Cortactin phosphorylation regulates cell invasion through a pH-dependent pathway

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Invadopodia are invasive protrusions with proteolytic activity uniquely found in tumor cells. Cortactin phosphorylation is a key step during invadopodia maturation, regulating Nck1 binding and cofilin activity. The precise mechanism of cortactin-dependent cofilin regulation and the roles of this pathway in invadopodia maturation and cell invasion are not fully understood. We provide evidence that cortactin–cofilin binding is regulated by local pH changes at invadopodia that are mediated by the sodium–hydrogen exchanger NHE1.

Introduction

Complications arising from cancer invasion and metastasis are the major cause of death in cancer patients. To escape from the primary tumor site, cancer cells must migrate through the ECM and intravasate through the basement membrane underlying blood vessels (Yamaguchi and Condeelis, 2007; Gimona et al., 2008). These complex and regulated processes are associated with the formation of specialized actin-rich membrane structures that degrade the ECM called invadopodia (Chen, 1989; Gimona et al., 2008). Several actin regulatory proteins, including N-WASP, cortactin, Arp2/3, and cofilin, are associated with invadopodium formation and function (Wang et al., 2007; Ayala et al., 2008), and each of these genes is up-regulated selectively in invasive breast carcinoma cells (Wang et al., 2004). Cortactin promotes invadopodium formation and maturation in many cancer cells (Artym et al., 2006; Ayala et al., 2008; Oser et al., 2009) and potentiates breast cancer metastasis in animal models (Li et al., 2001). Evidence suggests that invadopodia exhibit discrete stages of maturation, including the assembly of a cortactin-rich invadopodial precursor, cortactin phosphorylation and generation of barbed ends (actin polymerization), cortactin dephosphorylation (stabilization), and ECM degradation (Oser et al., 2009).

Cortactin phosphorylation is a key regulatory step of invadopodium maturation. Tyrosine phosphorylation of cortactin by a kinase cascade involving the Src and Arg nonreceptor tyrosine kinases (Tehrani et al., 2007; Mader et al., 2011) promotes recruitment of the Nck1 adaptor protein, N-WASP, and cofilin (DesMarais et al., 2009; Oser et al., 2009), leading to an increased Arp2/3 complex–mediated actin polymerization at invadopodia (Uruno et al., 2001; Weaver et al., 2001; Oser et al., 2010). Recent work has shown that phosphorylation of tyrosines 421 and 466, but not 482, is essential for both Nck1 binding and for barbed end formation at invadopodia (Oser et al., 2010). Together, these results suggest that cortactin-dependent regulation of N-WASP, Nck1, and cofilin is essential for actin polymerization in invadopodia. Cortactin is also known to regulate leading edge persistence (Bryce et al., 2005), matrix metalloproteinase (MMP) secretion (Clark and Weaver, 2008), and matrix degradation, contributing to cellular invasiveness.

Furthermore, cortactin tyrosine phosphorylation mediates the recruitment of NHE1 to the invadopodium compartment, where it locally increases the pH to cause the release of cofilin from cortactin. We show that this mechanism involving cortactin phosphorylation, local pH increase, and cofilin activation regulates the dynamic cycles of invadopodium protrusion and retraction and is essential for cell invasion in 3D. Together, these findings identify a novel pH-dependent regulation of cell invasion.

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Abbreviations used in this paper: CCD, charge-coupled device; EIPA, (N-ethyl-N-isopropyl)amiloride; FBE, free barbed ends; FRET, fluorescence resonance energy transfer; IP, immunoprecipitate; KD, knockdown; MMP, matrix metalloproteinase; pH, intracellular pH; SE-FRET, sensitized emission FRET; TEM, transmission electron microscopy; WT, wild type.

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though cofilin KD cells exhibit only minor defects in matrix invasion through an 8-µm-pore Transwell (Fig. S1 A), even these findings, cofilin KD cells (Fig. S2 C) exhibited impaired invasion independently of degradation (Fig. 1 C). In agreement with and cofilin regulate invadopodial elongation in MDA-MB-231 cells coated with Matrigel (Fig. 1 C). These data suggest that cortactin phosphorylation and cofilin are required for invasion of actin available to initiate actin polymerization (DesMarais et al., 2004, 2005). In so doing, cofilin regulates cell migration behavior, cell directionality (Ghosh et al., 2004; Sidani et al., 2007), and, ultimately, cell invasion (Wang et al., 2007; Oser and Condeelis, 2009; van Rheenen et al., 2009). Cofilin activity is regulated via diverse mechanisms including phosphorylation/dephosphorylation (Moriyama et al., 1996), PIP2 binding (van Rheenen et al., 2007; Leyman et al., 2009), and local pH changes mediated by NHE1 (Frantz et al., 2008). NHE1 is a ubiquitously expressed transmembrane protein that regulates intracellular pH (pHi) by exchanging extracellular sodium for intracellular protons (Kemp et al., 2008).

We provide evidence here that cortactin phosphorylation promotes recruitment of NHE1 to regulate pH in invadopodia. We demonstrate that increased pH disrupts cortactin binding to cofilin, thus releasing cortactin’s inhibitory grip on cofilin. Finally, we demonstrate that these pH changes regulate invadopodial maturation and invadopodium-mediated invasion in 3D. Our findings reveal a novel mechanism by which cortactin phosphorylation and NHE1 control pH as a key step in cancer cell invasion.

Results

Cortactin phosphorylation and cofilin are required for invasion

We used a previously described 1-µm Transwell assay (Schoumacher et al., 2010) to characterize the role of cortactin phosphorylation in MDA-MB-231 invasion. Protrusive structures that crossed to the underside of the membrane (>12 µm; Fig. 1 A) were quantified as the relative number of invading structures (Fig. 1 B). Cells expressing wild-type (WT) cortactin extended cortactin-rich invasive structures that crossed to underside of the Matrigel-coated 1-µm-pore Transwell (Fig. 1 A). However, cells expressing only cortactin mutant lacking all three major tyrosine phosphorylation sites (3YF) or in which cofilin was knocked down (KD) exhibited severe defects in invasion (Fig. 1, A and B). Trans-membrane protrusion was inhibited by the protease inhibitor GM6001, which indicates a requirement for MMP activity in the process (Fig. 1 B).

Cortactin 3YF and cofilin KD cells still exhibited severe defects in trans-membrane protrusion when surfaces were uncoated with Matrigel (Fig. 1 C). These data suggest that cortactin and cofilin regulate invadopodial elongation in MDA-MB-231 independently of degradation (Fig. 1 C). In agreement with these findings, cofilin KD cells (Fig. S2 C) exhibited impaired invasion through an 8-µm-pore Transwell (Fig. S1 A), even though cofilin KD cells exhibit only minor defects in matrix degradation (Fig. S1 B). Together, these results show that both cortactin phosphorylation and cofilin are required for invadopodia elongation and invasion.

Cortactin-cofilin interaction decreases during invadopodium maturation

Given their requirement for invadopodia elongation and invasion, we next determined whether cortactin–cofilin interactions change during invadopodium maturation. Cofilin is known to be the primary generator of actin FBE in invadopodia, whereas cortactin phosphorylation is a key regulator of cofilin activity (Oser et al., 2009). To test whether cortactin phosphorylation status changes during invadopodial maturation, we analyzed the presence of phosphorylated cofilin (P-cofilin) in both mature invadopodia and invadopodium precursors (Fig. S1 C). 95% of degrading invadopodia (mature invadopodia) have a clear accumulation of P-cofilin, whereas ~25% of invadopodial precursors without degradation activity show an accumulation of P-cofilin (Fig. S1, C and D; Bowden et al., 2006). We found that EGF stimulation leads to a decrease in fluorescence resonance energy transfer (FRET) between cofilin and cortactin that coincides with the cortactin phosphorylation peak after EGF stimulation in MDA-MB-231 cells (Fig. S1 E), as we have described previously in MTLn3 cells (Oser et al., 2009). Using the degradation-based invadopodia maturation analysis, we also measured cofilin–cortactin interactions in precursor versus mature invadopodia using acceptor photobleaching FRET (AP-FRET) measurements. P-cortactin–rich mature invadopodia show a reduction in cofilin–cortactin FRET compared with invadopodium precursors (Fig. 1, D and E), which is in agreement with previous observations (Oser et al., 2009). These results suggest that a dynamic loss of cortactin–cofilin interaction correlates with invadopodium maturation. Elevated cortactin–cofilin FRET is observed in both precursors and mature invadopodia of 3YF cortactin–expressing cells (Fig. S2, A and B), which indicates that cortactin phosphorylation is associated with decreased cortactin–cofilin interaction and that both events correlate with invadopodial maturation.

Cofilin–cortactin interaction at invadopodia is regulated by local pH changes

These results suggest that cortactin phosphorylation disrupts its interaction with cofilin. Thus, we conducted in vitro binding experiments to explore whether the cortactin phosphorylation was sufficient to disrupt cortactin–cofilin binding. As shown in Fig. 2 (A and B), cofilin binds efficiently to P-cofilin, with a dissociation constant ($K_d = 3.0 \pm 0.8 \mu M$) similar to what was reported for the 3YF cortactin (2.8 ± 0.34 µM; Oser et al., 2009). These data indicate that cortactin phosphorylation does not disrupt its binding to cofilin directly, but must influence cofilin binding via an indirect mechanism.

Frantz et al. (2008) demonstrated that cofilin activity is inhibited by phosphoinositide binding in a mechanism regulated by local pH changes. Cofilin undergoes pH-dependent structural changes, decreasing phosphoinositide binding at a higher pH. We tested whether pH regulates cofilin binding to cortactin. Cofilin binds to cortactin with a $K_d$ of $1.4 \pm 0.4 \mu M$ at pH 6.8 (Fig. 2, C and D). However, cortactin–cofilin binding...
Figure 1. Cortactin tyrosine phosphorylation and cofilin are required for invadopodia elongation. (A) Representative 3D reconstructions of the 1-µm Transwell experiment. Data are based on three or more independent experiments. (B) The number of protrusive structures crossing the bottom of the membrane (>12 µm) normalized to the cell area on top of the membrane is shown (>10 fields per group, n = 4; ***, P < 0.0001). (C) Quantification of cell protrusion through a 1-µm Transwell without Matrigel. The GM 6001 inhibitor was used to show that cell protrusion was independent of degradation (>10 fields/group, n = 4; ***, P < 0.0001). Cortactin tyrosine phosphorylation regulates Cofilin interaction in invadopodia. (D) Representative cofilin–cortactin FRET efficiency images of cells expressing either WT or 3YF cortactin. The white circle indicates the bleached spot. The top right insets show close-ups of the original images (indicated by the boxed regions; n = 5). (E) Quantification of cofilin–cortactin FRET/donor at mature invadopodia and precursors in MDA-MB-231 cell lines expressing WT or 3YF cortactin (endogenous cortactin KD). Light gray bars represent FRET on mature invadopodia and dark gray bars represent FRET on invadopodia precursors (*, P < 0.002; n = 5, >30 invadopodia/group). Error bars indicate SEM.

affinity is reduced at pH 7.2 (Fig. 2, E and F). We could not reach saturation of binding at the tested concentrations in buffer at pH 7.2, and thus we were only able to place a lower limit on the $K_d$ at pH 7.2 (limit $K_d$ ≥ 4.1 ± 1.3 µM). The pH range used in these experiments is in agreement with the previously reported pH-dependent binding of cofilin to PIP$_2$ (decreased binding at pH 7.5; Bailly and Jones, 2003; Frantz et al., 2008). This binding is specific, as several proteins do not bind cortactin under the conditions used here (Lapatina et al., 2009; Oser et al., 2010). Moreover, these binding affinities are within the
Figure 2. Cortactin-cofilin binding in vitro is not dependent on cortactin phosphorylation, but is pH sensitive. (A and B) Coomassie-stained gels (A) and quantification of the binding signal of cofilin to phosphorylated cortactin (B; P-cortactin; $K_d = 3.0 \pm 0.8$ µM); data points to $K_d$ calculation = 16. (C and D) Coomassie-stained gels (C) and quantification of the binding signal of cofilin to cofilin at pH 6.8 (D). $K_d = 1.4 \pm 0.4$ µM; data points to $K_d$ calculation = 22. (E and F) Coomassie-stained gels (E) and quantification of the binding signal of cortactin to cofilin at pH 7.2 (F). $K_d > 4.1 \pm 1.3$ µM;

G Mature Invadopodia

H Cortactin Intensity and pH changes at invadopodia

I Invadopodia precursors

J Cortactin Intensity and pH changes at invadopodia precursors
Together, these data suggest that cortactin–cofilin interaction in invadopodia is not directly affected by cortactin phosphorylation status but is pH dependent over the same range that pH varies in the invadopodium.

**Barbed end formation in invadopodia is regulated by pH**

Considering the observations that the cofilin–cortactin interaction is regulated by pH in vitro and that invadopodium maturation is accompanied by an increase in pH, we hypothesized that local pH changes regulate cofilin release from cortactin and the resulting increase in cofilin-dependent FBE in invadopodia.

To test this hypothesis, we measured FBE when the pHi was stabilized to 6.8 or 7.2 using the ionophore nigericin and standard high K+ media (Chaillet and Boron, 1985; Kaplan and Boron, 1994). Upon EGF stimulation, control cells that were not treated with nigericin and maintained at pH 7.2 responded with an increase in FBE at invadopodia (Fig. 3, A and B; expected range for interactions involving actin binding proteins (Pollard and Cooper, 1986; Cao et al., 2006).

The clear impact of pH changes on cofilin–cortactin binding led us to investigate pH fluctuations at invadopodia as a potential mechanism regulating cofilin–cortactin binding and cofilin activity. To characterize the local pH changes at invadopodial precursors (nondegrading) and mature invadopodia (degrading), MDA-MB-231 cells were plated on a fluorescent matrix and loaded with the pH indicator SNARF 5F. In mature invadopodia (Fig. 2 G), the peak of cortactin fluorescence (Fig. 2 H, red line) correlated with an increase in pH as determined by the SNARF 5F pH biosensor (Fig. 2 F, black line). On average, the pH in the invadopodium core was 7.15 ± 0.01 compared with a pH of 6.91 ± 0.03 in the surrounding cytoplasm. In contrast, analysis in invadopodium precursors showed that there is no significant pH change at the cortactin core of nondegrading precursors compared with the surrounding cytoplasm (pH range 6.97–7.05; Fig. 2, I and J).

Together, these data suggest that cortactin–cofilin interaction in invadopodia is not directly affected by cortactin phosphorylation status but is pH dependent over the same range that pH varies in the invadopodium.
Figure 4. Cortactin phosphorylation regulates NHE1 recruitment to the invadopodia. (A) Representative images of endogenous NHE1 and cortactin colocalization. (B) Representative image of MDA-MB-231 cells transiently expressing the WT NHE1-HA and WT cortactin. (C and D) The colocalization experiments were repeated in cells expressing KRA-NHE1–WT cortactin (C) and WT NHE1–3YF cortactin (D). Insets show enlarged views of the boxed regions. (E) Quantification of the cortactin–NHE1 colocalization. Results are based on the analysis of 15 cells/group; *, P < 0.01. (F) Quantification...
mean pH of control cells was 7.08 ± 0.015). Cells that were equilibrated with a lower pH (6.8) failed to increase FBE in response to EGF, whereas the equilibration of the pH to 7.2 alone induced a significant increase in FBE formation (Fig. 3, A and B). We show here that the pH increase from 6.8 to 7.2 observed during invadopodium maturation (Fig. 2, E and F) results in a significant increase in actin FBE. Moreover, NHE1KD cells failed to generate FBE after EGF stimulation (Fig. 3 C). These results support our hypothesis that invadopodium maturation and the generation of FBE within are regulated by pH.

Tyrosine phosphorylation regulates cortactin-NHE1 interaction at invadopodia

The sodium hydrogen exchanger NHE1 is a key regulator of several pH-dependent cellular processes (Stock and Schwab, 2009). Recent studies suggest that NHE1 localizes to invadopodia (Busco et al., 2010) and has an important role in cancer invasion (Reshkin et al., 2000; Bourguignon et al., 2004; Busco et al., 2010). We found that endogenous NHE1 (Fig. 4 A) or ectopically expressed HA-tagged WT NHE1 (Fig. 4 B) colocalizes with cortactin in mature invadopodia in MDA-MB-231 cells. To confirm these results, we conducted a colocalization analysis using HA tag NHE1 constructs. An ERM binding–deficient NHE1 mutant (KRA-NHE1) with uniform membrane localization (Denker et al., 2000) also showed a reduced colocalization with cortactin (Fig. 4, C and E). We examined whether cortactin phosphorylation regulates recruitment of NHE1 to the invadopodia by measuring the colocalization of the 3YF cortactin mutant and NHE1. Cells expressing 3YF cortactin showed reduced cortactin–NHE1 colocalization at invadopodia (Fig. 4, D and E). Similarly, WT NHE1 showed increased colocalization with cortactin in mature invadopodia compared with invadopodia precursors (Fig. 4 F). We performed coimmunoprecipitation experiments to measure interactions between NHE1 and cortactin. NHE1 WT coimmunoprecipitated with WT cortactin, however; the non-phosphorylatable cortactin 3YF mutant exhibited a decreased interaction with NHE1 compared with WT cortactin (Fig. 4, G and H). These results suggest that cortactin phosphorylation increases NHE1 recruitment to invadopodia.

To test if NHE1 regulates cofilin–cortactin interaction at invadopodia, we analyzed the cortactin–cofilin AP FRET both in NHE1KD cells and also in rescue experiments with WT NHE1 and KRA NHE1. Considering that we observed a consistently higher FRET in invadopodia precursors compared with mature invadopodia, we used the precursor/mature invadopodia FRET ratio to report the results. As shown in Fig. 4 I, in control cells there was a twofold higher precursor/mature invadopodia FRET. The precursors/mature invadopodia FRET ratio was decreased in NHE1KD cells and fully rescued by the expression of WT NHE1 but not KRA NHE1 (Fig. S2 F).

In agreement with these findings, NHE1 KD cells showed a reduced pHi, which was rescued by the expression of WT NHE1 (Fig. S3 A). Furthermore, cells expressing 3YF cortactin exhibited reduced average invadopodial pH, as NHE1 is not recruited to the invadopodia (Fig. S3 B) and NHE1 inhibition using 5-(N-ethyl-N-isopropyl)amiloride (EIPA) decreases pHi (Fig. S3 C). Together, these results show that NHE1 regulates the pH-dependent interaction between cortactin and cofilin in vivo.

**Cortactin phosphorylation regulates the dynamic oscillatory behavior of invadopodia**

We propose that cortactin phosphorylation regulates NHE1 recruitment to invadopodia, resulting in release and activation of cofilin, leading to free barbed end generation within invadopodia. To quantify the dynamic recruitment and accumulation of cortactin and cofilin, we performed fluctuation analysis on time-lapse images of the cofilin and cortactin content in invadopodia. Both WT cortactin and cofilin exhibited periodic fluctuations in total intensity at invadopodia of steady-state MDA-MB-231 cells (Fig. 5, A and B). This oscillatory pattern was not caused by microscope focal changes because the analysis is done using 3D acquisition followed by the analysis of the maximum projection stack (Fig. 5 A, bottom inset). Correlation analysis showed that both WT cortactin and cofilin displayed an oscillation pattern of ~800 s, as shown by the second peak on the curve (Fig. 5, C and D, arrows). 3YF cortactin expression inhibited the well-defined oscillatory pattern observed in WT cortactin-expressing cells (Fig. 5, E–G). In accordance with our model, we hypothesized that the oscillations in cortactin and cofilin at invadopodia were associated with pH fluctuations. We found that significant pH oscillations are observed at invadopodia (Fig. 5 H, bottom left) compared with the control areas in the cytoplasm (Fig. 5, H and I, bottom right). The coefficient of pH variation (standard deviation/mean, Fig. 5 I) shows that the pH oscillates more at invadopodia compared with other cytoplasmic areas of the same cell.

To test the role of NHE1 in the dynamic accumulation of cortactin and cofilin at the invadopodium, we treated MDA-MB-231 cells with the NHE1 inhibitor EIPA, which blocks NHE1-mediated pH regulation (Fig. S3 C), and analyzed cortactin and cofilin oscillations at invadopodia. Inhibition of NHE1 disrupts the stability of the invadopodium core, inducing its disassembly (Fig. S4, A and B). After addition of EIPA, the spatially stable fluorescent signals that characterize the invadopodial core disappear (Fig. S4, A and B). Together, these results define invadopodia as a very dynamic structure,
Figure 5. Invadopodia display a dynamic oscillatory behavior in 2D. (A) Representative images of MDA-MB-231 cells expressing WT cortactin-RFP and cofilin-GFP. The cells were imaged for 4 h and the fluctuation in the relative fluorescence intensity of cofilin and cortactin was analyzed and plotted. The insets show the boxed region. (B) Representative traces of WT cortactin and cofilin pixel intensity representative of 1 invadopodia from 12 analyzed. (C and D) The fluctuations in WT cortactin and cofilin fluorescence intensity were analyzed using autocorrelation (see Material and methods). The autocorrelation for WT cortactin is shown in C and the corresponding cofilin-GFP autocorrelation is shown in D. Data are based on the analysis of 12 invadopodia in three independent experiments. (E–G) The experiment was repeated in cells expressing 3YF cortactin-RFP and cofilin-GFP. (E) Representative traces of cofilin and 3YF cortactin pixel intensity (1/10). Autocorrelation analysis of the fluorescence intensity of 3YF cortactin is shown in F and the corresponding cofilin-GFP signal is shown in G. Red traces represent the standard error of the mean. (H) Representative images of cells expressing WT cortactin and the SNARF-5F pH indicator. Inner white circles represent the invadopodial core and the outer circles represent the invadopodium periphery. (I) Quantification of the coefficient of pH variation. n = 18 invadopodia in three experiments; ***, P < 0.001. Error bars indicate SEM.
characterized by cycles of cofilin and cortactin accumulation and pH increase. Furthermore, they provide evidence that cortactin phosphorylation determines the oscillatory behavior of invadopodia.

To corroborate these dynamic studies, we also analyzed the recovery of both cortactin and cofilin to invadopodia after photobleaching. As seen in Fig. S4 C, the expression of 3YF cortactin decreases the mobile fraction of both cofilin and cortactin, indicating that the invadopodium is less dynamic. These results indicate that cortactin phosphorylation regulates the dynamic recruitment of cofilin to invadopodia.

Coractin phosphorylation, NHE1, and cofilin are essential for invadopodium elongation

The results described here identify a new pH-dependent mechanism that regulates invadopodia maturation and actin FBE generation. To analyze this mechanism in a more physiologically relevant environment, we investigated the role of cortactin tyrosine phosphorylation, cofilin, and NHE1 in cells moving in three dimensions, using a 3D invasion assay. This assay consisted of embedding cells in a thick Geltrex matrix and analyzing the transition of an early round cell to a polarized invasive cell over 72 h. After 24 h of 3D culture, MDA-MB-231 cells formed small, well-defined protrusive structures enriched in cortactin, Tks5, and MT1-MMPs (Fig. S5, A and B; and Video 1). By using the degradation marker DQ collagen IV or DQ-BSA-Green, we observed that these cortactin and Tks5-rich protrusive structures at the cell front had degradation activity and were therefore considered invadopodia (Fig. 6 A). Multiple well-defined invadopodia were observed in cells expressing WT cortactin 48 h after matrix embedding (Fig. 6 B), and these invadopodia elongated into long (>12 µm) structures, eventually leading to a polarized cell phenotype (~72 h). A high-magnification transmission electron microscopy (TEM) image of an invadopodium (Fig. 6 B, lower right inset) shows a well-defined actin-rich protrusion surrounded by areas of matrix degradation (Fig. 6 B). Compared with WT cells, 3YF cortactin cells were rounder and had shorter invadopodia (Fig. 6, C and D). Closer examination revealed that these cells had less organized invadopodia, which was also confirmed by high-magnification TEM imaging (Fig. 6, C [bottom right inset] and D). Cofilin KD and NHE1 KD cells also showed shorter invadopodia as seen in the 3YF cortactin–expressing cells (not depicted). Thus, cortactin phosphorylation, cofilin, NHE1, and matrix proteolysis are all required for the formation of long (>12 µm) invadopodia (Fig. 6 D).

We also tested the role of NHE1 in invadopodial elongation. Similarly to what we observed in the 3YF cortactin and cofilin KD cells (Fig. 1, B and C), NHE1 KD cells failed to elongate through a 1-µm Transwell filter in the absence of Matrigel (Fig. 6 E). Cell elongation was rescued by the expression of WT NHE1 but not the KRA NHE1 mutant (Fig. 6 E). Cell invasion in the presence of Matrigel was also inhibited in NHE1 KD cells (Fig. 6 F). These data suggest that the primary role of NHE1 at invadopodia is not associated with matrix degradation but rather linked to the regulation of focused actin polymerization.

Coractin phosphorylation is required for the dynamic protrusion of invadopodia in 3D

Using the 3D invasion assay, we observed that proteolysis-dependent cell invasion is accomplished by a dynamic invadopodia protrusion/retraction cycle at the cell front (Fig. 7 A and Video 2). Interestingly, we observed a population of cells that actively degrade the matrix (Fig. 7 A) and a second population that would migrate through predegraded tracks (Fig. S5 C and Video 3; Takino et al., 2006). The mean migration speed of the degrading cells was 0.16 ± 0.02 µm/min, whereas the speed of cells migrating through the degraded areas was 0.85 ± 0.2 µm/min. Also, the protrusive structures in these cells did not resemble the structures observed on the leading edge of cells that were carving the tunnels (Fig. 7 A). Approximately 24 h after seeding, MDA-MB-231 cells extended long protrusions with a clear accumulation of cortactin at the tip (Fig. 7 A and Video 2). This invasive structure collapsed and elongated again, creating a well-defined cycle (Fig. 7 B). A few cortactin/Tks5-positive spots are also observed away from the leading invasive protrusion, and we propose that these are invadopodia precursors. Kymograph images of WT cortactin, 3YF cortactin, and NHE1KD cells (Fig. 7 C) showed that WT cells have a dynamic recycling of the invadopodia that are coupled to cell movements, whereas 3YF cortactin–expressing cells and NHE1KD cells have more stable structures with fewer cycling events. A similar phenotype was also observed for the cofilin KD cells (unpublished data). The kymograph images were analyzed in detail using the autocorrelation function of the intensity profiles for WT and 3YF cortactin. Fig. 7 D shows that WT cortactin cells had multiple peaks in the autocorrelation curve, indicating a recurrence of protrusion cycles. The correlation analysis of the 3YF cortactin cells (Fig. 7 E) and NHE1KD cells (Fig. S5 D) indicated slower and/or less defined protrusion events compared with WT. These results show that cortactin phosphorylation regulates cell invasion in 3D through the regulation of a dynamic invadopodia protrusion/retraction cycling at the cell front. Together, we provide evidence that cortactin phosphorylation regulates the dynamic cycles of cofilin and cortactin accumulation and pH increase at invadopodia, leading to an increased FBE generation. We propose that this mechanism explains the dynamic protrusion/retraction cycling of invadopodia at the cell front in 3D.

Discussion

Both cortactin phosphorylation and cofilin activity are critical for invadopodium maturation and cancer cell invasion (Oser et al., 2009, 2010; Mader et al., 2011), but how cortactin phosphorylation regulates cofilin activity and especially how this mechanism regulates cell invasion in 3D was unclear. Here, we demonstrate that cortactin phosphorylation during invadopodia maturation recruits NHE1, thereby inducing a local increase in pH. The higher pH releases cofilin from cortactin and generates FBE required for invadopodium elongation. We also demonstrate that invadopodia are very dynamic structures, displaying a well-defined oscillation in pH and cortactin and cofilin content.
Figure 6. Cofilin and cortactin phosphorylation are required for invadopodia protrusion. (A) Representative images of MDA-MB-231 cells expressing WT cortactin-RFP incubated in Geltrex + DQ Collagen IV (1:10) or Geltrex + BSA-Green 488 degradation marker. Images are representative of >50 cells analyzed in three independent experiments. (B) Representative images of WT cortactin cells embedded in the 3D matrix. The cells were imaged by confocal microscopy between 48 and 72 h after plating. The image is a 3D reconstruction of a representative field. Arrows and insets represent cells with invadopodia. The bottom right panel shows a representative TEM image of an invadopodium in a WT cortactin cell. (C) Representative images of 3YF cortactin cells embedded in the 3D matrix. The cells expressing 3YF cortactin-GFP were incubated as described (see Materials and methods). Arrows and bottom left insets represent high-magnification images of cells with invadopodia. The bottom right panel shows a representative TEM image of an invadopodium in a 3YF cortactin cell.
Figure 7. **Invadopodia are dynamic structures in 3D.** (A) Representative images of MDA-MB-231 cells expressing WT cortactin-RFP incubated with Gel-trex and BSA-Green 488 for 48 h. The cells were imaged for 8 h and the images represent snapshots of a time-lapse movie. The degradation marker channel was processed to highlight the degradation areas. (B) The cartoon shown represents the dynamics of invadopodia protrusion observed in these cells. Multiple cycles of invadopodia elongation and retraction are observed during cell invasion. Results are based on the analysis of 15 cells in three experiments. (C) Representative kymograph of a WT cortactin cell (left), a 3YF cortactin cell (middle), and a NHE1KD cell (right) migrating in 3D. The plots underneath represent the variations in pixel intensity of the protrusions over time. (D) Quantification of the WT cortactin protrusion autocorrelation. Cells expressing RFP-tagged WT cortactin were mixed with 10 mg/ml Geltrex and incubated for 24 h. Cells were imaged by between 24 and 48 h after plating. (E) Quantification of the 3YF cortactin protrusion autocorrelation. Cells expressing RFP-tagged 3YF cortactin were mixed with Geltrex and incubated for 24 h. Cells were imaged between 24 and 48 h after plating. n = 9 cells (WT) and 10 cells (3YF) in three independent experiments. Red traces represent the standard error of the mean.
We provide evidence that these events occur in a physiologically relevant 3D matrix system to control invasive migration. Our work demonstrates that cortactin phosphorylation–cofilin–NHE1 pathway regulates invadopodium elongation and dynamics at the cell front in 3D.

**Cortactin phosphorylation regulates cofilin activity through a pH-dependent pathway**

Cofilin is known to be the key generator of FBE in invadopodia, and its activation is known to be associated with cortactin phosphorylation at invadopodia. In particular, cortactin phosphorylation is correlated with dissociation of the cofilin–cortactin interaction, allowing cofilin to generate FBE (Oser and Condeelis, 2009; Oser et al., 2009). We have analyzed here the specific mechanism linking cortactin phosphorylation to the regulation of cofilin activity and found that the binding and release dynamics of cofilin from cortactin is not directly regulated by phosphorylation of cortactin. Rather, cortactin phosphorylation regulates cofilin binding indirectly, by controlling an NHE1-dependent pH increase that releases cofilin from cortactin’s inhibitory grip in invadopodia (Fig. 8).

The proposed inhibitory binding model is also supported by the following observations. First, we have found that cortacin concentration in MDA-MB-231 cells is in the micromolar range (1 µM, see Fig. S2 G). Additionally, the invadopodium compartment is unique and it contains on average approximately four times more cortactin than the mean cell levels, whereas cofilin shows only an approximately twofold increase over mean cell levels (Fig. S2 H). Although cofilin is found at a concentration of ~6 µM in breast cancer cells (Chan et al., 2000), the majority of it is bound to actin or PIP2. Only an estimated 4% of cofilin is free in the cell (Tania et al., 2011), which suggests that there is enough cortactin available to sequester free cofilin in invadopodia. Altogether, these observations support our model of cofilin sequestration by cortactin at invadopodia.

Substantial evidence supports the finding that local pH increase is an evolutionarily conserved mechanism that regulates cytoskeleton dynamics and cell migration (King et al., 1994; Denker and Barber, 2002; Lovy-Wheeler et al., 2006). We describe here a novel pathway, where invasive cancer cells use a fine tuned pH regulation at invadopodia to regulate free barbed end generation. Cofilin is known to be a pH biosensor and its pH sensitivity is mediated by the protonation of the His133 (Frantz et al., 2008). Although we have strong evidence supporting an electrostatic interaction between cofilin and cortactin, the exact determinants that mediate this interaction and its disruption by elevated pH are unclear.

**NHE1 is a key player in cancer invasion**

NHE1 catalyzes an exchange of extracellular Na⁺ for intracellular H⁺, and its activity increases in response to growth factors (Putney et al., 2002; Frantz et al., 2007) and integrin engagement (Tominaga and Barber, 1998). NHE1 is necessary for directed migration in different cells, including fibroblasts (Denker and Barber, 2002) and leukocytes. Increasing evidence suggests that cancer cell migration and invasion rely on multiple pH- and extracellular pH (pHe)-dependent pathways (Stüwe et al., 2007; Stock and Schwab, 2009) that regulate cell adhesion, cell volume, cytoskeleton dynamics, and ECM degradation (Stock et al., 2008; Stock and Schwab, 2009; Busco et al., 2010). More specifically, NHE1 is known to affect breast cancer invasion (Reshkin et al., 2000; Cardone et al., 2005; Cardone et al., 2007), melanoma migration (Stock et al., 2005), and ECM degradation through the regulation of pHe (Busco et al., 2010). We have used KD + rescue experiments to show that NHE1 regulates cortactin–cofilin interaction at invadopodia. NHE1 regulates the alkalization of the invadopodium, inducing cofilin activity and increasing actin filament free barbed end generation at mature invadopodia.

Upon receptor activation, cortactin is phosphorylated downstream of Src and Arg kinases (Mader et al., 2011). We propose that cortactin phosphorylation recruits NHE1 to the invadopodium, but the specific mechanism behind this step is unknown. The anchoring of NHE1 through the distal charged domain (550–564) has been demonstrated to determine the location of NHE1 (Denker et al., 2000). How this is linked to cortactin phosphorylation and then induction of the local pH changes that will drive cofilin off cortactin is unknown but may involve binding of both NHE1 and cortactin to F-actin (Weed et al., 2000; Weed and Parsons, 2001). Alternatively, NHE1 might interact with the cortactin-Nck1 invadopodium core through its binding to Nck-interacting kinase (NIK; Yan et al., 2001). Multiple mechanisms are known to regulate NHE1 exchanger activity, including phosphorylation by Rho-associated protein kinase (ROCK; Tominaga and Barber, 1998), p38 MAPK (Khaled et al., 2001), and NIK (Yan et al., 2001). Further analysis of the cortactin–NHE1 interaction is needed to elucidate this novel mechanism. The activation of NHE1 exchanger activity is also regulated downstream of growth factor receptors (Meima et al., 2009), which could represent a common EGF-dependent cortactin phosphorylation pathway. Recent work suggests that NHE1 localizes to invadopodia, where it acidifies the extracellular space, increasing ECM degradation (Busco et al., 2010). In addition to the proposed acidification of the peri-invadopodial extracellular space, we provide evidence here that the primary role of NHE1 may be to regulate barbed end formation and invadopodial elongation.

A model for invadopodial dynamics in 3D nonfibrillar matrices

The characterization of invadopodia in 3D models has been the interest of many research groups (Linder, 2007; Sabeh et al., 2009; Wolf et al., 2009; Li et al., 2010; Tolde et al., 2010). Invadopodial structure in 3D can differ significantly, depending on the cell type or the matrix model used. Using fibrillar collagen matrices or dermis slices, Wolf et al. (2009) have described cell invasion, proposing that invadopodia correspond to the lateral spikes associated with the sides but not the front of pseudopod extensions. However, this remains a speculation because no defined invadopodial markers were used in these studies. Similarly to what we describe here, Packard et al. (2009) showed that matrix degradation is localized at the leading front of migrating tumor cells. It is possible that as cells go through different ECM compositions, the mode of motility could change,
making the organization of the cell front different in each specific situation (Wolf et al., 2003; Van Goethem et al., 2010).

In our investigation, we analyzed the invasion of MDA-MB-231 cells through a reconstituted basement membrane using a real 3D cell embedding technique and time-lapse imaging. We identified invadopodia using well-accepted invadopodium characteristics (Blouw et al., 2008; Crimaldi et al., 2009; Oser et al., 2010). First, invadopodia are actin-rich invasive protrusions. Second, the invadopodium is associated with matrix degradation. Finally, the scaffolding proteins cortactin and Tks5 are enriched in invadopodia. Considering these definitions and experimental conditions, we observed that the invadopodium compartment that is normally seen as a defined ventral surface structure in 2D forms a complex and dynamic invasive structure at the cell front in 3D. Also, a careful analysis of our movies indicates that invadopodial extension determines
the direction of invasion. Although the crosstalk between directional sensing and the invadopodium has been suggested (DesManais et al., 2009), the physical separation between these structures forced by 2D culturing methods always imposed a barrier to the detailed analysis of this mechanism.

Invadopodia are defined as actin-rich protrusions that localize matrix-degrading activity to cell–substratum contact points, converging sites of signaling, proteolytic, adhesive, cytoskeletal, and membrane trafficking pathways (Weaver, 2006). A comprehensive description of invadopodia in 3D awaits further work, but we propose that the distinct lamellipodium and invadopodium compartments observed in 2D are merged into one invasive structure at the cell front in 3D that preserves the characteristics of an invadopodium. Recently, Poincloux et al. (2011) found that MDA-MB-231 cells invade Matrigel based on cell rear contractility and a round phenotype (Poincloux et al., 2011). We observed a population of MDA-MB-231 cells that sustained a round phenotype but the majority of the cells in our experiments extended invadopodia and acquired a polarized phenotype after 48 h of 3D culturing. We believe that having the cells embedded in the matrix and the longer incubation times/analysis (>48 h) explain the differences observed in cell invasion.

Using our 3D model, we were able to observe the transition from a round cell to an invasive, polarized cell. This process is based on the formation of one leading edge invasive structure that goes through multiple cycles of protrusion and retraction. Overall, the defined frequency of cycles is dependent on cortactin phosphorylation as indicated by the random elongation frequency observed in the 3YF cortactin mutant. This evidence is the first to suggest that invadopodia are very dynamic structures in 3D, in agreement with what has been proposed in other 2D models (Oser et al., 2009; Artym et al., 2011). Our results suggest that cortactin phosphorylation, NHE1, and cofilin are not only required for the focal elongation of invadopodia but also the well-defined protrusion/retraction cycling observed in 3D. Interestingly, we also show that both cofilin and cortactin go through cycles of accumulation at invadopodia in 2D, resembling the frequency observed in 3D. The expression of the cortactin phosphorylation mutant 3YF induced a disorganized fluctuation of both cortactin and cofilin at invadopodia. Together, these results support that cortactin phosphorylation/dephosphorylation at invadopodia is the molecular mechanism behind the rhythmic protrusion/retraction necessary for cell invasion.

Materials and methods

Cell line

MDA-MB-231 cells were cultured in 10% FBS/DME with antibiotics. Where indicated, cells were starved in 0.5% FBS/0.8% BSA/DME for 12–16 h and then stimulated with 2.5 μM EGF (Invitrogen). All live cell experiments were imaged at 37°C in L15 media.

DNA constructs, RNAi, and transfection

KRA-NHE1-HA and WT-NHE1-HA were a gift from D. Barber (University of California, San Francisco, San Francisco, CA; Denker et al., 2000). KRA-NHE1-HA mutant contains K-A and R-A mutations in the distal charged domain, regulating ERM binding (Denker et al., 2000) and ion exchange activity (Aharonovitz et al., 2000). Mouse WT cortactin and cortactin 3YF mutants were cloned as described previously (Iapetska et al., 2009). Stable cell lines were generated using the retroviral expression vector pLXSN (BD; Oser et al., 2010). Where indicated, cofilin-GFP, Tks5-GFP (Sylli et al., 2009), and WT or 3YF cortactin-RFP (Oser et al., 2009) constructs were transiently transfected into MDA-MB-231 cells using the Nucleofection kit V protocol (Lonza) 24 h before each experiment. Control nonsilencing siRNA was from Qiagen. For long time-lapse imaging and 3D experiments that required long incubations, cells with stable expression of WT or 3YF cortactin-GFP and stable KD of endogenous cortactin were used. To knock down the endogenous cortactin in these cells, the cortactin intron sequence 5′-GGACAAAGTGTTAAGACG3′ was subcloned into psuper-Retro [Oligogene] and transfected into Apatropic Phoenix packaging cell lines. Retroviral supers were used to infect the GFP-cortactin WT or mutant cell lines, which were then selected with 10 μg/ml puromycin. Human-specific cortactin siGenome Smart Pool (M010508-00-0005) and SiGenome Smart Pool NHE1 specific (S055277-0005) were obtained from Thermo Fisher Scientific. 106 MDA-MB-231 cells were transfected with 2 μM siRNA using the Nucleofection kit V protocol 72 h before each experiment. For rescue experiments, the NHE1 constructs were transfected back into NHE1 KD cells 24 h before the experiment.

Antibodies and inhibitors

Cortactin (ab33333) and rabbit NHE1 (ab67314) were obtained from Abcam; mouse anti-NHE1 (MAB3140) was obtained from Millipore. Chicken NHE1-A was obtained from Alpha Diagnostics. Arp2 (sc-H-84) and Tks5 (sc-30122) were obtained from Santa Cruz Biotechnology, Inc.; pY421 was obtained from Sigma-Aldrich; and anti-biotin FITC was obtained from Jackson ImmunoResearch Laboratories. Anti-cofilin (AE774) is a custom-made antibody (Yamaguchi et al., 2005). Alexa Fluor-conjugated secondary antibodies were from Invitrogen. For immunoprecipitation, cortactin 4F11 was obtained from Millipore. For immunoblots, cortactin (ab33333) was obtained from Abcam; anti-β-actin (AC15) and pY421 were obtained from Sigma-Aldrich. Chicken NHE1 antibody was from D. Barber. All Western blot analyses were performed using the secondary antibodies from Li-Cor (mouse-680 and rabbit-800). EIPA was from Sigma-Aldrich and GM6001 was from Enzo Life Sciences.

3D cell imaging

25,000 cells were resuspended in cold 10 mg/ml Geltrex growth factor reduced basement membrane mix (Invitrogen). The mixture (80 μl) was plated in a 10-mm Mattek dish well for 20 min at 37°C. The dish was carefully filled with 2 ml 10% FBS/DME and incubated for 24–48 h. Before imaging, the media was replaced with 10% FBS/L15. Matrix degradation was analyzed in real time using the DQ BSA Green self-quenched BODIPY probe (Busco et al., 2010). In brief, the BSA green probe was mixed with Geltrex (30 μg/ml final concentration) before the cells were resuspended. Where indicated, DG Collagen IV (Invitrogen) was added to the Geltrex at a 1:10 dilution before the cells were resuspended. In motility invasion experiments, the cells were imaged for 8 h using a microscope (DeltaVision; Applied Precision) and an EM charge-coupled device (CCD) camera (Photometrics). A 40-µm z stack (1-µm step) image was acquired every 5 min [60x, NA 1.42 oil objective lens]. For quantification of invadopodia length and 3D reconstructions, the samples were imaged using a SP5 AOPS confocal microscope (Leica; 20x, NA 0.7 multi-immersion objective lens) 48 h after incubation. Mean invadopodia length was calculated using the 3D reconstructions and included invasive cell protrusions associated with degradation in any direction. Images were analyzed using Velocity 5.5 (PerkinElmer). Where indicated, a look up table filter was applied to the BSA-green channel to help visualize the degradation area.

Fluctuation analysis

All fluctuation analysis was done using custom software written in IDL (ITT Visual Information Solutions). Invadopodia fluctuation analysis can be divided into two steps: (1) generating the raw count rate trace from the invadopodium with a tracking algorithm and (2) autocorrelating the count rate trace offline with a multi-tau correlation algorithm (Schätzle, K. 1985. New concepts in correlator design. Institute of Physics Conference Series. Hilger, London. 77:175–184). The tracking algorithm consists of a Gaussian mask localization procedure and a method for assembling particle trajectories, both described previously (Cracker and Orier, 1996; Thompson et al., 2002; Larson et al., 2005). For fluctuation analysis based on the intensity of the invadopodium, the precise measure of integrated intensity in every frame is critical, and the preceding algorithms have been modified for fluctuation measurements in several ways. First, the intensity of the site is determined after a local background correction. The local background is determined by defining a border square with an edge size that is seven times the point spread function width, and the values at this position are fit to a plane of arbitrary orientation using a linear regression. The intensity of January 22, 2018
the site is then determined using a Gaussian mask fit on the background-subtracted spot. Second, the invadopodium trajectory must account for frames where the site disappears completely. If no site is found by the localization algorithm, the tracking procedure goes back to the last frame where a site was found and uses that position as the putative location of the site to calculate the intensity with the Gaussian mask (Thompson et al., 2002). This approach necessitates an approach for calculating intensity that does not depend explicitly on the existence of a visible site. In other words, using the Gaussian mask, one need only specify a position and a width of the point spread function to calculate intensity. Once the invadopodium is visible again, the tracking algorithm resumes based on this new position. This same procedure is used to generate the background trace from the cytosol. For the background trace, an arbitrary position in the cytosol lacking an invadopodium, is chosen as a reference point, and intensity is calculated for each frame in the time stack.

After generation of a count rate trace using the tracking algorithm, the trace is autocorrelated using a multi-tau approach. This approach has been described in detail elsewhere (Schätzl, K. 1985. New concepts in correlator design. Institute of Physics Conference Series. Hilger, London. 77:175–184; Wohland et al., 2001) and is the basis of most commercial autocorrelator cards. In brief, the count rate, which is recorded in linearly spaced, invariantly sized frames (i.e., the intensity of a site as measured from a single frame of the time series), is rebinned to reduce noise at low autocorrelation delays. Rebinning consists of combining adjacent bins to increase the number of counts in the bin, and the bin size increases at longer delays. The bin size is chosen so that the variance is regularly sized samples and determines the signal-to-noise ratio and the resolution of the decay. For invadopodium autocorrelation, we used a doubling interval of four bins, meaning the size of the bin increases from one frame to two frames to four frames to eight frames, etc., at intervals of four times the fundamental imaging frequency. Therefore, if the imaging frequency is 0.125 Hz, the bin size at the 8s delay is half the bin size at the 32s delay. We refer the reader to Fig. 2 in Wohland et al. (2001).

Invadopodia protrusion/retraction analysis was also done using the autocorrelation analysis described. In brief, kymographs of cells invading through a 3D matrix were generated using ImageJ 1.44k. Measurements were made using a line tool crossing the protrusions on the kymograph image, and a pixel intensity profile plot was generated. The plots consisted of multiple peaks in intensity corresponding to the invadopodium protrusions. These plots were used for the autocorrelation analysis.

Cortactin–cofilin FRET experiments
MDA-MB-231 cells stably expressing WT cortactin–GFP or 3YF cortactin–TagRFP in which endogenous cortactin was transiently knocked down using siRNA lacking an invadopodium were plated on Alexa Fluor 405–labeled gelatin for 4 h and fixed with 3.7% PFA. Immunofluorescence was performed using chicken Cofilin 774 + Alexa Fluor 488–goat anti–chicken (donor) and cortactin ab-33333 + Alexa Fluor 555–goat anti–mouse (acceptor). All the AP FRET was performed as described previously (Oser et al., 2010). In brief, invadopodia containing colocalized cortactin and cofilin were bleached (561 s laser power, 10 iterations, bleach time ~8.4s) and the changes in fluorescent signal in the 488 nm channel were measured and analyzed. Images were acquired using a laser scanning microscope (LSM 5 LIVE Duoscan; Carl Zeiss) with a CCD camera using LSM 5 Live Duoscan software. FRET efficiency was calculated as E = 1 – (Donor/Pre/Donor post). All images were corrected for laser fluctuations, sample bleaching, and background. For sensitized emission FRET (S盆地FRET) experiments, MDA-MB-231 cells expressing Cofilin–GFP and WT cortactin–TagRFP were starved overnight as described previously and stimulated with 2.5 μM EGF. The live sensitized emission imaging was imaged on a microscope (IX70; Olympus) with 60× NA 1.4 oil objective lenses and a CCD (Sensicam QE; PCO), and the analysis was performed as described previously (Feige et al., 2005; Oser et al., 2010). In brief, control images spaced intervals were made using a line tool crossing the protrusions on the kymograph image, and a pixel intensity profile plot was generated. The plots consisted of multiple peaks in intensity corresponding to the invadopodium protrusions. These plots were used for the autocorrelation analysis.

In vitro binding
Binding experiments were performed as described previously (Oser et al., 2009). Recombinant cortactin protein was covalently coupled to AminoLink Beads (1 mg/ml gel bed; 16.3 μM; Thermo Fisher Scientific) according to the manufacturer’s instructions. Cofilin was buffer exchanged into binding buffer (50 mM Hepes, pH 6.8, 20 mM NaCl, 0.01% NP-40, and 5% glycerol) and serially diluted 1:3; 20 μl was saved as input sample, and 490 μl of each dilution was added to 50 μl cortactin bead slurry or 50 μl blank bead slurry. The reaction was incubated for 30 min (4°C), the supernatant was removed, and beads were resuspended with binding buffer. Bound protein was recovered with 40 μl Laemmli sample buffer and analyzed on 12% SDS-PAGE gels, stained with Coomassie blue R-250, and scanned using a densitometer (Bio-Rad Laboratories). For the specific pH-binding experiments, cofilin was covalently coupled to the beads and cortactin was added to the binding buffer. Integrated density measurements of the bands and equal background area were made using QuantityOne software (Bio-Rad Laboratories). The Kd was determined as described previously (Oser et al., 2009). Binding curves were generated using the standard one-site specific binding function of GraphPad Prism software (GraphPad Software).

pH measurements
Monitoring of pH was performed using the cell-permeable SNARF-5F pH indicator (Invitrogen) according to the manufacturer’s instructions. In brief, 50,000 cells were plated on a thin Alexa Fluor 405–labeled gelatin matrix 24 h before the experiment. The pH indicator SNARF-5F (Invitrogen) was resuspended in DMEM/pluronic F-127 (Invitrogen) and mixed with the cells at a final concentration of 5 μM for 30 min, 37°C. Cells were washed twice in 10% FBS/1:5 and imaged using a confocal microscope (SPS AOB; Leica). The pH indicator was excited at 514 nm and emission was collected at 640 nm (625–655) and 580 nm (565–595). The 640/480 emission ratio was used to calculate the pH. In situ calibration was performed as described by the manufacturer. In brief, cells were incubated in a High K medium + 10 μM Nigericin (1 μM Nigericin, 105 mM KCl, 20 mM CaCl2, 1 mM MgSO4, 10 mM Hepes, 10 mM glucose, and TRIS for pH adjustment) with specific pH (6.6, 6.8, 7, 7.2, 7.4). All the image analysis was done using Velocity Imaging: Image background corrected and a ratio image of 680 nm emission/580 nm (640/580) emission was generated. The mean cell 640/580 ratio of the calibration samples was used to generate a standard curve for pH calculations. Invadopodia were detected in the cortactin–GFP channel and matrix degradation was detected at using the Alexa Fluor 405–labeled matrix. At the end, a four-channel image was generated (405, matrix; 488, cortactin; 580, pH emission 1; and 640, pH emission 2). For the quantification of invadopodia pH, a linear measurement (~15 pixels in length by 3 pixels wide) crossing the center of the cortactin spot was created for each invadopodia (Velocity measurements tool). The three pixels representing the peak intensity of cortactin were defined as the invadopodium core (pixel size xy = 0.39 μm, z step = 2.0 μm). The peak pH value in the 3 × 3 pixel matrix representing the invadopodium core was used to line up the structures for subsequent analysis. The same analysis was done in irrelevant areas of the cell and returned no changes in pH, confirming the accuracy of this technique. In time-lapse experiments, the ratio of invadopodium core pH and surrounding cytoplasm pH was used to compensate for photobleaching and laser fluctuations.

Colocalization experiments
For detection of endogenous NHE1, cells expressing WT cortactin or 3YF cortactin (100,000) were plated on thin 405-labeled gelatin matrix and fixed after 4 h in 3.7% PFA. Cells were immunostained for cortactin (ab-33333 + Alexa Fluor 555–goat anti–mouse) followed by secondary antibodies (1:300 Alexa Fluor anti–rabbit 488). Images were acquired using an inverted microscope (IX81; Olympus; 60x, NA 1.4 oil objective lens). For colocalization analysis of HA-tagged NHE1 mutants, cells were transfected with the indicated constructs and plated on a gelatin matrix and fixed as described earlier. Samples were imaged using a confocal microscope (SPS AOB; Leica), and colocalization was analyzed as described previously (Oser et al., 2010). In brief, images were processed using ImageJ Spot enhancing filter 2D (3.0 pixel Gaussian filter) and threshold levels were set to select only invadopodia (entropy threshold). The images were then analyzed for colocalization (Mander’s coefficient) using the Jaccop colocalization plug-in for ImageJ.

Barbed end assay
The barbed end assay was performed as described previously (Oser et al., 2009). In brief, MDA-MB-231 cells were starved, stimulated with 2.5 μM EGF, and permeabilized with a permeabilization buffer (20 mM Hepes, pH 7.5, 138 mM KC1, 4 mM MgCl2, 3 mM EGTA, 0.2 mg/ml saponin, 1 mM ATP, and 1% BSA) containing 0.4 M biotin–actin (Cytoskeleton) for 1 min at 37°C. Cells were fixed in 3.7% formaldehyde for 5 min followed by blocking in PBS containing 1% FBS, 1% BSA, and 3 μM phalloidin. The samples were then stained with FITC anti-biotin to visualize barbed ends and Arp2 to identify invadopodium precursors. The actin–biotin intensity at invadopodium precursors was quantified as mean gray value (MGV) at invadopodium precursors minus MGV of the background. The data were normalized to the control condition for each experiment (Oser et al., 2009).
Co-immunoprecipitation

Immunoprecipitation experiments were performed as described previously (Oser et al., 2009). In brief, 106 MDA-MB-231 cells were plated on plastic tissue culture dishes 48 h before the experiment. The samples were stimulated with EGF (2.5 nM, 3 min), washed with cold PBS, and lysed in buffer containing 10 mM Hepes, 50 mM NaCl, 5 mM EGTA, 5 mM MgCl₂, 1% Triton X-100, and phosphatase (NaF and NaVO₃) and protease inhibitors. The lysates were centrifuged at 18,000 g for 10 min at 4°C and precleared with protein A/G agarose beads (sc-2003) for 30 min. The immunoprecipitate (IP) was mixed overnight with 2 µg control antibody 4F11 or 2 µg control IgG1 antibody conjugated to protein A/G agarose beads. The IPs were washed three times in lysis buffer and analyzed by Western blotting. All Western blot analyses were done using Quantity-One (Bio-Rad Laboratories). Images were corrected for background levels and results are expressed as co-IP normalized to the IP optical density for each sample.

Electron microscopy

MDA-MB-231 cells expressing WT or 3YF cortactin-TagRFP where endogenous cortactin was transiently knocked down were plated on Gelrex (10 mg/ml)-coated Mattek dishes. Samples were fixed 24 h after incubation using 2.5% glutaraldehyde in 0.1 M sodium cacodylate buffer, postfixed with 1% osmium tetroxide followed by 1% uranyl acetate then by dehydration through a graded series of ethanol, and embedded in LX112 resin (LADD Research Industries). Ultrathin (80-nm) sections were cut on an ultramicrotome (Ultratome UCT; Reichert), stained with uranyl acetate followed by lead citrate, and viewed on a transmission electron microscope (1200EX; JEOL) at 80 kv. To confirm that the matrix-free regions on the matrix were not artifacts, we compared areas around invadopodia and distant cell–matrix interface areas searching for matrix-free areas. The majority of matrix-free areas were associated with invadopodia. (Fig. S5 E)

Transwell invasion assay

Transwell assays were performed as described previously (Oser et al., 2010). In brief, 8.0 µm Transwell supports (Costar) were coated with 50 µl of reduced growth factor Matrigel (2.5 mg/ml) for 1 h at 37°C. 50 µl of reduced growth factor Matrigel (2.5 mg/ml) for 1 h at 37°C. Transwell assays were performed as described previously (Oser et al., 2005). The membranes were allowed to equilibrate in DME at 37°C for 1 h. 50,000 cells were resuspended in 200 µl 0.5% FBS/DME and plated in the upper chamber. The membranes were imaged using an SP5 microscope FITC-gelatin (1/20 dilution) and applied to the upper surface of the membrane. Similarly, 1 µm Transwell assays were adapted from Schoumacher (2000). First, 1 µm Transwell supports (Costar) were coated with 50 µl of reduced growth factor Matrigel (2.5 mg/ml) for 1 h at 37°C. 50,000 cells were resuspended in 200 µl 0.5% FBS/DME and plated in the upper chamber. The bottom chamber was filled with 1 ml 10% FBS/DMEM. Similarily, 1 µm Transwell assays were adapted from Schoumacher et al. (2010). In brief, 2.5 mg/ml Matrigel was mixed with 0.2 mg/ml FITC-gelatin (1/20 dilution) and applied to the upper surface of the membrane. Membranes were equilibrated as described and 50,000 cells were plated. Where indicated, cells were plated in Transwell membranes without Matrigel coating and allowed to invade for 24 h followed by fixation in 3.7% PFA. Intact membranes were imaged using an SPS microscope (Leica; 3D stacks of the membrane section). Cell invasion in Matrigel-coated membranes was calculated as the number of invadopodia crossing to the underside of the membrane normalized to the cell area on the top of the membrane. Cell elongation on non-Matrigel-coated membranes was calculated using a ratio of cell area in the bottom and cell area on the coated membranes was calculated as the number of invadopodia crossing to the underside of the membrane normalized to the cell area on the top of the membrane. Cell elongation on non-Matrigel-coated membranes was calculated using a ratio of cell area in the bottom and cell area on the top of the membrane to calculate invasive. In all Transwell experiments, the cells were stained with either Alexa Fluor 488 or Alexa Fluor 555 Phalloidin for 30 min before imaging. The MMP inhibitor GM6001 (25 µM) was added as indicated.

FRAP

FRAP curves were acquired on the Zeiss 5 Live Duoscan (Carl Zeiss) with dual bleaching using the 488 nm and 561 nm lasers, and dual fluorescence recovery monitoring in both red (550–615 nm) and green (495–525 nm) channels. Images were acquired with a time structure of 5 prebleach scans and 115 postbleach scans at time intervals of 250 ms, resulting in a total acquisition of ~30 s. The integrated intensity of the invadopod at each time point was determined using a Gaussian mask tracking algorithm (Zhang et al., 2005) to generate the raw FRAP curves. The curves were normalized to prebleach values, and the size of the bleach square was fixed at 10 × 10 pixels. Total cellular fluorescence was also measured to correct for possible cellular photobleaching. For the illumination intensities used to monitor fluorescence recovery over this time scale, we observed minimal photobleaching, obviating the need for additional normalization of the FRAP curve. The resulting normalized curves were fit using a nonlinear least squares fitting routine to the following function:

\[ F(t) = A(1 - e^{-kt}) + C, \]

where \( F(t) \) is the integrated fluorescence of the invadopod at time \( t \); \( k \) is the recovery time constant; \( A \) is the relative fractions of \( k \); and \( C \) is a constant offset to account for the bleach depth. All image processing was done with custom software implemented in IDL and curve fitting was done using GraphPad 5.03 (GraphPad Software).

Correlation coefficients between time constants determined for the same invadopodium but in different channels were calculated according to:

\[ P = \sum \frac{(k_{ji} - \bar{k})(F_{ji} - \bar{F})}{\sigma_k \sigma_F} \]

where \( k \) are the time constants described above \( [j = 1, 2] \) for a particular channel (red, green) and \( i \) is the summation variable that accounts for individual invadopodia. The mean value of the population is denoted by a superscript bar.

Statistical analysis

In experiments containing multiple groups, ANOVA was performed associated to Bonferroni tests. Where indicated, statistical analysis was calculated using the unpaired, two-tailed Student’s t test. Statistical significance was defined as P < 0.05. Error bars represent the SEM.

Online supplemental material

Fig. S1 complements the data shown in Fig. 1, including colocalization analysis, matrix degradation, and SE-FRET analyses. Fig. S2 shows the KD efficiency and expression levels of proteins tested. Fig. S3 shows the effects of NHE1 KD and EIPA inhibitors on invadopodia and cell pH. Fig. S4 shows the effect of EIPA in invadopodia stability and the FRAP analysis of cortactin and coinfit at the invadopodia. Fig. S5 further describes the migration of MDA-MB-231 cells in 3D. Videos 1–3 show time-lapse images of MDA-MB-231 cells migrating in 3D. Online supplemental material is available at http://www.jcb.org/cgi/content/full/jcb.201103045/DC1.

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