S100A4 regulates macrophage invasion by distinct myosin-dependent and myosin-independent mechanisms

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INTRODUCTION
S100A4 is a member of the S100 family of Ca\(^{2+}\)-binding proteins. The S100 proteins, which are found exclusively in vertebrates, are typically symmetric homodimers. Each subunit contains two Ca\(^{2+}\)-binding motifs: a C-terminal canonical EF-hand and an N-terminal pseudo EF-hand that is unique to S100 proteins (Bresnick et al., 2015). Although S100 family members exhibit a high degree of sequence and structural similarity, they are not functionally interchangeable, and distinct S100 proteins participate in a wide range of biological processes, including proliferation, migration and/or invasion, inflammation, and differentiation (Marenholz et al., 2004; Donato et al., 2013). Functional diversity within the S100 family is accomplished through differences in the metal-binding properties of individual family members; their ability to form homodimers, and in some cases, heterodimers; their interactions with distinct protein partners; and their distinct patterns of tissue/cell expression (Heizmann and Cox, 1998; Santamaria-Kisiel et al., 2006; Donato et al., 2013).

S100A4 is highly expressed in fibroblasts (Takenaga et al., 1994) and immune cells, including monocytes and macrophages (Li et al., 2010; Hashimoto et al., 2013), T-cells (Weatherly et al., 2015), and other bone marrow–derived cells (Erlandsson et al., 2013; Scheiter et al., 2013; Bruhn et al., 2014). Furthermore, S100A4 expression in breast, lung, prostate, colorectal, and brain tumor cells has been linked to invasive behavior (Bresnick et al., 2015). A major mechanism of S100A4 action in motile cells involves its regulation of actomyosin-mediated motility. Biochemical studies have shown that S100A4 binds with high affinity to the nonmuscle myosin-IIA heavy chain and mediates myosin-IIA filament depolymerization (Li et al., 2003; Kiss et al., 2012, 2016). Consistent with the biochemical activity of S100A4 in vitro, the loss of S100A4 results in myosin-IIA filament overassembly and a decrease in the migratory capacity of multiple cell types (Dulyaninova and Bresnick, 2013). Importantly, the binding of S100A4 to myosin-IIA is Ca\(^{2+}\) dependent. Thus, S100A4 functions as a calcium sensor whose regulation of the actomyosin cytoskeleton is linked to dynamic changes in intracellular calcium levels.

Our previous studies showed that bone marrow–derived macrophages (BMMs) isolated from mice with a genetic deletion of S100A4 exhibit defects in directional motility and chemotaxis in vitro and reduced thioglycollate-induced recruitment to the peritoneal cavity in vivo (Li et al., 2010). The recruitment of inflammatory monocytes to sites of infection requires exit from the bone marrow and
spleen, entry into the bloodstream, and migration across the endothelium at sites of infection (Shi and Pamer, 2011). Upon migration into infected tissue, monocytes differentiate into dendritic cells and classically activated M1 macrophages, and migrate though the dense extracellular matrix to confront pathogens. The ability of monocytes/macrophages to undergo transendothelial migration and invade target tissues depends on podosomes (Medina et al., 2015). These actin-rich structures perform both adhesive and invasive functions, allowing macrophages to penetrate through endothelial layers and the extracellular matrix (Gimona et al., 2008; Linder and Wiesner, 2015).

Podosomes contain an actin-rich core that is surrounded by a ring structure consisting of proteins associated with cellular adhesions (Linder et al., 2011). Individual podosomes are linked by actomyosin filaments, and sensing of mechanical stress by the myosin-mediated contractile apparatus regulates podosome dynamics (Linder and Wiesner, 2016). Podosomes drive cell invasion by coupling actin-mediated protrusion to microtubule-mediated secretion of matrix metalloproteases (MMPs) (Wiesner et al., 2010). Individual podosomes organize into larger structures called rosettes, whose appearance correlates with the ability to degrade extracellular matrix (Van Goethem et al., 2011; Guiet et al., 2012).

The formation of podosome rosettes is enhanced by calcium influx via specific ion channels (Clark et al., 2006; Siddiqui et al., 2012) and is disrupted by pharmacological inhibition of myosin-II (Collin et al., 2006, 2008). Given that S100A4 is a Ca^{2+}-dependent regulator of myosin-IIA, we examined its role in the formation of matrix-degrading structures in BMMs. Our studies identify S100A4 as an important and novel regulatory component in the organization and degradative function of podosome rosettes in macrophages. Importantly, S100A4 acts through both myosin-IIA-dependent and myosin-IIA-independent mechanisms to regulate rosette size, degradative capacity, and dynamics, such that S100A4 loss leads to a profound defect in macrophage invasion.

RESULTS

S100A4 is required for macrophage invasion and regulates MMP9 secretion

Our previous studies showed that primary BMMs from S100A4^{−/−} mice exhibit defects in colony-stimulating factor 1 (CSF-1)-stimulated chemotactic migration, due in part to myosin-IIA overassembly (Li et al., 2010). In addition, these studies revealed that S100A4 deletion reduces macrophage recruitment in a thioglycollate-induced model of peritonitis. Consistent with our published in vivo recruitment studies, S100A4^{−/−} BMMs exhibited a 44% reduction in Matrigel invasion (Figure 1).

While defects in the directional motility of S100A4^{−/−} BMMs (Li et al., 2010) could account for the observed reductions in invasion, we considered whether additional mechanisms might contribute to this phenotype. In particular, a number of studies have reported that S100A4 knockdown in tumor cells reduces MMP9 mRNA levels (Saleem et al., 2006; Zhang et al., 2011), and genetic deletion of MMP9 inhibits BMM invasion and matrix degradation (Fleetwood et al., 2014). To evaluate whether alterations in MMP9 or other proteases contribute to the reduction in the invasive capacity of S100A4^{−/−} BMMs, we used an angiogenesis antibody array. Conditioned medium derived from S100A4^{−/−} BMMs contained increased levels of MMP8 and MMP9 compared with medium from wild-type BMMs (Figure 2A). These changes did not reflect a general effect on the secretory pathway, as few differences in other secreted proteins were observed (unpublished data). An examination of MMP9 by gelatin zymography also revealed a significantly increased accumu-
S100A4 regulates podosome rosette-mediated gelatin degradation

The finding that S100A4−/− BMMs secreted more MMP9 was surprising, given that these cells exhibited defects in Matrigel invasion. To further explore the mechanisms contributing to impaired invasion by S100A4−/− BMMs, we directly examined the capacity of these cells to degrade matrix. When cells were plated on a fluorescent gelatin substrate, the area of degraded matrix was reduced by 61% in S100A4−/− BMMs compared with wild-type BMMs (Figure 4, A and B). This suggests that MMP9 secretion by S100A4−/− BMMs is not effectively coupled to matrix degradation.

In murine bone marrow–derived macrophages, the formation of podosome rosettes is required for matrix degradation and three-dimensional migration (Cougoule et al., 2010). To determine whether the difference in the matrix degrading activity of S100A4−/− BMMs was due to alterations in podosome rosette formation, we examined the number and size of podosome rosettes. In cells plated overnight on gelatin, there was a 60% reduction in the number of S100A4−/− BMMs that formed podosome rosettes as compared with wild-type BMMs (Figure 5, A and B). In addition, an evaluation of rosette area revealed that the number and size of podosome rosettes in wild-type BMMs was due to alterations in podosome rosette formation, we examined the difference in the matrix degrading activity of S100A4−/− and wild-type BMMs. Data represent the mean ± SEM from five independent blots. Statistical analysis was performed using an unpaired Student’s t test.

When CSF-1 and serum starved wild-type BMMs were plated on gelatin, we observed that 69% of wild-type BMMs had podosome rosettes, whereas only 28% of S100A4−/− BMMs exhibited podosome rosettes (Figure 6, A and B). Stimulation with an equal volume of complete BMM medium (60 ng/ml CSF-1 and 7.5% serum final concentration) induced the rapid disassembly of podosome rosettes in wild-type BMMs, with only 5% of cells exhibiting rosettes 10 min after stimulation. In contrast, S100A4−/− BMMs exhibited a delay in rosette disassembly, with 19 and 7.7% of S100A4−/− BMMs exhibiting podosome rosettes 10 and 20 min after stimulation, respectively (Figure 6, A and B). These data demonstrate that not only do S100A4−/− BMMs have fewer and smaller podosome rosettes, but the kinetics of rosette disassembly is slowed.
Blebbistatin treatment rescues defects in podosome organization and matrix degradation

Our biochemical studies previously showed that myosin-IIA is over-assembled in S100A4−/− BMMs (Li et al., 2010). Consistent with these data, myosin-IIA exhibited a linear punctate pattern in the cortex of S100A4−/− BMMs that was not observed in wild-type BMMs (Figure 7A). Although myosin-IIA is not a component of the podosome core, it localizes to the radial F-actin network that surrounds individual podosomes (Bhuwania et al., 2012), and actomyosin filaments regulate the organization of podosome clusters in dendritic cells (Meddens et al., 2016). An examination of podosome rosettes revealed that myosin-IIA is highly enriched in these structures (Figure 7B). Despite the extensive myosin-IIA filaments detected in the cortex of S100A4−/− BMMs, we did not observe any gross alterations in the localization or organization of myosin-IIA within the small podosome rosettes in these cells.

We have previously shown that low doses of blebbistatin, a small molecule inhibitor that stabilizes the myosin-II motor in a weak actin-binding state (Kovacs et al., 2004; Limouze et al., 2004), can counteract the actomyosin overassembly induced by S100A4 loss (Li et al., 2010). We therefore used blebbistatin as a tool to probe for myosin-IIA–dependent and myosin-IIA–independent mechanisms with respect to MMP9 secretion and matrix degradation in S100A4−/− BMMs. Treatment of S100A4−/− BMMs with the active isomer of blebbistatin (Supplemental Figures 2 and 3). We therefore tested whether the myosin-IIA–independent role for S100A4 in Matrigel invasion could involve α-tubulin K40 acetylation. To determine whether alterations in tubulin acetylation affect the invasive capacity of wild-type BMMs, we treated wild-type BMMs with tubacin, an HDAC6-selective inhibitor. Doses of 2.5–5 µM tubacin increased α-tubulin K40 acetylation in wild-type BMMs to levels observed in S100A4−/− BMMs (Figure 9, A and B). This increase in tubulin acetylation resulted in the stabilization of podosome rosettes to serum/CSF-1–mediated disassembly (Figure 9C), similar to what was seen in S100A4−/− BMMs (Figure 6B), and also correlated with a 30–50% reduction in the invasion of wild-type BMMs (Figure 9D) that was comparable to the decrease in invasion observed for S100A4−/− BMMs (Figure 1). Higher levels of microtubule acetylation (unpublished data) and greater reductions in invasion (Figure 9D) were seen with 25 µM tubacin. These data demonstrate that post-translational modification of microtubules is a critical regulator of BMM invasion, as increases in microtubule acetylation by S100A4 loss or tubacin treatment are sufficient to inhibit invasion.

**DISCUSSION**

In this study, we demonstrate that S100A4, a Ca2+-regulated mediator of myosin-IIA assembly, modulates the matrix-degrading capacity of macrophages via both myosin-IIA–dependent and myosin-IIA–independent mechanisms. Studies in a number of cell types have suggested that calcium influx mediates the assembly and activity of invadopodia and podosomes (Miyauchi et al., 1990; Siddiqui et al., 2012; Sun et al., 2014). Although the direct visualization of localized calcium signals in association with these matrix-degrading structures has been elusive (Visser et al., 2013), more recent studies have identified calcium oscillations as critical for invadopodium precursor assembly (Sun et al., 2014). Moreover, in macrophages, calcium influx supports short-lived calcium transients at the leading edge of polarized cells (Evans and Falke, 2007) where podosomes and podosome rosettes reside (Evans et al., 2003; Siddiqui et al., 2012). Consistent with their regulation by calcium, podosomes and invadopodia have been reported to be modulated by a number of calcium channels, including the store-operated calcium channel Orai1, the calcium release–activated calcium channel, and small-conductance calcium-activated potassium channels (Siddiqui et al., 2012; Sun et al., 2014), as well as Ca2+-regulated proteins such as calpain, Pyk2, calmodulin, Iba1 (Calle et al., 2006; Gil-Henn et al., 2007; Siddiqui et al., 2012), and now S100A4.

Podosomes are recognized as mechanosensors whose activity and dynamics are modulated by actomyosin contractility...
Mechanosensing is mediated by two populations of actomyosin filaments: lateral filaments within individual podosomes that link the top of the podosome core to the adhesion protein ring structure and filaments that connect individual podosomes into higher-order superstructures (Linder and Wiesner, 2015). An appropriate balance of actomyosin contractility is critical for the maintenance of podosomes, as podosome dissolution is caused by disruption of myosin-II binding to F-actin by blebbistatin or inhibition of myosin regulatory light-chain phosphorylation (Kopp et al., 2006; Collin et al., 2008; van Helden et al., 2008). Similarly, podosome formation is inhibited by expression of kinase-dead TRPM7, a cation channel fused to an α-kinase that phosphorylates the myosin-IIA heavy chain to mediate myosin-IIA filament assembly through heavy chain–directed mechanisms is critical for tuning actomyosin contractility, which could impact individual podosomes via the modulation of oscillatory behavior (van den Dries et al., 2013; Labernadie et al., 2014) and the organization of podosomes into higher-order structures such as clusters and rosettes (Meddens et al., 2016).

Our studies with blebbistatin also reveal myosin-IIA–independent effects of S100A4 loss that include enhanced α-tubulin K40 acetylation and MMP9 secretion. Because microtubule acetylation stimulates the trafficking of kinesin-associated vesicles in vivo (Bhuwania et al., 2014) and MMP9-containing vesicles traffic along stabilized microtubules (Hanania et al., 2012), we suspect that the increase in MMP9 secretion is likely a secondary consequence of the increase in microtubule acetylation and stability observed in S100A4−/− BMMs.

How might loss of S100A4 modulate microtubule acetylation? Microtubule acetylation is stimulated by the acetyltransferase α-TAT1 (MEC-17) and inhibited by the deacetylases HDAC6 and Sirtuin-2 (North et al., 2003; Sadoul and Khochbin, 2016). Paxillin, a focal adhesion– and podosome-associated scaffold protein, has been shown to regulate microtubule acetylation by directly inhibiting HDAC6 (Deakin and Turner, 2014). We have previously shown that S100A4 deletion has profound effects on paxillin (Li et al., 2010); pY118-paxillin is increased in CSF-1–stimulated S100A4−/− BMMs, and there is a dramatic decrease in pY118-paxillin associated with detergent-resistant cytoskeletons. This suggests that S100A4 may play a role in the targeting of paxillin to cytoskeletal structures, which could impact HDAC6 localization or activity.

In addition, S100A4 could regulate microtubule dynamics via interactions with liprin B1 (Kriaevskaya et al., 2002), a scaffolding protein associated with cortical assemblies that cluster near focal adhesions to mediate microtubule stability (van der Vaart et al., 2013; Chiaretti and de Curtis, 2016; Dong et al., 2016). Recent studies have identified liprin B1 as a Sirtuin-2–interacting protein (Budayeva and Cristea, 2016). While it is not known how liprin B1 regulates Sirtuin-2, loss of S100A4 could affect liprin B1/Sirtuin-2 interactions and Sirtuin-2 activity or subcellular targeting. Thus, S100A4 could modulate microtubule acetylation through a number of mechanisms whose investigation is beyond the scope of the current study.

While the precise link between S100A4 and microtubule acetylation remains unknown, changes in microtubule stability could dramatically affect podosome dynamics and their organization into higher-order structures like rosettes. At the level of individual podosomes, microtubule acetylation decreases microtubule–podosome contacts and delivery of MMP-containing vesicles and MMP9 secretion.
also decreases podosome number in human macrophages (Bhuvania et al., 2014). While this might suggest that disruption of microtubule–podosome contacts is responsible for impaired invasion in S100A4−/− BMMs, blebbistatin rescues matrix degradation by individual podosomes but does not rescue invasion.

The finding that S100A4 loss stabilizes podosome rosettes, as does tubacin treatment of wild-type macrophages, suggests an alternative hypothesis, in which the stabilization of podosome rosettes inhibits three-dimensional (3D) invasion. Consistent with this idea, microtubule destabilization in human primary macrophages promotes podosome disassembly (Linder et al., 2000). Furthermore, in osteoclasts, where podosomes assemble into peripheral belts that form sealing rings required for degradation of bone matrix, microtubule acetylation increases the formation of podosome belts (Biosse Duplan et al., 2014) and enhances belt stability (Destaing et al., 2005). Similarly, loss of microtubule acetylation in Pyk2−/− osteoclasts is associated with decreased podosome-belt formation (Gil-Henn et al., 2007). In these studies, podosome-belt stabilization caused by microtubule acetylation correlates with an increase in two-dimensional (2D) matrix degradation. However under 2D culture conditions on calcite, the podosome belts observed in osteoclasts are relatively stable structures (Batsir et al., 2017), perhaps due to the high density of the extracellular matrix they encounter (Collin et al., 2006). In contrast, in Src-transfected fibroblasts plated on gelatin or transforming growth factor β1–stimulated THP-1 cells plated on fibronectin, podosome rosettes and superstructures are highly dynamic, with striking changes in both shape and localization within the seconds to minutes timescale (Cox et al., 2011; Kedziora et al., 2016). While stabilization of podosome rosettes by microtubule acetylation in osteoclasts might increase rosette degradative capacity, human macrophages moving through a 3D collagen gel rapidly disassemble and reassemble podosomes (lifetime of ∼5 min) (Van Goethem et al., 2011), and enhanced stabilization of rosettes could interfere with this remodeling. This is analogous to observations from NIH 3T3 cells, in which microtubule acetylation increases the stability of focal adhesions and decreases invasion (Tran et al., 2007). In future experiments, live-cell imaging of podosome rosettes will test whether a disruption of rosette dynamics is responsible for the reduced invasion by S100A4−/− BMMs.

In summary, we find that S100A4 plays complex roles in the regulation of macrophage function. The defect in invasion by S100A4−/− BMMs involves S100A4 regulation of both the actomyosin and microtubule cytoskeletons. The myosin-IIA–dependent and myosin-IIA–independent functions of S100A4 appear to converge on the regulation of podosome rosettes, and the formation and dynamics of these structures emerge as critical regulators of macrophage motility and invasion.

FIGURE 6: Podosome rosettes are more stable in S100A4−/− BMMs. (A) Representative images of podosome rosette disassembly in S100A4−/− and wild-type BMMs. BMMs were serum and CSF-1 starved, plated on gelatin, and then stimulated with medium (final concentrations: 60 ng/ml CSF-1 and 7.5% serum). Starved BMMs (0 min) have rosettes (arrows), which disassemble into individual podosomes (F-actin punctae). Inset: individual podosomes within a rosette. Scale bar: 50 µm. (B) Quantification of percent of S100A4−/− and wild-type BMMs with podosome rosettes. Data represent the mean ± SEM from three independent experiments. Inset: the number of cells with rosettes plotted as a percent of the number at t = 0.

MATERIALS AND METHODS

Antibodies and reagents

The nonmuscle myosin-IIA rabbit antibody was from Biomedical Technologies (cat. no. BT-567). Vinculin (cat. no. V9131), β-actin (cat. no. A1978), α-tubulin (cat. no. T6074), and acetylated K40 α-tubulin (cat. no. T7451) antibodies were from Sigma. The MMP9 polyclonal antibody against the catalytic domain of rat MMP9 was from Millipore (cat. no. AB19016). The MMP9 polyclonal antibody against mouse full-length MMP9 (Ala-20–Pro-730) was from R&D Systems (cat. no. AF909-SP). Recombinant human proMMP9 (93 kDa) was from R&D Systems (cat. no. 911-MP-010). Rhodamine phalloidin was from Invitrogen. The active and inactive blebbistatin
isomers were purchased from Calbiochem; and BFA, tubacin, and nocodazole were purchased from Sigma.

Isolation of bone marrow–derived macrophages

S100A4−/− mice were maintained and bred in a pathogen-free barrier facility. All experiments were performed according to protocols approved by the Animal Welfare Committee at the Albert Einstein College of Medicine. BMMs were isolated from 12- to 14-wk-old mice as described previously (Li et al., 2010). Mature macrophages were cultured in complete BMM medium (α-MEM containing 0.02 mg/ml asparagine, 2 mM l-glutamine, 100 U/ml penicillin G, 100 µg/ml streptomycin sulfate, 15% fetal bovine serum, and 10,000 U/ml human recombinant CSF-1 [kindly provided by E. Richard Stanley, Albert Einstein College of Medicine]). Day 8–11 BMMs were used for all experiments.

Transwell invasion assays

BMMs were starved for 5 h in serum and CSF-1–free complete BMM medium. In all, 100,000 BMMs were seeded into the upper chambers of 8-µm Transwells coated with growth factor–reduced Matrigel (Bicoid #354483, Corning) in 400 µl of starvation medium. The Transwells were placed into a 24-well plate containing 500 µl of complete BMM medium, and the cells were allowed to migrate for 24 h at 37°C. For control samples, starvation medium was placed into the lower well. For invasion assays in the presence of blebbistatin, 5 µM of the active (−) or inactive (+) isomer was added to both the upper and lower wells. For invasion assays in the presence of tubacin, 2.5, 5, or 25 µM tubacin was added to both the upper and lower wells. After molecular weight cutoff. The antibody arrays were used to analyze the conditioned media according to the manufacturer’s protocol.

Gelatin zymography

Conditioned medium was prepared as described in the preceding section. Samples were diluted in nonreducing sample buffer and separated on a 9.5% Tris-Tricine gel (0.95 M Tris, 0.10% SDS, 0.95% glycerol) containing 0.1% gelatin with a running buffer that contained 0.1 M Tris, 0.1 M Tricine, and 0.1% SDS. Gels were then washed twice for 30 min each in 2.5% Triton X-100 and incubated overnight at 37°C in developing buffer (50 mM Tris, pH 8.0, 2% Na2HPO4, 0.02% NaN3, 2% gelatin, 20% glycerol) containing 0.1% BSA. Gels were examined for active enzymes, and the gels were stained with Coomassie staining solution for automated counting of nuclei by setting up a threshold. Each nucleus present on the lower surface of each Transwell was scored as one invasion event. Invasion toward complete BMM medium was expressed as the fold change relative to invasion in the absence of serum and CSF-1 in the lower well.

Antibody arrays

Mouse Angiogenesis Proteome Profiler Arrays were purchased from R&D Systems (ARY015). Conditioned medium was prepared from 1.5 × 106 BMMs seeded onto 60-mm tissue-culture plates. Cells were allowed to attach and washed three times with PBS, and 2 ml of serum-free complete BMM medium was added to the plate. After 24 h, conditioned medium was collected, filtered with a 0.22-µm Millex-GV filter unit, and concentrated approximately eightfold with an Amicon Ultra Centrifugal Filter (3000-Da molecular weight cutoff). The antibody arrays were used to analyze the conditioned media according to the manufacturer’s protocol.

MMP9 activity assay

MMP9 activity in conditioned media collected from BMMs (as described earlier) was determined using the MMP9 Biotrak Activity Assay (GE Healthcare Life Sciences) following the manufacturer’s protocol. For measurement of total MMP9 activity (pro- and active MMP9), 1 mM p-aminophenylmercuric acetate was added to the samples. An MMP9 standard curve (0.00625–0.05 ng of recombinant human proMMP9) was used to determine the concentration of active MMP9 in the conditioned media.

RNA isolation and RT-qPCR

RNA was purified with TRIzol (Roche) according to the manufacturer’s protocol and reverse transcribed with Moloney murine
performed in duplicate. Statistical analysis was performed using Data represent the mean ± SEM from three independent experiments. (C) Podosome rosette area in S100A4−/− and wild-type BMMs. Data represent the mean ± SEM from two to four independent experiments. (D) Matrigel invasion of S100A4−/− and wild-type BMMs toward growth medium. BMMs were starved for 5 h in serum and CSF-1–free complete BMM medium before being seeded into the upper chambers of 8-µm Transwells coated with growth factor–reduced Matrigel. Stimulated invasion is expressed as the fold change relative to unstimulated cells. NT, not treated; B(+), inactive isomer of blebbistatin; B(−), active isomer of blebbistatin. Data represent the mean ± SEM from three independent experiments performed in duplicate. Statistical analysis was performed using one-way ANOVA.

For acetylated α-tubulin/total α-tubulin immunoblots, cells were seeded as described for the gelatin zymography. For MMP9 immunoblots, 750,000 BMMs were plated on 60 mm tissue culture dishes in complete BMM medium. For MMP9 antibodies were used for immunoblots. MMP9 from BFA-treated samples were prepared as described in the preceding section. Samples were separated by 10% SDS–PAGE, transferred to Immobilon-FL polyvinylidene difluoride (PVDF; Millipore), and probed with MMP9 antibodies. Two different MMP9 antibodies were used for immunoblots. MMP9 from BFA-treated cells could only be detected with the MMP9 antibody from R&D Systems, which was made against mouse full-length MMP9. However, this antibody poorly recognizes the pro- and active forms of MMP9 in untreated cells. The MMP9 antibody from Millipore, made against the MMP9 catalytic domain, was used to detect pro- and active MMP9 in cell lysates from untreated cells.

**Immunofluorescence and microscopy**

Typically, cells were fixed in 3.7% formaldehyde in PBS for 15 min and permeabilized with 0.5% Triton X-100 in fixative for 3 min. For immunostaining of α-tubulin, cells were fixed and permeabilized in 100% methanol for 20 s at −20°C. For studies evaluating nocardazole-mediated microtubule depolymerization, cells were treated with 10 µM nocardazole for 5 min before fixation. Cells were blocked in PBS containing 1% bovine serum albumin (BSA) for 1 h and incubated with primary antibodies in PBS containing 1% BSA for 1 h using the following final dilutions: α-tubulin (1:200), myosin-IIA (1:100), and vinculin (1:200). Cells were then incubated with fluorochrome-conjugated secondary antibodies (1:500) and rhodamine phalloidin (1:300) in PBS containing 1% BSA for 30 min. Coverslips were mounted using DAPI Fluoromount-G.

Epifluorescence images were acquired with a Plan-Apo 60x, 1.4 NA objective (Olympus) and HiQ band-pass filters (Chroma Technology). All images were acquired using MetaMorph software and a CoolSNAP HQ interline 12-bit, cooled CCD camera.
FIGURE 9: Tubacin treatment inhibits Matrigel invasion of wild-type BMMs. (A) Representative immunoblot of α-tubulin K40 acetylation in whole-cell lysates from wild-type BMMs treated with DMSO or tubacin and from untreated S100A4−/− BMMs. (B) Quantification of α-tubulin K40 acetylation in whole-cell lysates from untreated and tubacin-treated wild-type BMMs and untreated S100A4−/− BMMs. Acetylation is expressed as the ratio of acetylated α-tubulin to total α-tubulin normalized to wild-type BMMs. Data represent the mean ± SEM from four independent blots. Statistical analysis was performed using one-way ANOVA. (C) Rosette disassembly assay. Wild-type BMMs were starved, plated, treated with DMSO or 5 µM tubacin for 90 min, and stimulated as in Figure 6B. Data represent the mean ± SEM from three independent experiments. (D) Matrigel invasion of tubacin-treated wild-type BMMs toward growth medium. BMMs were starved for 5 h in serum and CSF-1–free complete BMM medium before being seeded into the upper chambers of 8-µm Transwells coated with growth factor–reduced Matrigel, in the presence of DMSO or 2.5–25 µM tubacin. Stimulated invasion is expressed as the fold change relative to unstimulated cells. Control: vehicle-treated BMMs. Data represent the mean ± SEM from three independent experiments performed in duplicate. Statistical analysis was performed using one-way ANOVA.

Gelatin degradation assays

Acid-washed glass coverslips were treated with 0.01% poly-L-lysine for 10 min at room temperature followed by washing with PBS. Coverslips were then coated with 200 µg/ml Oregon Green 488–conjugated gelatin (Life Technologies) for 15 min at room temperature and washed extensively with PBS. BMMs were starved in serum-free complete BMM medium for 3 h, and then 1.5 x 10⁵ BMMs were plated on the coverslips in the same medium. After 1 h, serum was added, and the cells were incubated for an additional 17 h. Cells were then fixed with 3.7% paraformaldehyde in PBS for 15 min and stained with rhodamine phalloidin and DAPI. At least 10–20 fields per condition were imaged using an UPlanFl 20×, 0.3 NA objective. The degradation area per field was measured by thresholding the images using ImageJ software to determine the total area in the field that lacked fluorescence. The percent degradation area was defined as the total degraded area per field divided by the total cell area per field based on rhodamine phalloidin staining. The number of cells in the field was determined by counting DAPI-stained nuclei. Three independent experiments were performed with a minimum of 700 cells per condition examined for each experiment. For studies with blebbistatin, cells were incubated overnight with 5 µM of the active (−) or inactive (+) isomers of blebbistatin. Three independent experiments were performed with a minimum of 230 cells per condition examined for each experiment.

Analysis of podosome rosette formation and size

BMMs were treated, plated, fixed, and stained as described for gelatin degradation assays. At least 20 fields per condition were imaged using an UPlanFl 20×, 0.3 NA objective and 1.5× magnification. Podosome rosettes were defined as ring-like structures composed of podosomes. The number of podosome rosettes per field was divided by the total cell number per field, which was determined by DAPI staining. Three independent experiments were performed with a minimum of 460 cells per condition examined for each experiment.

For the analysis of podosome rosette size, BMMs were cultured overnight on 200 µg/ml gelatin, fixed, and stained with rhodamine phalloidin and DAPI. For studies with blebbistatin, cells were incubated overnight with 5 µM of the active (−) or inactive (+) isomers of blebbistatin. At least 20 fields per condition were imaged using a Plan-Apo 60×, 1.4 NA objective. The area (in pixels) of podosome rosettes was measured using ImageJ software and the Trace tool. Three independent experiments were performed with a minimum of 86 rosettes per condition traced in each experiment.

Podosome disassembly assays

BMMs were serum and CSF-1 starved for 2 h and then plated on 200 µg/ml gelatin in starvation medium for 1 h to allow cells to form podosome rosettes. BMMs were then stimulated by the addition of an equal volume of complete BMM medium (60 ng/ml CSF-1 and 7.5% serum final concentration) to promote rosette disassembly. For studies with tubacin, cells were incubated with DMSO or 5 µM tubacin for 90 min before stimulation. Cells were fixed before stimulation (0 min) and at 5, 10, and 20 min after stimulation, and then stained with rhodamine phalloidin and DAPI. At least 15 fields per condition were imaged using a Plan-Apo 60×, 1.4 NA objective. The number of podosome rosettes per field was counted and divided by the total cell number per field. Three independent experiments were performed with a minimum of 70 cells per time point examined for each experiment.
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