The nucleating activity of the Arp2/3 complex promotes the assembly of branched actin filaments that drive plasma membrane protrusion in migrating cells. Arp2/3 complex binding to nucleation-promoting factors of the WASP and WAVE families was previously thought to be sufficient to increase nucleating activity. However, phosphorylation of the Arp2 subunit was recently shown to be necessary for Arp2/3 complex activity. We show in mammary carcinoma cells that mutant Arp2 lacking phosphorylation assembled with endogenous subunits and dominantly suppressed actin filament assembly and membrane protrusion. We also report that Nck-interacting kinase (NIK), a MAP4K4, binds and directly phosphorylates the Arp2 subunit, which increases the nucleating activity of the Arp2/3 complex. In cells, NIK kinase activity was necessary for increased Arp2 phosphorylation and plasma membrane protrusion in response to epidermal growth factor. NIK is the first kinase shown to phosphorylate and increase the activity of the Arp2/3 complex, and our findings suggest that it integrates growth factor regulation of actin filament dynamics.

Introduction

The seven subunit actin-related protein (Arp) 2/3 complex nucleates actin filaments that drive diverse cell processes such as vesicle trafficking, plasma membrane protrusion during cell migration, and motility of some pathogens in host cells (Goley and Welch, 2006). Hence, resolving how activity of the Arp2/3 complex is regulated is important for a basic understanding of diverse cell processes and for targeting aberrant actin-dependent cell behaviors in diseases. Arp2/3 complex binding to nucleation promoting factors (NPFs) such as N-WASP was previously thought to be sufficient to increase nucleating activity (Welch and Mullins, 2002). However, we recently showed that the Arp2 subunit must be phosphorylated for the complex to be activated by NPFs (LeClaire et al., 2008; Narayanan et al., 2011; Choi et al., 2013). The necessary role for Arp2 phosphorylation for nucleation activity suggests that the Arp2/3 complex is a coincidence detector requiring both binding to NPFs and phosphorylation of Arp2 for increased activity. Coincidence detection is emerging as a common regulatory mechanism for many actin-binding proteins, including NPFs (Rohatgi et al., 2001; Rivera et al., 2009) and cofilin (Frantz et al., 2008; Magalhaes et al., 2011). In support of coincidence regulation of the Arp2/3 complex, we show that expression of a phosphorylation-defective Arp2 mutant complexes with endogenous Arp2/3 complex subunits and dominantly suppresses increased actin filament assembly and membrane protrusion in response to EGF.

Despite the confirmed importance of phosphorylated Arp2 in the nucleation activity of the Arp2/3 complex, kinases phosphorylating Arp2 have not been identified. We report that the Nck-interacting kinase (NIK), a Ste20/MAP4K4 serine/threonine kinase, phosphorylates Arp2 and primes the Arp2/3 complex for activation by NPFs. NIK and its orthologues have a conserved role in regulating actin cytoskeleton-dependent cell processes. NIK activity is necessary for mesoderm migration (Xue et al., 2001) and for epithelial cell membrane protrusion (Baumgartner et al., 2006) and invasion (Wright et al., 2003). The NIK orthologue misshapen in Drosophila melanogaster functions in determining epithelial polarity, neuronal targeting, and cell invasion (Su et al., 1997; Cobreros-Reguera et al., 2010), and the orthologue MIG-15 in Caenorhabditis elegans controls axonal navigation (Poinat et al., 2002) and neuroblast

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migration (Chapman et al., 2008). However, a substrate for NIK or an orthologue that directly regulates actin filament assembly has not been identified. We show that NIK activity is necessary for increased Arp2 phosphorylation and membrane protrusion in response to EGF and also that NIK directly binds the Arp2/3 complex. These findings identify a new regulator for increasing Arp2/3 complex activity and suggest that NIK phosphorylation of Arp2 is a mechanism linking growth factor signaling to actin filament assembly.

Results and discussion

Arp2-T237/238A-Y202A dominantly suppresses EGF-increased actin filament assembly and membrane protrusion

Previous studies to reveal cell processes regulated by Arp2/3 complex have mostly used RNA interference or genetic depletion to eliminate expression of individual subunits (Di Nardo et al., 2005; Nicholson-Dykstra and Higgs, 2008; Suraneni et al., 2012; Wu et al., 2012). However, a limitation of these approaches is that expression of other subunits is abolished because subunit stability depends on an assembled complex. Small molecule inhibitors of Arp2/3 complex activity (Nolen et al., 2009) retain an intact complex but disrupt cortical localization of the complex (Yang et al., 2012). We asked whether these limitations could be resolved by heterologous expression of a mutant nonphosphorylatable Arp2 to dominantly inhibit endogenous Arp2/3 complex activity.

To test this prediction, we stably expressed Arp2 wild type (WT) and a T237/238A-Y202A mutant (TTY/A), tagged at the C terminus with tandem HA and V5 epitopes, in MTLn3 rat carcinoma cells. We used the triple phospho-site Arp2 mutant because we previously showed that phosphorylation of either Thr237/238 or Tyr202 functions as an OR gate for NPF-stimulated Arp2/3 complex activity (LeClaire et al., 2008; Choi et al., 2013). Immunoblotting cell lysates with antibodies to V5 showed similar expression levels of recombinant Arp2-WT and -TTY/A, and neither construct changed the abundance of endogenous ARPC1, ARPC2, or Arp2 (Fig. 1 A). ARPC1 coprecipitated with HA-immune complexes of Arp2-WT and -TTY/A (Fig. 1 B), which suggests that recombinant Arp2 assembles with endogenous Arp2/3 complex subunits.

In MTLn3 cells maintained in growth medium, HA immunolabeling was seen at the distal margin of cells expressing Arp2-WT and -TTY/A but was not in vector control cells, although all cells showed a nonspecific cytoplasmic signal (Fig. 1 C). Co-labeling for endogenous ARPC1 showed localization at the cell distal margin and overlap with HA labeling of both Arp2-WT and -TTY/A in merged images (Fig. 1 C), which indicates that localization of the Arp2/3 complex is not disrupted by expression of Arp2-TTY/A.

Expression of Arp2-TTY/A dominantly suppressed increased actin filament assembly with EGF. In MTLn3 cells, EGF induces a marked increase in F-actin (Chan et al., 1998; Yang et al., 2012), which we confirmed by measuring fluorescence of fixed cells labeled with rhodamine phalloidin. In WT cells, a time-dependent increase in F-actin with EGF was maximal at 5 min (Fig. 1 D). At all time points with EGF, F-actin abundance was not different with expression of Arp2-WT compared with WT cells but was significantly less with expression of Arp2-TTY/A and with latrunculin B (1 µM), which binds and sequesters G-actin (Fig. 1 D). In the absence of EGF, the mean F-actin abundance was not different with Arp2-TTY/A or latrunculin compared with WT cells.

Compared with WT cells and Arp2-WT, expression of Arp2-TTY/A decreased the extent and rate of membrane protrusions with EGF. In growth medium, there were no obvious morphological differences between WT cells and cells expressing Arp2-WT or Arp2-TTY/A, as indicated in Fig. 1 C. However, marked differences were seen with Arp2-TTY/A in quiescent cells, as indicated in cells expressing the actin filament reporter Lifeact-GFP (Riedl et al., 2008). Images captured by time-lapse microscopy showed that quiescent WT and Arp2-WT cells had marked cortical actin bundles and a ruffled plasma membrane. With EGF, the cortical actin bundle remained mostly unchanged but lamellipodia rapidly extended (Fig. 2 A). Quiescent cells expressing Arp2-TTY/A mostly lacked a distinct cortical actin bundle and had fewer and smaller ruffles but an increase in filopodia containing actin filaments in linear arrays (Fig. 2 A). With EGF, extension of lamellipodia was markedly reduced but there was little change in the number or length of filopodia. We also saw this stellate phenotype with filopodia in MTLn3 cells treated with CK666 (Fig. 2 A), and, as we previously reported, in Drosophila S2 cells with RNAi knockdown of Arp2 and expression of Arp2-T237/238A-Y202A (LeClaire et al., 2008). Additionally, the stellate phenotype is seen in fibroblasts lacking ARPC2 (Wu et al., 2012) or ARPC3 (Suraneni et al., 2012) subunits. Hence, inhibiting the Arp2/3 complex may unmask actin assemblies regulated by other nucleators such as formins, which can generate contractile actin filaments. Consistent with this prediction, cells with Arp2-TTY/A and CK666 were smaller and appeared more contracted. Additional analysis showed that an increase in the area of WT and Arp2-WT cells with EGF was significantly less with Arp2-TTY/A and CK666 (Fig. 2 B). Compared with WT cells, the rate of membrane protrusion was significantly faster, with Arp2-WT at 2 and 3 min and significantly slower with Arp2-TTTY/A at 3, 4, and 5 min (Fig. 2 C). It is unclear why the phenotype of cells expressing Arp2-TTY/A is different with quiescence compared with being maintained in growth medium but may reflect an unresolved role for Arp2 phosphorylation. Although cells expressing Arp2-TTY/A had a slower proliferation rate than WT and Arp2-WT cells (Fig. S1), they were viable, which is in contrast to the lethality of Dicyostelium discoideum cells lacking this subunit (Zaki et al., 2007). The ability of Arp2-TTY/A to assemble with endogenous Arp2/3 complex subunits and suppress EGF-induced actin filament assembly and membrane protrusion without disrupting localization of the complex suggests a new strategy to determine functions of the Arp2/3 complex in diverse cell processes.

The Arp2/3 complex is an NIK substrate

To identify kinases that phosphorylate Arp2, we used in vitro kinase assays with Arp2/3 complex purified from Acanthamoeba
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castellani and dephosphorylated with Antarctic phosphatase (AP). We tested a subset of kinases previously shown to associate with the Arp2/3 complex or regulate actin or membrane dynamics, including p21-activated kinase (PAK), the Src tyrosine kinase (Src), focal adhesion kinase (FAK), and NIK. Neither PAK1, a Ste20 family member like NIK, nor FAK phosphorylated any Arp2/3 complex subunit, although they did phosphorylate the generic substrate myelin basic protein (unpublished data). Src phosphorylated Arp3 and Arp2 subunits (Fig. S2 A), but as described below it had no effect on nucleating activity. In contrast, NIK phosphorylated Arp3, Arp2, and ARPC2 subunits of the A. castellanii Arp2/3 complex and recombinant human Arp2/3 complex generated by baculovirus expression (Fig. 3 A). NIK also phosphorylated the ARPC5 subunit of human but not A. castellanii Arp2/3 complex. Phosphorylation of subunits was not detected in the absence of NIK or with a mutant kinase-inactive

Figure 1. Arp2-T237/238A-Y202A coprecipitates with endogenous Arp2/3 complex subunits and dominantly suppresses EGF-increased actin filament assembly. (A) Immunoblot of lysates from WT MTLn3 cells (WT) and MTLn3 cells stably expressing recombinant Arp2 WT (Arp2-WT) and mutant T237/238A-Y202A (Arp2-TTY/A) tagged with VA and HA epitopes show that expression of recombinant Arp2 does not change the abundance of endogenous ARPC1, ARPC2, or Arp2. (B) Immunoblot of proteins in anti-HA immune complexes shows coprecipitation of endogenous ARPC1 with recombinant Arp2 WT and mutant TTY/A. (C) Immunolabeling with anti-HA and anti-ARPC1 antibodies indicates colocalization at the distal margin of plasma membrane protrusions. Bar, 5 µm. (D) Total F-actin in MTLn3 cells in the absence and presence of EGF for the indicated times shows a similar time-dependent increase in WT cells and cells expressing Arp2-WT that is significantly attenuated in cells expressing Arp2-TTY/A or WT cells treated with latrunculin. Data are means ± SEM (error bars) of triplicate determinations in four separate cell preparations. *, P < 0.05; **, P < 0.01; ***, P < 0.001 compared with WT at the same time point (Student’s t test).
NIK (NIK-D152N) that lacks ATP binding but retains substrate binding (Baumgartner et al., 2006; Fig. 3 A).

To test whether NIK, a Ser/Thr kinase, phosphorylates Arp2-Thr237/238, we used as a substrate recombinant Arp2/3 (rArp2/3) complex containing Arp2-WT or -TTY/A, generated by baculovirus expression as we described previously (Narayanan et al., 2011). Recombinant Arp2-TTY/A had no effect on the stoichiometric assembly of rArp2/3 complex subunits (Fig. 3 B, gel). Active NIK but not NIK-D152N phosphorylated Arp3, Arp2, and ARPC2 subunits of WT rArp2/3 complex (Fig. 3 B). With mutant Arp2-T237/238A-Y202A, NIK phosphorylation of Arp2 was abolished but phosphorylation of Arp3 and ARPC2 was unchanged compared with WT. Kinetic analysis revealed a $K_m$ of $37.7 \pm 8 \mu M$.  

Figure 2.  
Expression of recombinant Arp2-TTY/A inhibits membrane protrusion with EGF.  
(A) Time-lapse images of MTLn3 cells expressing Lifeact-GFP in the absence (0 min) and presence (5 min) of EGF show differences in morphology and changes in plasma membrane protrusions for WT, Arp2-WT, Arp2-TTY/A, and WT cells treated with the Arp2/3 inhibitor CK666. Cells expressing Arp2-TTY/A or treated with CK666 had prominent filopodia in the absence of EGF that did not increase in size or abundance with EGF. Also shown are overlapped tracings of the cell periphery in the absence (red and green outlines) and presence of EGF (yellow and blue outlines) that were used to quantify cell area, as well as high-magnification images of the cell periphery at 5 min with EGF (shown on the right, taken from the indicated cyan boxes). Bars: (left) 5 µm; (right) 2 µm. (B) WT and Arp2-WT cells had increased areas after 5 min with EGF that was significantly less with Arp2-TTY/A and WT cells treated with the Arp2/3 complex inhibitor CK666. Data are from four separate cell preparations, with each filled circle representing a single cell (for WT, $n = 39$; for Arp2-WT, $n = 34$; for Arp2-TTY, $n = 36$; for CK666, $n = 14$). (C) The rate of membrane protrusion was significantly faster with Arp2-WT at 2 and 3 min and significantly slower with Arp2-TTY/A at 3, 4, and 5 min compared with WT cells. *, $P < 0.05$; **, $P < 0.01$; ***, $P < 0.001$ compared with WT (Student’s unpaired $t$ test).
NIK phosphorylates Arp2/3 complex

NIK-phosphorylated Arp2/3 complex has increased nucleating activity

Figure 3. NIK phosphorylates and binds the Arp2/3 complex. (A) In vitro, WT NIK but not kinase-inactive NIK-D152N phosphorylates Arp3, Arp2, and ARPC1 subunits of the Arp2/3 complex purified from A. castellanii and from baculovirus expression of human recombinant Arp2/3 complex (rArp2/3). (B) NIK phosphorylation of Arp2 but not Arp3 and ARPC3 subunits is not seen with rArp2/3 complex containing a mutant Arp2-T237/238A-Y202A. (C) Arp2/3 complex coupled to CH-Sepharose and incubated with recombinant NIK 1–340, NIK 1–500, and NIK 1–703 shows that the NIK region of amino acids 340–500 binds the Arp2/3 complex. (D) Schematic of NIK showing the kinase domain and proposed Arp2/3 complex binding domain, which contains abundant acidic (red) and basic (blue) residues. (E) Arp2 coprecipitates in Myc immune complexes of MTLn3 cells transiently expressing Myc-tagged NIK-D152N by not with Myc-tagged WT NIK or empty adenovirus (Ad).
NIK activity is necessary for EGF-increased Arp2 phosphorylation and membrane protrusion

Arp2 phosphorylation increases in MTLn3 mammary carcinoma cells treated with EGF (LeClaire et al., 2008) and in D. discoideum cells treated with cAMP (Choi et al., 2013); however, a kinase mediating either effect has not been identified. To test a possible role for NIK, we used metabolically labeled MTLn3 cells stably expressing shRNA for NIK or a nontargeting (NT) shRNA, confirming the specificity of decreased NIK expression by immunoblotting (Fig. 5A).

Incorporation of $[^{32}P]$orthophosphate into Arp2/3 complex subunits was determined in immune complexes obtained using ARPC1 subunit antibodies. Quiescent control and NT cells had a low basal level of phosphorylated Arp2 that was markedly increased with EGF by 60 s (Fig. 5A). A second phosphorylated polypeptide

Figure 4. NIK restores the actin nucleation activity of the Arp2/3 complex. (A) Pyrene actin assays show that WT NIK but not kinase-inactive NIK-D152N restores activity of alkaline phosphatase–treated Arp2/3 complex with but not without the VCA domain of Scar. (B) The NIK kinase domain [NIK, 1–288] lacking Arp2/3 complex binding sites activates Arp2/3 complex to ~30% of the rate of Arp2/3 complex phosphorylated by full-length NIK. The NIK Arp2/3 binding domain [NIK, 340–500] lacking the kinase domain has no effect on Arp2/3 complex activity. (C) Actin–nucleation activity of Arp2/3 complex containing T237/238A-Y202A Arp2 is not restored with NIK. (D) Src does not restore activity of dephosphorylated Arp2/3 complex. The data shown for pyrene actin traces are from a single representative experiment out of three repeats.

the Arp2/3 complex binding site increased Arp2/3 activity to 30% of untreated Arp2/3 complex, which suggests that NIK-increased nucleating activity is in part dependent on binding the Arp2/3 complex (Fig. 4 B). However, NIK 340–500 had no effect on the nucleating activity of untreated or AP-treated Arp2/3 complex (Fig. 4 B), which indicates that the binding domain without the kinase domain is not sufficient to activate the complex. Additionally, rArp2/3 with Arp2-TTY/A was inactive in the presence of Scar-VCA both with and without NIK (Fig. 4 C), further confirming that the Arp2 phosphorylation sites T237/238-Y202 are necessary for NIK-increased Arp2/3 complex nucleating activity. Although Src phosphorylated several Arp2/3 complex subunits (Fig. S1 A), it did not restore the nucleation activity of the AP-Arp2/3 complex (Fig. 4 D). The functional significance of Src-increased phosphorylation and whether Src phosphorylates Arp2-Y202 remain to be determined.

NIK activity is necessary for EGF-increased Arp2 phosphorylation and membrane protrusion

Arp2 phosphorylation increases in MTLn3 mammary carcinoma cells treated with EGF (LeClaire et al., 2008) and in D. discoideum cells treated with cAMP (Choi et al., 2013); however, a kinase mediating either effect has not been identified. To test a possible role for NIK, we used metabolically labeled MTLn3 cells stably expressing shRNA for NIK or a nontargeting (NT) shRNA, confirming the specificity of decreased NIK expression by immunoblotting (Fig. 5A). Incorporation of $[^{32}P]$orthophosphate into Arp2/3 complex subunits was determined in immune complexes obtained using ARPC1 subunit antibodies. Quiescent control and NT cells had a low basal level of phosphorylated Arp2 that was markedly increased with EGF by 60 s (Fig. 5A). A second phosphorylated polypeptide
Consistent with NIK increasing phosphorylation and activity of the Arp2/3 complex in vitro, the EGF-induced increase in MTLn3 cell area seen in controls was significantly attenuated with expression of NIK shRNA (Fig. 5, D and C; and Fig. S3 B). Additionally, the rate of membrane protrusion with EGF was significantly less with NIK shRNA compared with NT shRNA (Fig. 5 F). We previously showed that NIK-dependent membrane protrusion with EGF is in part dependent on NIK phosphorylation of ERM proteins (Baumgartner et al., 2006). The ERM protein ezrin stabilizes membrane blebs.
by tethering cortical actin filaments to the plasma membrane (Charras et al., 2006). Hence, we propose that in response to EGF, NIK activates the Arp2/3 complex to initiate membrane protrusions, which are stabilized by NIK activation of ERM proteins.

Our findings identify NIK as the first kinase shown to both phosphorylate Arp2/3 complex subunits and increase nucleation activity. Of particular interest is that in addition to NIK binding the Arp2/3 complex, it also binds and phosphorylates ERM proteins (Baumgartner et al., 2006) and NHE1 (Yan et al., 2001), which also regulate actin dynamics at the plasma membrane (Denker and Barber, 2002; Patel and Barber, 2005; Frantz et al., 2008; Neisch and Fehon, 2011). As its acronym indicates, NIK binds the receptor tyrosine kinase adaptor protein Nck, and Nck binds to and activates N-WASP (Rohatgi et al., 2001; Rivera et al., 2009). Together, these findings suggest that NIK coordinates actin filament and membrane dynamics in response to activated receptor tyrosine kinases.

Materials and methods

MTLn3 cell culture and infection

MTLn3 rat adenocarcinoma cells (provided by J. Condeelis, Albert Einstein College of Medicine, New York, NY) were maintained in MEM without nucleosides supplemented with 10% FBS at 37°C and 5% CO2. Cells were made quiescent by decreasing FBS medium from 10% to 0.2% for 4 h. For expression of recombinant Arp2, cells were infected with lentivirus expressing WT Arp2 and Arp2-T237/238A-Y202A tagged at the C terminus with tandem HA and V5 epitopes. Expression was selected by resistance to blasticidin and confirmed by immunoblotting and immunofluorescence. NT and NIK shRNA were obtained from SABiosciences with the sequence 5′-CTC-GGGAGGTTGGAAGATGAGTTC-3′ to silence NIK expression. Plasmids were transfected into cells with a 5:2 ratio of Fugene HD (Promega) with a 2 µg shRNA construct. Cells positive for GFP expression were isolated by flow cytometry after 48 h and cultured for an additional 48 h before analysis. The plasmid containing mEGFP-N1-LifeAct was provided by R. Wedlich-Söldner (Max Planck Institute of Biochemistry, Martinsried, Germany). The EGFP-N1-LifeAct sequence was cloned into a pENTR TOPO plasmid (Life Technologies) according to the manufacturer’s instructions (Haynes et al., 2011). For lentiviral transduction, MTLn3 cells were infected in growth medium supplemented with 4 µg/ml Polybrene (Sigma-Aldrich). Infected cells were selected with 15 µg/ml blasticidin (Life Technologies). Adenovirus was prepared by cloning WT NIK and D152N NIK into pAD-TRACK-CMV and recombining with 15 µg/ml blasticidin (Life Technologies). Adenovirus was prepared by infecting 75 T75 tissue culture flasks (Baumgartner et al., 2006).

MTLn3 cells plated in 24-well dishes. Latrunculin B (Life Technologies) was added 1.5 min before EGF treatment. At the indicated times, cells were fixed and stained as described above for immunofluorescence and incubated with rhodamine-phalloidin (1:500; Life Technologies) for 30 min. After washing, cells were incubated for 5 min with Hoechst 33342 (1:10,000; Life Technologies) to label nuclei. Labeling was measured with a SpectraMax M5 plate reader (Molecular Devices), with rhodamine fluorescence normalized to Hoechst signal and data expressed relative to WT cells in the absence of EGF.

Cell proliferation

MTLn3 cells maintained in growth medium were counted 48 h after plating 10 × 104 cells/well in 24-well plates. Doubling time was determined and data are shown for individual wells from four separate cell plating preparations. Significant differences compared with WT MTLn3 cells were determined with an unpaired t test.

Video microscopy

MTLn3 cells plated in MatTek 35-mm glass-bottom dishes (MatTek Corporation) were incubated using the microscope system described above in a climate-controlled chamber at 37°C and 5% CO2. Changes in cell area were measured as described previously (El-Sibai et al., 2008), with the following modifications to quantify changes in cell size after stimulation with growth factors. Images of cells were captured at 20-s intervals, beginning 2 min before adding 5 nM EGF and ending 10 min after adding EGF. Image analysis was performed in NIS Elements software (Nikon), including time-dependent changes in cell size and lamellipodium, designated as the region from the cell body to the distal margin of the plasma membrane. Fold increase in cell [CA]2 area was calculated by the equation 

\[
\Delta \text{Area} = \frac{\text{Area}_{\text{EGF}} - \text{Area}_{\text{WT}}}{\text{Area}_{\text{WT}}}
\]

where Area is the area of the lamellipodium measured 5 min after EGF stimulation and \( \text{Area}_{\text{WT}} \) is the area of the lamellipodium measured before EGF stimulation. Approximately 30 cells were measured for each condition.

Immunoblotting, immunolabeling, and immunoprecipitation

Immunoblotting was performed by lysing cells in ice-cold Nonidet P-40 lysis buffer, collecting postnuclear supernatants, and separating 40 µg of protein by SDS-PAGE. Proteins were transferred onto PVDF membranes, blocked in 5% BSA, and probed with primary antibodies for 60 min. We used antibodies for a mouse monoclonal V5 (1:5,000; Life Technologies), mouse monoclonal HA (1:1,000; ABAM), mouse monoclonal tubulin (1:2,000; Sigma-Aldrich), rabbit monoclonal ARPC2 (1:1,000; EMD Millipore), rabbit polyclonal ArpC1 (1:5,000, generated to C terminus of ArpC1) by OpenBiosystems; Thermo Fisher Scientific), and rabbit polyclonal ZC1-3a for NIK (1:3,000; Baumgartner et al., 2006). After washing, membranes were incubated with HRP-coupled secondary antibodies (1:5,000; Jackson ImmunoResearch Laboratories) for 1 h at room temperature, and immunoreactivity was visualized using enhanced chemiluminescence (Thermo Fisher Scientific) with reactions imaged using a Chemidoc XRS (BioRad Laboratories).

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Actin polymerization

Cellular Factin in the absence and presence of EGF was determined in MTLn3 cells plated in 24-well dishes. Latrunculin B (Life Technologies) was added 1.5 min before EGF treatment. At the indicated times, cells were fixed and stained as described above for immunofluorescence and incubated with rhodamine-phalloidin (1:500; Life Technologies) for 30 min. After washing, cells were incubated for 5 min with Hoechst 33342 (1:10,000; Life Technologies) to label nuclei. Labeling was measured with a SpectraMax M5 plate reader (Molecular Devices), with rhodamine fluorescence normalized to Hoechst signal and data expressed relative to WT cells in the absence of EGF.

In vitro actin polymerization was measured using 4 µM monomeric nonmuscle actin containing 5% phosvitin and actin in KMEI (50 mM KCl, 1 mM MgCl2, 1 mM EGTA, and 0.1 mM imidazole, pH 7.0; LeClaire et al. 2008). Polymerization was determined in the absence or presence of 5 nM Arp2/3 complex, 500 nM D152N NIK, or mutant NIK. Pyrene actin was excited in a spectrophotometer (RF-5301PC; Shimadzu) at 365 nm, and fluorescence was measured at 407 nm at 1-s intervals. Purified Arp2/3 complex was dephosphorylated by treatment with 1 U AP (New England Biolabs, Inc.) in Tri-Hpf buffer (50 mM Tris, pH 8.0, 1 mM MgCl2, and 0.1 mM ZnCl2) for 1.5 h at 30°C.

Cell proliferation

MTLn3 cells maintained in growth medium were counted 48 h after plating 10 × 104 cells/well in 24-well plates. Doubling time was determined and data are shown for individual wells from four separate cell plating preparations. Significant differences compared with WT MTLn3 cells were determined with an unpaired t test.

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where Area is the area of the lamellipodium measured 5 min after EGF stimulation and \( \text{Area}_{\text{WT}} \) is the area of the lamellipodium measured before EGF stimulation. Approximately 30 cells were measured for each condition.
In vitro kinase and binding assays

Kinase reactions were performed in kinase reaction buffer (25 mM Hapes, pH 7.5, 1 mM DTT, 10 mM MgCl2, 2 mM MnCl2, 50 mM nonradioactive ATP, and 5 μCi [32P]ATP) for 20 min at 30°C. Reactions contained 200 ng NIK generated by baculovirus/insect cell expression, as described previously [Baumgartner et al., 2006]. In brief, the NIK gene (NM_00125220) was ligated into pFASTBac and recombined into a bacmid genome in DH10Bac cells (Life Technologies). Bacmid encoding NIK was isolated and transfected into Sf21 cells, and supernatants containing baculovirus were collected. Viral stocks were used to infect Sf21 cells in suspension. Recombinant NIK was purified by nickel affinity chromatography. Native Arp2/3 complex purified from A. castellani was purified (provided by D. Mullins, University of California, San Francisco, San Francisco, CA). WT Arp2/3 complex was generated by baculovirus/insect cell expression as described previously [Gourrier et al., 2001], or rArp2/3 containing mutant rArp2T237/238A-Y202A was generated as described previously [Narayanan et al., 2011]. In brief, Sf21 cells were infected with baculoviruses containing cDNA encoding Arp2/3 complex subunits. Infected cells were grown in suspension for 48°C for 48 h. Recombinant Arp2/3 complexes were purified on Talon affinity resin (Takara Bio Inc.). Purified Arp2/3 complex was passed over a Superdex 200 FPLC column (GE Healthcare) to isolate fully assembled complexes. A range of Arp2/3 concentrations from 1 nM to 500 nM per reaction were performed to determine kinase kinetics. Reactions were stopped by the addition of Laemmli buffer, proteins were separated by SDS/PAGE, and gels were dried and exposed to autoradiography.

Binding assays were performed in binding buffer (50 mM Tris, pH 7.4, 150 mM NaCl, 1 mM DTT, 100 μM ATP, 1% BSA, and 5% glycerol) containing rArp2/3 complex coupled to CH-Sepharose (GE Healthcare) and the indicated NIK truncations. Truncated NIK proteins were generated by in vitro expression using a cell-free expression kit (Life Technologies) with Express-Tag EXPRESS Protein Labeling Mix (PerkinElmer) containing [35S]methionine. Arp2/3-coupled Sepharose was incubated with the labeled protein for 30 min at 25°C. Sepharose was pelleted by centrifugation, washed with binding buffer, and analyzed by SDS-PAGE. Gels were dried and visualized to be detected by autoradiography.

The dissociation constant (Kd) of NIK 340–500 for Arp2/3 complex was determined by incubating glutathione-bound GST-NIK 340–500 with progressive dilutions of Arp2/3 complex in Arp2/3 binding buffer (2 mM Tris, pH 8.0, 0.2 mM ATP, 0.1 mM CaCl2, and 0.5 mM DTT) from 2,000 ng/ml to 25 ng/ml for 30 min at 37°C. Sepharose-bound protein complexes were centrifuged at 3,000 g, resuspended in Laemmli buffer, and separated by SDS-PAGE. Gels were stained in Coomassie brilliant blue, imaged on an Odyssey Li-Cor scanner, and quantified using ImageJ. Protein binding curves were generated using GraphPad Prism.

Online supplemental material

Fig. S1 shows that the expression of Arp2/3-3TY slows the growth rate of MTLn3 cells. Fig. S2 shows Arp2/3 complex phosphorylation and binding. Fig. S3 shows that NIK regulates the Arp2/3 actin nucleation rate and binding. Fig. S4 shows Arp2/3 complex phosphorylation and binding. Protein binding curves were generated using GraphPad Prism.
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