–ERK in the context of membrane nanoclusters but redistributes to the cytosol when these structures dissolve (Tian et al., 2007). It is tempting to speculate that the activated Raf-1 molecules leaving the membrane may be those that bind Rok–α in vivo.

Raf-1reg binds to Rok–α and inhibits its kinase activity

The R89L and CC/SS mutations may prevent or weaken Ras binding, thus precluding the conformational change that makes Raf-1reg accessible for Rok–α; alternatively, they may be more directly involved in the interaction with Rok–α. To distinguish between these possibilities, we introduced the R89L and CC/SS mutations in Raf-1reg, which lacks the Raf-1 kinase domain.
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In contrast to FL Raf-1 R89L, Raf-1reg R89L retained the ability to coimmunoprecipitate with Rok-α (Fig. 3 A). Thus, binding of Ras-GTP to FL Raf-1 is required solely to disrupt the interaction between the regulatory and kinase domains of Raf-1. Interfering with Ras binding did not increase complex formation with Rok-α, indicating that Ras and Rok-α do not compete for Raf-1.

In contrast, Raf-1reg CC/SS, which binds to Ras but not to the Raf-1 kinase domain (Cutler et al., 1998), failed to associate with Rok-α, implying that the CRD plays a critical role in Raf-1–Rok-α complex formation (Fig. 3 B).

Raf-1 CRD might restrain the activity of Rok-α by binding directly to its kinase domain. Indeed, recombinant GST–Raf-1reg interacted with Rok-α–K in vitro, pulling down ~25% of the Rok-α–K input, whereas GST–Raf-1reg CC/SS was much less efficient (Fig. 3 C). GST–Raf-1reg, but not a CC/SS mutant, reduced Rok-α–K activity in an in vitro kinase assay (Fig. S2). Moreover, Raf-1reg inhibited recombinant Rok-α–K in a dose-dependent manner (≥70% inhibition at approximately equimolar concentrations of Raf-1reg and MLC2; Fig. 3 D). The calculated half-maximal inhibitory concentration of 2.65 µM is fairly high, but this does not prejudice the physiological relevance of the interaction per se, as exemplified by the even lower affinity (20 µM) of the Raf CRD for Ras-GTP (Williams et al., 2000). Besides, it is unclear how a half-maximal inhibitory concentration calculated in vitro using recombinant proteins relates to the physiological setting.
Raf-1reg, but not the CRD mutant, also corrected all defects of Raf-1 KO cells: it significantly improved migration (Fig. 4 C), normalized cell shape, cortical actin bundles, and ezrin phosphorylation (Fig. 4, D and E). Finally, Raf-1reg reduced Fas surface clusters and cell death in Raf-1 KO cells (Fig. S3, A and B). These results demonstrate the biological relevance of the interaction between Rok-α and Raf-1reg and formally rule out a contribution of Raf-1 kinase activity to the regulation of cell shape, migration, and Fas expression.

Raf-1reg and Rok-α reg inhibit Rok-α-K in vivo

Our data suggest that the activity of the Rok-α kinase domain, restrained in cis by its own regulatory domain (Rok-αreg) before activation (Amano et al., 1999), is inhibited in trans by to the binding affinity of the two FL, posttranslationally modified proteins in vivo. Indeed, when expressed at near-endogenous levels in KO MEFs, Raf-1reg wild type (WT), and much less so CC/SS, associated with Rok-α and decreased its kinase activity to levels similar to those observed in WT MEFs (Fig. 3 E).

Concentration of the partners in relevant subcellular compartments will also drive protein–protein interaction in vivo. FL Raf-1 and Rok-α accumulate in membrane protrusions (Fig. 1 D), and both Raf-1 (Ehrenreiter et al., 2005) and Raf-1reg colocalize with Rok-α on filamentous structures (Fig. 4 A) corresponding to the vimentin cytoskeleton. Vimentin is a direct substrate of Rok-α, which by phosphorylating it contributes to its depolymerization (Sin et al., 1998). Vimentin collapses in juxtanuclear aggregates in Raf-1–deficient cells, a phenotype rescued by Raf-1reg (Fig. 4 B). Thus, Raf-1reg is sufficient to mediate the correct localization of Rok-α to the vimentin cytoskeleton and to inhibit Rok-α activity, preventing the collapse of these intermediate filaments. In addition to the vimentin defects, Raf-1 KO cells are contracted and characterized by cortical actin bundles. They contain higher amounts of phosphorylated ezrin than WT cells, and their migration is impaired (Fig. 4, C–E). Finally, the death receptor Fas is found in characteristic clusters on the surface of Fas of Raf-1 KO cells, which are more sensitive to Fas-induced cell death (Fig. S3, A and B). All of these defects are caused by Rok-α hyperactivity and can be rescued by chemical inhibition of Rok-α, by expressing dominant-negative Rok-α, or by silencing the Rok-α gene (Ehrenreiter et al., 2005; Piazzolla et al., 2005).

Raf-1reg, but not the CRD mutant, also corrected all defects of Raf-1 KO cells: it significantly improved migration (Fig. 4 C), normalized cell shape, cortical actin bundles, and ezrin phosphorylation (Fig. 4, D and E). Finally, Raf-1reg reduced Fas surface clusters and cell death in Raf-1 KO cells (Fig. S3, A and B). These results demonstrate the biological relevance of the interaction between Rok-α and Raf-1reg and formally rule out a contribution of Raf-1 kinase activity to the regulation of cell shape, migration, and Fas expression.

**Figure 2. Activated Raf-1 preferentially interacts with Rok-α.** (A–C) EGF increases the Rok-α–Raf-1 interaction. (A) MEFs were stimulated with 10 ng/ml EGF, and endogenous Rok-α was immunoprecipitated at the indicated time points. (B and C) Fluorescence lifetime (τ), GFP intensity, and RFP intensity in MDA-MB-468 transfected with GFP-FL Raf-1 and mRFP-FL Rok-α upon stimulation with 100 ng/ml EGF. (C) Percentage of FRET efficiency is shown. Error bars indicate SEM (n > 3). (D) Activated Ras promotes Rok-α–Raf-1 interaction. COS-1 cells were transfected with HA-tagged FL Rok-α, constitutively active Ras (RasV12), or membrane-tethered SOS-F, resulting in the constitutive activation of endogenous Ras and the corresponding vectors (V). UT, untransfected COS-1 cells; *, endogenous Ras. Black lines indicate that intervening lanes have been spliced out. (E–G) Ras binding and subcellular localization affect Rok-α–Raf-1 interaction. (E) COS-1 cells were transfected with HA-tagged FL Rok-α and the indicated FL Raf-1 mutants. HA immunoprecipitates were analyzed and quantified as described in Fig. 1. (F) Fluorescence lifetime, GFP intensity, and mRFP1 intensity in MCF-7 cells transfected with GFP-FL Raf-1 WT or CC/SS mutant (donor) and mRFP1–Rok-α–K (acceptor). The cell marked with the asterisks was excluded from the cumulative FRET efficiency analysis in G as a result of insufficient photon counts (see Materials and methods). (G) Percentage of FRET efficiency is shown. (A, E, and G) Error bars indicate SD of three experiments. *, P < 0.05; **, P < 0.01; ***, P < 0.005. Bars: (B) 20 µm; (F) 30 µm.
Raf-1reg once activation has occurred. A computational model of the Rok-α CRD, based on the structure of the autoinhibitory CRD of Raf-1 (Mott et al., 1996), is compatible with this idea (Fig. 5 A). More importantly, both Rok-αreg and Raf-1reg inhibit the activity of cotransfected Rok-α-K in vivo, reducing the phosphorylation of Rok-α downstream targets by a comparable extent (Fig. 5 B). Finally, Rok-αreg and Raf-1reg, but not Rok-αreg CC/SS, reduced the hyperphosphorylation of ezrin as a result of hyperactive endogenous Rok-α in Raf-1 KO cells (Fig. 5 C). Thus, the regulatory domains of Rok-α and Raf-1 are similarly effective in regulating Rok-α activity in vivo, supporting a model in which activated Raf-1 modulates Rok-α.

Figure 3. Raf-1reg interacts with Rok-α and inhibits its kinase activity. (A and B) CRD but not RBD mutation disrupts the binding of Raf-1reg to Rok-α. HA immunoprecipitates (IP) were analyzed as in Fig. 1. The mean ± SD of at least three experiments is shown. **, P < 0.01. (C) Rok-α-K interacts in vitro with Raf-1reg WT but not with Raf-1reg CC/SS. 2 µg GST–Raf-1reg WT, or GST–Raf-1reg CC/SS on glutathione Sepharose beads were incubated with 25 ng Histagged Rok-α-K. Rok-α-K and GST proteins were detected by immunoblotting with His and GST antibody. 3.125–25 ng recombinant Rok-α-K was loaded as a reference on the same gel. One representative experiment out of three is shown. Black lines indicate that intervening lanes have been spliced out. (D) Dose-dependent inhibition of Rok-α-K by purified Raf-1reg. 0.29–9.3 µM Raf-1reg was incubated with 100 ng Rok-α-K (0.05 µM) before a Rok kinase assay with 7 µM recombinant MLC2 (recMLC2) as a substrate. The mean ± SD of three experiments is shown. (E) Raf-1reg inhibits Rok-α activity in vivo. The activity of endogenous Rok-α (eRok-α), immunoprecipitated from WT and KO MEFs, and from KO MEFs transfected with Raf-1reg WT or Raf-1reg CC/SS was assessed as in D. Rok-α activity, expressed as pMLC/MLC ratio and normalized by the amount of Rok-α present in the assay, is indicated below each lane. Rok-α activity of Raf-1 KO MEFs is set to 100%.

Figure 4. Raf-1reg colocalizes with Rok-α and rescues all phenotypes of Raf-1 KO MEFs. (A) Raf-1 KO MEFs expressing Raf-1reg were identified by staining with antibodies against Raf-1. The localization of Raf-1 and Rok-α in migrating MEFs was determined by immunofluorescence. (B–D) Raf-1reg improves cytoskeletal organization and migration. (B) Raf-1reg rescues vimentin cytoskeleton collapse in Raf-1 KO MEFs. Raf-1 KO MEFs cotransfected with pEGFP and pCMV (V) or pCMV Raf-1reg were stained with vimentin antibodies and analyzed by confocal microscopy. UT, untransfected cells. (C) Migration of Raf-1 KO MEFs transfected with the indicated pEGFP constructs was assessed using 10% FCS as a chemoattractant. The percentage of transfected cells migrating to the lower compartment of a Boyden chamber in 2.5 h is plotted. (D and E) Raf-1 KO MEFs were cotransfected with pEGFP and the indicated pCMV constructs, stained with phalloidin to visualize filamentous actin (D) or with anti-ezrinpT567 (E), and analyzed by confocal microscopy. The number of cells displaying cortical actin bundles (CAB) or prominent ezrin phosphorylation is plotted on the right. Arrowheads indicate EGFP-expressing cells. Error bars indicate SD of three experiments. **, P < 0.01.
Thus far, we don’t have any evidence that Rok-α modulates Raf-1 activity. The Raf-1 kinase domain does not bind to Rok-α, and the regulatory domain is not a Rok-α substrate in vitro (unpublished data). It is possible that Rok-α regulates Raf-1 by promoting its localization to intermediate filaments, thereby bringing it in the proximity of specific substrates. Further studies will be needed to clarify this issue.

Implications for transformation

Ras, Rho, and their downstream effectors are implicated in tumorigenesis. A good example of Ras–Rho cross talk is the suppression of Rho signaling by Ras/ERK in transformed cells, leading to increased motility. This is achieved either at the level of integrin-mediated Rho activation, which is impaired by the product of the ERK target gene fra-1 (Vial et al., 2003) or, more specifically, by uncoupling Rho activation from its downstream effector Rok. In particular, Rok expression can be reduced by ERK activation in Ras-transformed cells with high levels of active Rho (Sahai et al., 2001; Pawlak and Helfman, 2002b) and in v-src–transformed cells (Pawlak and Helfman, 2002a).

Our data identify a novel, ERK-independent mechanism by which Ras selectively regulates Rho signaling by promoting interaction between the top-tier kinases Raf-1 and Rok-α. We have recently shown the significance of this interaction in a model of Ras-driven epidermal tumorigenesis (Ehrenreiter et al., 2009) in which Ras causes transformation by inducing proliferation and survival (Sibilia et al., 2000) and by selectively blocking differentiation. We found that Ras mediates this block by promoting Raf-1–Rok-α interaction and the inhibition of Rok-α activity. If Raf-1 is ablated, both development and maintenance of the Ras-driven tumors are abrogated (Ehrenreiter et al., 2009). Understanding the mechanisms underlying the interaction between Raf-1 and Rok-α may hold promise for the design of novel, specific inhibitors for therapeutic treatments.

Materials and methods

Plasmids

The following plasmids were used in transient expression experiments: pX40-HA-FL Rok-α, ΔPH/CRD, Rok-α-K, Rok-αreg (Leung et al., 1996), pEFmyc FL Raf-1, pCMV5 FL Raf-1, Raf-1reg, Raf-1reg R89L (provided by W. Kolch, System Biology Institute, Dublin, Ireland; Kubicek et al., 2002; O’Neill et al., 2004), pEXV FL Raf-1, R89L, CC/SS, CAAX (provided by J.F. Hancock, University of Texas Medical School, Houston, TX; Roy et al., 1997), pEGFP Raf-1reg (provided by R.M. Lafrenie, Northern Ontario School of Medicine, Sudbury, Ontario, Canada; Zhang et al., 2002), pRsv FL Raf-1, Raf-1reg, and Raf-1 K (Bruder et al., 1997). For expression in bacteria, pGEX Raf-1reg (aa 1–187) was subcloned from pGEX Rok-αreg (aa 1–258; O’Neill et al., 2004), pXJ40-HA–Rok-α, Rok-α–K, Rok-αreg (Leung et al., 1996), pEFmyc FL Raf-1, pCMV5 FL Raf-1, Raf-1reg, Rok-αreg R89L (provided by W. Kolch, System Biology Institute, Dublin, Ireland; Kubicek et al., 2002; O’Neill et al., 2004), pEXV FL Raf-1, R89L, CC/SS, CAAX (provided by J.F. Hancock, University of Texas Medical School, Houston, TX; Roy et al., 1997), pEGFP Raf-1reg (provided by R.M. Lafrenie, Northern Ontario School of Medicine, Sudbury, Ontario, Canada; Zhang et al., 2002), pRsv FL Raf-1, Raf-1reg, and Raf-1 K (Bruder et al., 1997). For expression in bacteria, pGEX Rok-αreg (aa 1–187) was subcloned from pGEX Rok-αreg (aa 1–258; O’Neill et al., 2004) by PCR amplification and ligation. All CC/SS mutations were generated by site-directed mutagenesis and verified by sequencing. Monomeric RFP1 (mRFP1)–Rok-α constructs were generated by PCR amplification of pX40-HA–Rok-α and subcloned into the pcDNA mRFP1 vector. pGEX KG MLC2 and RhoA V14 Flag tagged were generated by PCR amplification of pXJ40-HA–Rok-α, Rok-α–K, Rok-αreg (Leung et al., 1996), pEFmyc FL Raf-1, pCMV5 FL Raf-1, Raf-1reg, Raf-1reg R89L (provided by W. Kolch, System Biology Institute, Dublin, Ireland; Kubicek et al., 2002; O’Neill et al., 2004), pEXV FL Raf-1, R89L, CC/SS, CAAX (provided by J.F. Hancock, University of Texas Medical School, Houston, TX; Roy et al., 1997), pEGFP Raf-1reg (provided by R.M. Lafrenie, Northern Ontario School of Medicine, Sudbury, Ontario, Canada; Zhang et al., 2002), pRsv FL Raf-1, Raf-1reg, and Raf-1 K (Bruder et al., 1997). For expression in bacteria, pGEX Raf-1reg (aa 1–187) was subcloned from pGEX Rok-αreg (aa 1–258; O’Neill et al., 2004) by PCR amplification and ligation. All CC/SS mutations were generated by site-directed mutagenesis and verified by sequencing. Monomeric RFP1 (mRFP1)–Rok-α constructs were generated by PCR amplification of pX40-HA–Rok-α and subcloned into the pcDNA mRFP1 vector. pGEX KG MLC2 and RhoA V14 Flag tagged were provided by E. Sahai (Cancer Research UK, London, England, UK) and A. Ridley (King’s College London, London, England, UK), respectively.

Cell culture and transfection

3T3-like MEFs derived from c–Raf-1−/− and WT embryos (Mikula et al., 2001), COS-1, MCF-7, and MDA-MB-468 cells (which express a high amount of EGF receptor; Filmus et al., 1985) were maintained in DME with 10% FCS and transiently transfected using Lipofectamine reagents (Invitrogen) according to the manufacturer’s instructions.

Figure 5. The regulatory domains of Raf-1 and Rok-α act as inhibitors of Rok-α kinase activity in vivo. [A] Comparison of the experimental solution structure of Raf-1 CRD (left) and the computational model of the Rok-α CRD (middle). Zinc cations are shown as spheres, and the side chains of the residues coordinating the cations are shown as lines: red in one metal site and blue in the other. [right] Superposition of the Raf-1 and Rok-α CRDs. [B] COS-1 cells [B] and MEFs [C] were transfected with the indicated constructs. 24 h after transfection, cells were lysed and analyzed by immunoblotting. KO, Raf-1 KO MEFs; *, unspecific band. [D] Model of the regulation of Rok-α by Raf-1. GTPase binding disrupts intramolecular interaction between the regulatory and kinase domains of Raf-1 and Rok-α, upon which Raf-1reg binds to the kinase domain of Rok-α, restraining Rho-induced Rok-α kinase activity. Inhibition in trans limits the phosphorylation of Rok-α downstream targets, regulating cell motility and differentiation.
Migration assay
Migration was assessed in a modified Boyden chamber as described previously [Ehrenreiter et al., 2005]. Migrating and nonmigrating EGF-transfected cells were visualized and quantified (≥450 cells/sample) by epifluorescence microscopy.

Immunofluorescence
Raf-1, Rok-α, vimentin, actin, ezrin567 and Fas were performed as described previously [Ehrenreiter et al., 2005; Piazza et al., 2005]. For Raf-1 and Rok-α staining, cells plated on fibronectin (Invitrogen) were permeabilized (0.01% Triton X-100), fixed in 4% PFA, and blocked with 0.2% gelatin before incubation with primary antibodies (Raf-1 and Rok-α; BD) and staining with the appropriate Alexa Fluor 488- or 594-conjugated secondary antibodies (Invitrogen). Rhodamine-conjugated phalloidin (Invitrogen) was used to visualize actin filaments.

To visualize vimentin, intermediate filaments cells were fixed in methanol containing 5 mM EDTA and permeabilized with 0.5% Triton X-100. Cells were subsequently stained with vimentin antibody (Sigma-Aldrich) followed by Alexa Fluor 594–conjugated secondary antibodies. For ezrin567 staining, cells were fixed in cold methanol/5 mM EDTA and blocked (10% goat serum/1% BSA) before incubation with phospho ezrin-radixin-moesin antibody (p567; Cell Signaling Technology) followed by Alexa Fluor 594–conjugated secondary antibodies.

For Fas staining, cells were fixed in cold methanol/5 mM EDTA for 10 min at room temperature followed by Alexa Fluor 594–conjugated secondary antibodies. Anti-Fas reagent (ProLong Anti-fade; Invitrogen) was used as a mounting medium.

Confocal microscopy was performed at room temperature with a microscope (Axiovert 100M; Carl Zeiss, Inc.) fitted with a Plan Apochromat 63x/1.40 NA oil objective and equipped with the confocal laser-scanning module (LSM 510; Carl Zeiss, Inc.). Immersion (518; Carl Zeiss, Inc.) was used as imaging medium. Images were acquired using the LSM 510 software (version 2.3; Carl Zeiss, Inc.). Representative Z-stacks are shown. 600 transfected cells were counted for the quantification.

Cell lysates, immunoprecipitation, and immunoblotting
Cells were washed with ice-cold PBS and lysed in 200 mM Tris-HCl, pH 7.4, 2 mM EDTA, and 1% Triton X-100 with protease and phosphatase inhibitors. Lysates and HA–Rok-α immunoprecipitates were prepared from subconfluent cells 24–48 h after transfection and analyzed by immunoblotting using the following antibodies: Rok-α (Millipore), HA (12CA5), Raf-1, SOS (BD), pCotilin53, Cotillin, pMLC2/519 [Santa Cruz Biotechnology, Inc.], pERK, pEzrin567, ezrin–radixin–moesin (Cell Signaling Technology), tubulin (Sigma-Aldrich), and pan-Raf-1567 (EMD). The amount of Raf-1 proteins in the immunoprecipitation was quantified by densitometry (ImageQuant [GE Healthcare] or AlphaEase [Alpha Innotech]) and normalized to the amount of immunoprecipitated Rok-α.

Protein expression and purification
GST–Raf-1reg proteins were expressed in Escherichia coli Rosetta (DE3; EMD) by induction with 1 mM IPTG and incubation in minimal medium (MB). The amount of recombinant Raf–1reg and MLC2 were obtained by thrombin cleavage eluted with 20 mM reduced glutathione in 50 mM Tris-HCl, pH 8.0. Recombinant Raf–1reg and MLC2 were obtained by thrombin cleavage (6 U/ml overnight at 4°C) as previously described (Wyckoff et al., 2006).

GST pull-down and Rok-α in vitro kinase assays
GST–Raf-1reg immobilized on glutathione Sepharose beads (GE Healthcare) and eluted with 20 mM reduced glutathione in 50 mM Tris-HCl, pH 8.0. The recombinant Raf–1reg and MLC2 were obtained by thrombin cleavage (6 U/ml overnight at 4°C) as previously described (Wyckoff et al., 2006).

FRET determination by multiphoton FLIM
Time domain FLIM was performed at room temperature with a multiphoton microscopy system comprised of a solid state–pumped (Verdi BW; Coherent, Inc.), femtosecond self-mode-locked Ti:Sapphire laser system (Mira; Coherent, Inc.), an in-house–developed scan head, and an inverted microscope (TE2000E; Nikon; Peter et al., 2005; Festy et al., 2007). FRET was monitored by the conventional equation: FRET efficiency = 1 - (F-R/F-R0)/(F-F0/F-F0), where F-F0 is the GFP–Raf–1 in cells that coexpress mRFP–Rok–α, and τ_F0 is the GFP–Raf–1 lifetime measured in the absence of an acceptor. Because ≤100-ps time resolution is achieved with our instrumentation, for a τ_F0 value of 2.35 ns, FRET efficiencies as low as 3% can be determined accurately. Pixel by pixel lifetime determination was achieved using a modified Levenberg–Marquardt fitting technique (Barber et al., 2005). The error in fitting the monoeponential decay model for fluorescence lifetime determination is <0.4% for signals with a peak of ≥500 photon counts. Also, in general, the lifetime of the interacting population (FRET species) can only be accurately determined with a peak photon count of ≥500 (Barber et al., 2009). We have therefore routinely excluded cells that have insufficient photon counts (<500 photons at the peak) from lifetime analysis.

Statistical analysis
All values are expressed as mean ± SD of at least three independent experiments unless indicated otherwise. P-values were calculated using the unpaired, two-tailed Student’s t-test. P ≤ 0.05 is considered statistically significant.

Online supplemental material
Fig. S1 shows that FL Raf-1 and Raf-1reg, but not the Raf-1 kinase domain, interact with FL Rok-α and ΔPH/CRD Rok-α. Fig. S2 shows that Raf-1, but not Raf-1 reg CC/SS, inhibits the kinase activity of Rok-α in vitro. Fig. S3 shows that expression of Raf-1 prevents the formation of Fos clusters at the cell surface of Raf-1 KO cells and reduces their sensitivity to Fas-induced cell death. Online supplemental material is available at http://www.jcb.org/cgi/content/full/jcb.200906178/DC1.

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