Abstract

Cell migration involves a multitude of signals that converge on cytoskeletal reorganization, essential for development, immune responses and tissue repair. Using knockdown and dominant negative approaches, we show that the microtubule-associated Ste20-like kinase SLK is required for focal adhesion turnover and cell migration downstream of the FAK/c-src complex. Our results show that SLK co-localizes with paxillin, Rac1 and the microtubules at the leading edge of migrating cells and is activated by scratch wounding. SLK activation is dependent on FAK/c-src/MAPK signaling, whereas SLK recruitment to the leading edge is src-dependent but FAK independent. Our results show that SLK represents a novel focal adhesion disassembly signal.

Introduction

Migration is required for numerous biological processes such as development, tissue repair and regeneration. Signal transduction events governing cell migration involve an ever-expanding number of molecules functioning in interconnected biochemical pathways regulating the turnover of adhesion complexes at the leading edge of migrating cells. Stimulation of cell adhesion and migration induces the formation of integrin-FAK-src complexes required for the recruitment and activation of a number of adaptor molecules leading to focal adhesion turnover and migration [1–3]. Indeed, FAK-null cells assemble large and stable adhesion complexes leading to migratory deficits [4]. Similarly, a FAK mutant at tyrosine 397, deficient for c-src binding, fails to induce focal adhesion disassembly in FAK-deficient fibroblasts [5–7]. Supporting this, src-family kinase-deficient cells or cells expressing kinase inactive v-src display larger focal adhesions that fail to disassemble [8,9].

In addition to an amino-terminal serine/threonine kinase domain, the Ste20-like kinase SLK bears a central coiled-coil domain and a carboxy-terminal AT1-46 [10] homology (ATH) domain [11,12] of unknown function. Elevated SLK expression and activity leads to rapid actin stress fiber disassembly in a Rac1-dependent manner [13]. We have previously shown that SLK localizes to vinculin-rich ruffles at the cell periphery in spreading fibroblasts, suggesting a role for SLK in adhesion dynamics [13]. Consistent with a role in cytoskeletal rearrangements, SLK has been shown to indirectly associate with the microtubule network [13] and is required for fusion of C2C12 myoblasts into differentiated myotubes [14]. Interestingly, SLK has also been shown to regulate cell cycle progression [15]. In addition, SLK overexpression has been shown to induce an apoptotic response [12]. Supporting a role for SLK in cell death and cellular stress, cleavage of SLK by caspase 3 results in its activation [11]. Similarly, anoxia-recovery also activates a SLK/p38-dependent apoptotic response [16].

Our previous studies showed that SLK overexpression induced a rapid actin stress fiber disassembly that could be partially rescued by co-expression of dominant negative Rac1 [13]. Furthermore, fibroblasts expressing an activated SLK c-terminal truncation failed to assemble large peripheral adhesions during spreading on fibronectin, suggesting that SLK is an important regulator of cytoskeletal dynamics [13]. Here we show that SLK co-localizes with microtubules and adhesion components at the leading edge of migrating cells. We demonstrate that SLK is activated following scratch wounding of fibroblast monolayers in a FAK-src-MAPK-dependent manner. We find that SLK knockdown or expression of a dominant negative version results in impaired microtubule-dependent adhesion turnover and delayed migration. Overall our results show that SLK is a novel regulator of focal adhesion turnover and cell migration.

Results

SLK is activated by monolayer wounding and is required for cell migration

We have previously shown that SLK can be co-precipitated with β-tubulin and that it localizes to membrane ruffles at the
periphery of spreading fibroblasts [13]. In addition, SLK appears to induce actin stress fiber breakdown through a Rac1-mediated pathway [13]. As the signaling pathways activated during cell spreading also regulate cell motility [1,2,17,18], we tested the possibility that SLK may play a role in cell migration.

To test this, we initially investigated the localization of SLK and other cytoskeletal markers following scratch wounding of fibroblast monolayers. Co-immunostaining of SLK with actin stress fibers shows that, in addition to a perinuclear distribution, it is also enriched at the leading edge of migrating cells but not along stress fibers (figure 1A–C). Similarly, at the leading edge, SLK was found to co-localize with paxillin and Rac1 in structures reminiscent of membrane ruffles (figure 1D–F and J–L). Interestingly, it did not localize to large focal adhesions as evidenced by the lack of co-localization with paxillin at these sites (figure 1D–F). Supporting our previous results demonstrating SLK-tubulin co-precipitation [13], SLK co-localized with the microtubule network at the leading edge (figure 1G–I). These observations suggest that SLK is recruited at the leading edge of migrating cells with other adhesion signaling proteins.

Scratch wounding of confluent monolayers has been shown to induce polarization and migration [19,20]. Therefore, we
investigated SLK kinase activity at various time points following scratch wound induced migration of fibroblast monolayers. Immunoprecipitation and in vitro kinase assays show that SLK kinase activity is markedly increased following scratch wounding of confluent fibroblasts, with a peak of activity at 60 minutes followed by a decline at 90 minutes (figure 2). Extended time courses up to 120 minutes have shown that SLK activity does not return to basal levels observed at time 0 (not shown). This is likely due to the continued cell migration that occurs following wounding. As previously reported, GSK3β was found to be inactivated (pGSK3βSer9) over the time course indicating that cell migration has been induced. WCL, whole cell lysate.

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Figure 2. SLK is activated by scratch wounding and is required for cell migration. Confluent MEF 3T3 fibroblasts plated on fibronectin (10 μg/ml) were stimulated to migrate by scratch wounding. SLK kinase activity was assayed in vitro and found to increase over time reaching a maximum activity by 60 minutes following wounding. Total immunoprecipitated SLK levels are shown (lower panel). As previously reported, GSK3β was found to be inactivated (pGSK3βSer9) over the time course indicating that cell migration has been induced. WCL, whole cell lysate.

Activation and redistribution of SLK requires FAK/src/ MAPK signaling

The assembly and activation of a FAK/c-src complex during cell motility appears to be required for efficient focal adhesion turnover [7,27]. In addition, the activated FAK/c-src complex can recruit multiple signaling adapters and activate downstream signaling through several pathways [2,3,28–31]. The activation of SLK following scratch wounding of monolayers appears to be a relatively late event (60 min; figure 2), suggesting that upstream signaling may be necessary prior to SLK activation. Because of the ultimate requirement for the FAK-c-src complex in cell migration, we tested whether src family kinases or FAK were required for SLK activation in a scratch wound assay.

To test the involvement of src family kinases in SLK regulation, monolayers were pretreated with the Src-family inhibitor PP2 (or PP3 control) and then subjected to wounding in the presence of the inhibitor. As shown in figure 6A, treatment with the control PP3 resulted in SLK activation 60 minutes following scratch wounding. Interestingly, treatment with PP2 resulted in an increase in SLK activity in unscratched confluent
monolayers that could not be further increased by wounding. Similar results were obtained in SYF cells and rescued when c-src was expressed (not shown). This suggests that src family kinases are required to negatively regulate SLK activity. Supporting this, our previous results have shown that overexpression of v-src inhibits SLK kinase activity in a CKII-dependent manner [32].

To investigate the role of FAK in SLK regulation, scratch wounding assays were performed on FAK(-/-) fibroblasts and

Figure 3. SLK knockdown or expression of a dominant negative SLK inhibits cell migration. Subconfluent MEF 3T3 cells were infected with Adenovirus vectors expressing DN SLK (Ad-HA-K.DC) or LacZ control and subjected to fibronectin (FN) transwell migration assays. (A) Western blot analysis of HA-K.DC expression. Cdc42 was used as a loading control. (B) Polycarbonate membranes were DAPI stained and cells on the underside were enumerated (C) in random fields and expressed as the average/field from triplicate wells. (D) MEF 3T3 cells were transfected with SLK siRNAs and analysed for SLK expression. Western blot analysis of treated lysates indicates that SLK siRNA at 10 pM resulted in a marked knockdown of SLK. Reprobing the membrane with a β-tubulin antibody was used as a control for loading (lower panel). (E–F) Cells were treated with SLK-specific or control siRNAs and assayed for migration through a chamber coated with bovine serum albumin (BSA) (10 μg/ml) (control) or fibronectin (FN) (10 μg/ml). In both cases a 60–70% reduction in migration was observed. (G) Confluent MEF3T3 cells were infected with Adenovirus vectors expressing a scramble or SLK shRNA and manually scratched with a pipet tip. Wound closure was followed for 12 h and the percent closure was evaluated.
wildtype controls. Following wounding of FAK wildtype monolayers, SLK kinase activity is upregulated to levels comparable to that of MEF3T3 cells (figure 6B). However, little or no SLK upregulation was observed following the wounding of FAK-null monolayers (figure 6B), suggesting that adhesion signaling is required and that SLK activation is not a secondary effect of monolayer wounding. Similarly, scratch wounding in the presence of the MEK1 inhibitor U0126 prevented SLK upregulation when compared to a DMSO control (figure 6C). For the same time course, scratch wounding resulted in ERK1/2 phosphorylation whereas it was markedly reduced in the presence of U0126 (figure 6C). Similar experiments in the presence of the p38 inhibitor SB203580 showed no effect on SLK activation (not shown). Overall, these results suggest that SLK activation by scratch-induced motility requires the FAK/c-src/MAPK signaling system.

Cell spreading and scratch wounding of fibroblast monolayers results in the recruitment of a proportion of SLK protein at the cell periphery or the leading edge, respectively (figure 1 and [13]). Microtubule dynamics and stability have been shown to be regulated by src family kinases in various systems, including stabilization by integrin-mediated FAK signaling [33–36]. Therefore, we investigated whether SLK recruitment to the leading edge was also FAK/c-src dependent. Monolayers of FAK-null or SYF (src/yes/fyn triple knock-out), as well as wildtype controls, were scratch wounded and co-immunostained for SLK and Rac1. Although FAK-null cells do not migrate efficiently, SLK was detected, along with Rac1, at the leading edge (figure 7C–D). Similarly, SLK and Rac1 were recruited to ruffles and the leading edge in U0126-treated cells (figure 7E and F). However, little or no SLK could be detected at the leading edge of wounded SYF monolayers (figure 7G–H). Similarly, Rac1 distribution was impaired in those cells. The distribution of both Rac1 and SLK was restored when c-src was re-expressed in SYF cells, suggesting that c-src expression is sufficient to recruit SLK at the leading edge in migrating cells.

Discussion

Our previous studies have shown that SLK co-localizes with vinculin and Rac1 in the membrane ruffles and lamellipodia of spreading fibroblasts [13]. In addition, we have shown that SLK can indirectly associate with the microtubule network and mediate actin stress fiber dissolution in a Rac1-dependent manner [15]. Here, we show that SLK can be co-localized with paxillin, α-tubulin and Rac1 at the leading edge of migrating cells and that its activity is upregulated by scratch wounding of fibroblast monolayers. A reduction in SLK levels or activity negatively affects cell migration and microtubule-dependent focal adhesion turnover, suggesting that SLK is required for cell motility. Cell migration by monolayer wounding stimulates SLK activity in a FAK/c-src/MAPK signaling dependent manner. In addition, efficient recruitment of SLK to the leading edge of migrating cells required c-src.
Overall, our data suggest that SLK is an important regulator of focal adhesions/contacts dynamics. The activation and recruitment of SLK at the leading edge of migrating cells may be necessary for the destabilization of focal contacts, a process required for further protrusive activity and motility [2,17,23,37]. Focal adhesion/contact disassembly would also destabilize the actin network, a phenotype previously reported to be induced by SLK overexpression [11,13].

The microtubule network has been shown to be tightly linked to cell adhesion and motility [26,36,38]. It has been reported that microtubules target focal contacts to modify their characteristics, including their disassembly [21,39]. In addition, the microtubule network has been reported to be stabilized by FAK and Rho GTPase signaling at the leading edge of migrating cells [21,39]. The association of SLK with the microtubule and its requirement for efficient focal adhesion turnover suggests that it is a novel microtubule-associated signal required for cell migration.

Interestingly, inhibition of src family kinases by PP2 results in an upregulation of SLK activity in confluent unscratched monolayers, suggesting that src family kinases negatively regulates SLK (see figure 6) which is alleviated by PP2 treatment. Consequently, SLK cannot be further activated by wounding. Supporting this, we have previously shown that SLK is not tyrosine phosphorylated and that v-src overexpression results in SLK downregulation through casein kinase II [32]. Although it cannot be activated, SLK can still be recruited to the leading edge of FAK-null cells. However, SLK recruitment in these structures is impaired in SYF cells (see figure 6).

One possibility is that the initial c-src recruitment and activation during focal contact assembly [42–46] recruits the microtubule network and SLK. Our previous results show that src can activate CKII, leading to direct downregulation of SLK [32]. Therefore, it is possible that SLK is kept inactive until src family kinases have been downregulated. Following the subsequent inactivation of c-src through csk or protein phosphatases [47–49], SLK activity can be upregulated through a MAPK pathway (figure 8). As the recruitment of SLK to the leading edge occurs in FAK-null but not in SYF cells, it is likely that microtubule recruitment through c-src is FAK independent. One possibility is that, in FAK(-/-) cells, some aspects of the Pyk2/c-src signaling complex can compensate for the loss of FAK in the recruitment of SLK [4,50]. However,
Pyk2 cannot substitute for FAK which appears to be necessary for SLK activation, through a MAP kinase-dependent pathway [31]. Overall our data show that SLK is activated by FAK/c-src/MAPK signaling during cell migration. Its recruitment to the leading edge in a c-src-dependent manner is required for focal adhesion turnover. Whether SLK activation occurs prior to recruitment or following c-src downregulation is still unclear. Similarly, whether it is mediated by direct MAPK phosphorylation or other MAPK-dependent events remains to be elucidated. Similarly, it is not known whether members of the Rho family of GTPases impinge on SLK activation or recruitment, or on its ability to destabilize actin. The identification of SLK substrates or downstream signaling systems will further our understanding on the role of SLK in cell migration.

Materials and Methods

Cell culture and migration assays

MEF 3T3, wildtype, FAK(-/-) and SYF (src/fyn/yes triple mutant) cells were all maintained in Dulbecco’s modified MEM (DMEM, Gibco) supplemented with 10% fetal bovine serum (FBS, Gibco), 2 mM L-glutamine (Gibco) and penicillin G (200 U ml⁻¹, Gibco) in a humidified 37°C incubator at 5% CO₂. For migration assays, MEF 3T3 cells were either infected with adenovirus or treated with control or SLK siRNAs and serum starved overnight (DMEM+0.5% FBS). Cells (1–3×10⁴) were then resuspended in DMEM containing 0.5% BSA and added to the top of a Boyden transwell migration chamber pre-coated with fibronectin (10 µg ml⁻¹) and allowed to migrate for 3–6 hours. Residual cells were removed from the top of the chamber and the filter was rinsed in PBS, fixed in 4% PFA for 10 minutes and stained with DAPI (0.5 µg ml⁻¹, Sigma). The cells that migrated to the underside of the filter were enumerated from 5 to 10 random fields using DAPI fluorescence. Cell counts were performed in triplicate for three independent experiments. Representative experiments are shown.

Scratch wound induced migration was performed as described [19]. Briefly, MEF 3T3 cells were plated on fibronectin-coated (10 µg ml⁻¹) dishes and serum-starved confluent monolayers were then scratched with a pipette tip until approximately 50% of the monolayer was removed. Cells were then washed with PBS, refed and collected at various time points. In some experiments, the monolayers were pre-incubated for 60 minutes with 10 µM of PP3, PP2 (EMD-Calbiochem) or U0126 (Cell Signaling) and scratched wounded in the presence of inhibitors or DMSO control. For microtubule-dependent adhesion turnover assays [25], serum-starved subconfluent MEF 3T3 cells were treated with 10 µM nocodazole for 3 hours. The cultures were then washed 4 times with serum-free medium and refed with DMEM/0.5% FBS for the duration of the time course. Cultures were harvested at different time points and surveyed for pFAK-Tyr397 (BioSource) by Western blotting.

SiRNA Knockdown and Adenovirus infections

MEF 3T3 cells plated at a density of 1–3×10⁵ in 60 mm plates were transfected with 5 or 10 pM SLK siRNA (Dharmacon) duplex (5’- GGUUGAGAUUGACAUAUUA) using Lipofectamine 2000 transfection reagent (Gibco) according to manufacturer-
ers recommendations. In some experiments, the cells were infected with an adenoviral vector expressing an SLK suppressor hairpin RNA (psiStrike; Promega) that consisted in the same siRNA sequence. Cells were collected 48 hours post-transfection and assayed for cell migration in transwell inserts and protein expression by western blot analysis. Control siRNAs consisted of Dharmacon’s non-targeting duplex. Similar results were obtained with scrambled SLK siRNAs. To monitor the effect of kinase deficient SLK on cell migration, adenoviral vectors expressing kinase inactive SLK (HA-KJCa: aa 1–373 with an ATP-binding site mutation; Lys 63 >> Arg) or a control (GFP or LacZ) were used to infect MEF 3T3 cultures [15]. Cells were infected at a MOI of 10 by the addition of the adenovirus directly to the cells in 0.25% FBS-DMEM 16 hours before migration assays. Expression of the SLK constructs was confirmed by anti-HA immunoblotting. LacZ and GFP expression was confirmed by immunoblotting with anti-beta-galactosidase antibody (Promega) and by epifluorescence, respectively. GFP expression was confirmed in live cells by epifluorescence.

Antibodies and immunofluorescence

The primary antibodies used in these studies were as follows: SLK polyclonal antibodies were as described previously [15]. Phospho-GSK3α/β (ser 9) and GSK3β (Cell Signaling), phospho FAK (Y397) (Biosource) and total FAK (Santa Cruz) Paxillin (BD Transduction labs), α-tubulin (Sigma), Rac1 (Santa Cruz) and phosphor ERK1/2 and total Erk1/2 (Santa Cruz) were used to infect MEF 3T3 cultures [15]. Cells were infected at a MOI of 10 by the addition of the adenovirus directly to the cells in 0.25% FBS-DMEM 16 hours before migration assays. Expression of the SLK constructs was confirmed by anti-HA immunoblotting. LacZ and GFP expression was confirmed by immunoblotting with anti-beta-galactosidase antibody (Promega) and by epifluorescence, respectively. GFP expression was confirmed in live cells by epifluorescence.

For immunofluorescence studies, MEF 3T3 cells were plated on coverslips coated with or fibronectin (10 µg/ml) and incubated overnight. The following day, monolayers were scratched and stained after 2–4 h. Briefly, the cells were rinsed with PBS, fixed in 4% PFA and blocked in PBS containing 5% goat or donkey serum overnight at 4°C in 5% BSA or skim milk powder in 1× TBST (50 mM Tris pH 7.4, 150 mM NaCl, 0.5% Nonidet P-40) and subjected to SDS-polyacrylamide gel electrophoresis (PAGE) or kinase assay. In vitro SLK kinase assays were performed following SLK immunoprecipitation as described previously [15]. Kinase reactions were stopped by the addition of 7 µl of 4× sodium dodecyl sulfate (SDS) sample buffer and electrophoresed on 8% SDS-PAGE. The gels were transferred to PVDF membranes and subjected to autoradiography followed by western blotting with anti-SLK antibody.

Author Contributions

Performed the experiments: SW PK MM. Analyzed the data: LS SW KR. Contributed reagents/materials/analysis tools: ZC CS KR MM. Wrote the paper: LS SW.

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