Endosomal WASH and exocyst complexes control exocytosis of MT1-MMP at invadopodia

Pedro Monteiro,1,2 Carine Rossé,1,2 Antonio Castro-Castro,1,2 Marie Iordonelle,1,2 Emilie Lagoutte,1,2 Perrine Paul-Gilloteaux,1,3 Claire Desnos,4,5 Etienne Formstecher,6 François Darchen,4,5 David Perrais,7,8 Alexis Gautreau,9 Maud Hertzog,1,2 and Philippe Chavrier1,2

1Research Center, Institut Curie, 75005 Paris, France
2Membrane and Cytoskeleton Dynamics and 3Cell and Tissue Imaging Facility, Centre National de la Recherche Scientifique UMR 144, 75005 Paris, France
3Université Paris Descartes and 4Centre National de la Recherche Scientifique, UMR8192, 75006 Paris, France
4Hybrigenics services SAS, 75014 Paris, France
5University of Bordeaux and 6Centre National de la Recherche Scientifique, Interdisciplinary Institute for Neuroscience, UMR 5297, 33000 Bordeaux, France
7Laboratoire d’Enzymologie et Biochimie Structurales, Centre National de la Recherche Scientifique UPR3082, 91198 Gif-sur-Yvette, France

Introduction

The matrix-degrading protrusions of the plasma membrane known as invadopodia are currently thought to form in invasive tumor cells when the extracellular matrix and cues from the tumor microenvironment, such as growth factors, trigger the assembly of F-actin into precursor structures through a signaling cascade involving Cdc-42 and Nck1 and the actin regulatory proteins neuronal Wiskott-Aldrich syndrome protein (N-WASP), Arp2/3 complex, cortactin, and coflin (Lorenz et al., 2004; Yamaguchi et al., 2005; Clark et al., 2007; Ayala et al., 2008; Oser et al., 2009, 2010; Murphy and Courtneidge, 2011). These precursors then mature into functional invadopodia upon accumulation of the trans-membrane type 1 matrix metalloproteinase (MT1-MMP). Artyom et al., 2006; Clark et al., 2007; Sakurai-Yageta et al., 2008; Liu et al., 2009; Williams and Coppolino, 2011). Based on these findings, we proposed that this cohort of proteins regulates the trafficking and exocytosis of MT1-MMP from late endocytic storage compartments to its invadopodial target plasma membrane (Poincloux et al., 2009).

A significant fraction of MT1-MMP is internalized from the cell surface as a means to regulate its surface level (Jiang et al., 2001; Uekita et al., 2001). In MDA-MB-231 human breast adenocarcinoma cells, we found the majority of intracellular MT1-MMP located in a late endosome compartment (Steffen et al., 2008). We and others reported that an exocytic machinery comprising cortactin, the vesicle-docking exocyst complex, and the SNARE protein vesicle-associated membrane protein 7 (VAMP7) is required for MT1-MMP delivery to invadopodia. However, the nature of the carriers that mediate plasma membrane

required for matrix degradation by an exocytic mechanism that involves tubular connections between MT1-MMP–positive late endosomes and the plasma membrane in contact with the matrix. This ensures focal delivery of MT1-MMP and supports pericellular matrix degradation and tumor cell invasion into different pathologically relevant matrix environments. Our data suggest a general mechanism used by tumor cells to breach the basement membrane and for invasive migration through fibrous collagen-enriched tissues surrounding the tumor.
delivery of MT1-MMP, the mechanism underlying MT1-MMP exocytosis in the biogenesis of invadopodia, and how exocytosis is possibly influenced by the composition and biophysical properties of the matrix remain poorly understood.

Recent studies have documented an essential role for actin cytoskeleton dynamics in endosome function (Derivery et al., 2009; Gomez and Billadeau, 2009; Morel et al., 2009; Puthenveedu et al., 2010; Carnell et al., 2011; Harrington et al., 2011). The mechanism emerging from these on-going studies indicates that actin–Arp2/3 assemblies organize early endosomal membranes into functional subdomains and contribute to cargo sorting and generation of transport intermediates. Some of these studies also highlighted the essential role of the newly identified Wiskott-Aldrich syndrome protein and Scar homolog (WASH) complex, a member of the WASP (Wiskott–Aldrich syndrome protein) family of Arp2/3 activators associated with the endosomal/lysosomal system and playing a major role in the polymerization of endosomal actin (Derivery et al., 2009; Gomez and Billadeau, 2009; Duleh and Welch, 2010). All together, these data support a critical role for WASH in linking Arp2/3 and F-actin–assisted elongation and fission of endosomal tubules with sorting and trafficking from the endosomal system to the cell surface (Derivery et al., 2009; Gomez and Billadeau, 2009; Puthenveedu et al., 2010; Carnell et al., 2011; Temkin et al., 2011; Zech et al., 2011; Gomez et al., 2012). Here, we describe a novel interaction of WASH with the exocyst complex on MT1-MMP–containing late endosomes in invasive breast tumor cells. Our data support a mechanism of exocytosis of MT1-MMP through late endosome-to-plasma membrane connections occurring at invadopodia and requiring WASH and exocyst complexes for their formation.

**Results**

**WASH and the exocyst complex interact on MT1-MMP-positive endosomes in breast tumor cells**

In a series of yeast two-hybrid screens aimed at isolating partners of the eight subunits of the exocyst complex, we identified interactions of both Exo84 and Sec3 exocyst subunits with the amino-terminal region of WASH. Overlapping regions from independent isolated clones defined WASH domains interacting with Sec3 (amino acids 9–109 of WASH) or with Exo84 (amino acids 15–258 of WASH; unpublished data). By using fluorescence microscopy of MDA-MB-231 cells overexpressing GFP-WASH and immunofluorescence microscopy of the endogenous protein and in agreement with the function of WASH as an activator of the Arp2/3 complex (Derivery et al., 2009; Gomez and Billadeau, 2009; Duleh and Welch, 2010), we observed WASH puncta closely associated with punctate accumulations of Arp2/3 complex, F-actin, and the F-actin–binding protein cortactin on MT1-MMP–positive endosomes (Fig. S1 A and Fig. 1, A and B). Exo84 was also detected as puncta closely apposed to and partially overlapping with WASH and cortactin-positive subdomains on MT1-MMP–containing endosomes (Fig. 1, C and D; and Fig. S1 B). Live-cell imaging revealed highly dynamic WASH puncta associated with cortactin (Video 1). The number of WASH-cortactin patches seemed to correlate with endosome size (Fig. 1 B; Gomez and Billadeau, 2009; Derivery et al., 2012). Endosomal puncta of WASH and cortactin were more closely associated (mean distance of ~2 pixels) as compared with WASH and Exo84-positive ones (~5 pixels; Fig. 1 E). Similarly, we observed another component of the WASH complex, FAM21, associated with cortactin domains on MT1-MMP–positive endosomes (Fig. S1 C). WASH was also apposed to punctate structures containing the late endocytic SNARE protein VAMP7 on the limiting membrane of these endosomes (Fig. S1 D), in agreement with the predominant localization of MT1-MMP to late endosomes (Steffen et al., 2008; Williams and Coppolino, 2011). WASH- and cortactin-positive puncta were similarly associated with MT1-MMP–containing endosomes in BT-549 breast cancer cells (Fig. S1 E). To further evaluate association of WASH with cortactin or Exo84 on MT1-MMP–containing endosomes, we performed a Duolink in situ proximity ligation assay (Söderberg et al., 2006). This analysis confirmed a strong association between WASH and cortactin and a weaker one between WASH and Exo84, with most of the Duolink signal coinciding with MT1-MMP–containing endosomes (Fig. 1, F and G). Finally, endogenous WASH protein in HeLa cell extracts coimmunoprecipitated with overexpressed full-length Exo84 and Sec3 and with endogenous Sec8 (Fig. 1 H), thus confirming the two-hybrid results and showing that WASH interacts with the exocyst complex in vivo. Together, these findings reveal the existence of WASH and Arp2/3–F-actin–cortactin–positive domains on MT1-MMP–containing late endosomes. The exocyst complex is closely apposed to and interacts with WASH-positive subdomains on MT1-MMP–positive endosomes, suggesting a coordinated function of these complexes in breast tumor cells.

**WASH and exocyst control actin assembly on MT1-MMP-positive endosomes**

To investigate the roles of WASH and the exocyst complex we first looked at the morphology of MT1-MMP–positive endosomes in MDA-MB-231 cells silenced for WASH or Exo84 with two independent siRNAs (Fig. S2, A and B). As previously reported (Steffen et al., 2008), MT1-MMP accumulated in large vacuolar compartments in the central region of the cell as well as in smaller vesicles spread throughout the cytoplasm (Fig. 2 A, class #1). Silencing of WASH resulted in drastic changes in the morphology of the MT1-MMP–containing endosomes: in ~60% of cells they aggregated in the center (Fig. 2, A and B, class #2) and in ~15% of them large clumps of MT1-MMP–positive structures appeared (Fig. 2, A and B, class #3). We observed a similar, albeit less marked, phenomenon in cells silenced for Exo84 (Fig. 2, A and B). Live-cell imaging revealed perinuclear aggregated MT1-MMP–positive endosomes in WASH-depleted cells, some with exaggerated tubular membrane extensions (Fig. 2 C and Video 2). This phenotype is reminiscent of endosome tubulation induced upon depletion of WASH seen in other cell types (Derivery et al., 2009; Gomez and Billadeau, 2009; Gomez et al., 2012). Given the proposed function of WASH-dependent actin/cortactin assembly in regulation of endosomal tubule dynamics, we explored the contribution of WASH and exocyst complex to the formation of Arp2/3–F-actin–cortactin punctate domains on MT1-MMP–positive late endosomes.
Silencing of WASH led to a large reduction of cortactin accumulation on endosomal compartments (Fig. 2, D and E). In contrast, silencing of N-WASP that is required for invadopodial actin assembly (Yamaguchi et al., 2005; Oser et al., 2009, 2010; Yu et al., 2012) had no effect on endosomal cortactin (and WASH)-positive microdomains (Fig. S3 C and Fig. 2, D and E). Interestingly, silencing of Exo84 was accompanied by a limited but significant increase of the size of WASH patches on MT1-MMP–positive endosomes and a marked lengthening of comet-shaped cortactin domains associated with WASH patches (Fig. 2, F and G). Similar observations were obtained upon depletion of Sec8 exocyst subunit (unpublished data). Together, these data indicate that WASH controls actin polymerization on MT1-MMP–positive endosomes and suggest that the exocyst complex associated with WASH may finely tune actin assembly required for regulation of endosomal membrane scission and remodeling (Derivery et al., 2009; Gomez and Billadeau, 2009; Park et al., 2013).
Figure 2. Regulation of MT1-MMP–positive endosome dynamics by WASH and exocyst complex. (A and B) MDA-MB-231 cells treated with the indicated siRNAs were plated on cross-linked gelatin and stained for endogenous MT1-MMP. Bars, 5 µm. Cells were scored according to the distribution of MT1-MMP–positive endosomes, i.e., normal distribution (#1, blue boxes in B), clustered endosomes (#2, green box), and aggregated endosomes (#3, red box). Values plotted in B are mean ± SEM of normalized endosome distribution from three independent experiments scoring ~100 cells for each siRNA. *, P < 0.05; **, P < 0.01; ***, P < 0.001 compared with siNT-treated cells (B). (C) MDA-MB-231 cells expressing MT1-MMP-mCh were treated as in A and time series were acquired by spinning-disk microscopy (see Video 2). Still images from the time series are shown. Arrows point to tubular extensions emanating from MT1-MMP–positive endosomes in cells depleted of WASH. Bars, 5 µm. (D) MDA-MB-231 cells treated with the indicated siRNA were processed for immunofluorescence microscopy with antibodies against cortactin (red) and WASH (green). Images were acquired by 3D deconvolution microscopy of the
WASH is present at docking sites of MT1-MMP-positive endosomes corresponding to invadopodia

Given that MT1-MMP is the key component for pericellular matrix degradation by invadopodia and that the exocyst complex is involved in MT1-MMP delivery to these structures (Sakurai-Yageta et al., 2008; Liu et al., 2009), we investigated whether WASH was also associated with invadopodia. Immunofluorescence microscopy of MDA-MB-231 cells plated on fluorescent gelatin revealed MT1-MMP–positive endosomes flanked by WASH-positive puncta in contact with the plasma membrane, some of which were associated with matrix degradation (Fig. 3, A–C). Observation by total interference reflection fluorescence microscopy (TIRFM) of cells plated on a thin layer of cross-linked gelatin coexpressing MT1-MMP:mCherry and GFP-tagged WASH or Exo84 showed that puncta of WASH and Exo84 associated with MT1-MMP–positive endosomes were in close contact with the adherent plasma membrane (Fig. 3, D and E). Similarly, WASH-positive puncta associated with MT1-MMP–containing endosomes were visible at the plasma membrane in BT-549 cells by TIRFM (Fig. 3 F). We also observed MT1-MMP–positive endosomes flanked by plasma membrane–apposed puncta containing Arp2/3, FAM21, cortactin, and Exo84 (Fig. 3, G and H; and Fig. S3, A and B). Matrix degradation was strictly confined to and superimposed upon WASH–FAM21–cortactin–Arp2/3–exocyst puncta adjacent to MT1-MMP–containing endosomes (Fig. 3, A, G, and H; arrowheads; and Fig. S3, A and B, arrowheads), strongly suggesting that these structures represent bona fide invadopodia. TIRFM of GFP-cortactin and MT1-MMP:mCh documented repeated docking of MT1-MMP–containing endosomes at the level of cortactin-positive invadopodia (Fig. 3, I and J; and Video 3). Interestingly, although all invadopodia were positive for N-WASP, association of WASH with cortactin-positive invadopodia was inversely correlated with the surface of matrix degradation underlying each invadopodium (Fig. 3 K and Fig. S3 C). These results suggest that WASH, associated with MT1-MMP–positive endosomes, interacts transiently with invadopodia and preferentially at an early stage of their lifetime (associated with smaller degradation area) in contrast to N-WASP that is permanently associated with invadopodia.

Invadopodia correspond to late endosome-to-plasma membrane connections

Based on the proposed role of WASH-mediated Arp2/3 activation in the formation of endosomal tubular membrane extensions (Puthenveedu et al., 2010; Gomez et al., 2012; Piotrowski et al., 2013), we hypothesized that WASH puncta, possibly in conjunction with the exocyst complex, may regulate the formation of transient continuities between late endosomes and the invadopodial plasma membrane to allow the delivery of MT1-MMP for matrix degradation. We added a tracer to the extracellular milieu to test whether the endosome lumen was accessible through such connections. MDA-MB-231 cells expressing MT1-MMP:mCh were plated on fluorescent gelatin and FITC-dextran was added for a 1-min pulse; then cells were examined immediately after washing out the tracer. A subset of MT1-MMP–positive endosomes were loaded with FITC-dextran during the pulse (Fig. 4 A, arrowheads), some exocytosed their MT1-MMP content while filling up with the dextran (Fig. 4 A, doublehead arrow). Cortactin-rich puncta and foci of matrix degradation were seen adjacent to FITC-dextran pulse-labeled endosomes (Fig. 4, B and C, arrowheads). In addition, after a 1-min pulse, fluorescent dextran–labeled endosomes were positive for Rab7 and VAMP7, arguing that these represent late endosomes making direct connection with the plasma membrane (Fig. 4, D and E). Furthermore, by live-cell TIRFM we observed regurgitation of the luminal FITC-dextran content of endosomes, some occurring through visible thin tubules extending from the endosome (Fig. 4 F). MT1-MMP–containing tubule emanating from plasma membrane–docked endosome was also visible by TIRFM (Fig. 4 G). These observations suggest the existence of tubular connections between MT1-MMP–positive late endosomes and the invadopodial plasma membrane that may mediate MT1-MMP delivery to the surface.

MT1-MMP exocytosis requires WASH and the exocyst complex

We used TIRFM to analyze the effect of silencing of WASH or Exo84 on the density of MT1-MMP:mCh–positive endosomes in the zone beneath the plasma membrane. Large numbers of MT1-MMP vesicles were visible at the adherent plasma membrane of MDA-MB-231 cells (Fig. 5 A). Silencing of WASH or Exo84 significantly reduced MT1-MMP–positive endosome density beneath the plasma membrane (Fig. 5, A and B), along with the observed collapse of MT1-MMP–positive endosomes in the cell center (Fig. 2, A and C). All together, these findings indicate that both WASH and the exocyst complex play an important role in polarization and tethers of MT1-MMP–positive endosomes at the plasma membrane. The contribution of WASH and Exo84 to the delivery of MT1-MMP to the invadopodial plasma membrane was assessed by quantifying the surface accumulation of MT1-MMP:pHluorin (Lizárraga et al., 2009); fluorescence of this construct is eclipsed in the acidic environment of the late endosome and rises upon exocytosis and exposure to the extracellular pH (Miesenböck, 2012). MT1-MMP:pHluorin accumulated at the level of cortactin-positive invadopodia on the ventral cell surface (Fig. 5, C and D). Knockdown of WASH or Exo84 led to a 50% reduction of surface middle plane of the cells. A region of interest (dashed line) was drawn manually around WASH-cortactin structures for quantification (shown in E). Bars, 5 µm. (E) Quantification of the integrated intensity of perinuclear endosomal cortactin and WASH in cells treated as in D and normalized to the integrated intensity in control cells in two independent experiments. ns, not significant; *, P < 0.05; **, P < 0.01 compared with control. (F) MDA-MB-231 cells treated with the indicated siRNAs were processed for immunofluorescence microscopy with cortactin and WASH antibodies and analyzed by high-resolution (~100-nm spatial resolution) 3D SIM. Bar, 1 µm. (G) Quantification of the projected surface of WASH and cortactin puncta in cells as described in F, normalized to mean projected surface of the puncta in control cells set to 100 (three independent experiments; n = 25 puncta for each condition; *, P < 0.05; ***, P < 0.001 compared with control).
Figure 3. Association of WASH endosomal puncta with invadopodia. (A) MDA-MB-231 cells plated on cross-linked FITC-labeled gelatin (blue) were stained with antibodies against MT1-MMP (red) and WASH (green) and imaged by 3D deconvolution microscopy. Image is a single section in the ventral plane of the cell in contact with gelatin. Insets corresponding to the boxed region show split channels with dashed line representing the contour of the MT1-MMP-containing endosome projected on the FITC-gelatin image. Arrowhead points to matrix degradation restricted to WASH puncta adjacent to MT1-MMP–positive endosome. Bars: (main) 5 µm; (inset) 1 µm. (B) Orthogonal section through the dashed line in A. From left to right: merged image, MT1-MMP (M), WASH (W), and FITC-gelatin (G) signals. (C) Fluorescence intensity profile through the dashed line in the boxed region in A (x-axis, in pixels; y-axis, in arbitrary units). (D and E) MDA-MB-231 cells expressing MT1-MMPmCh and GFP-WASH (D) or GFP-Exo84 (E) plated on cross-linked gelatin and...
accumulations of MT1-MMPpHluorin (Fig. 5 E), indicating that WASH and exocyst complex are critical for MT1-MMP exocytosis at invadopodia.

**WASH and exocyst complex are required for invasive migration through type I collagen**

Silencing of MT1-MMP is known to reduce drastically the ability of MDA-MB-231 cells to degrade a matrix of cross-linked gelatin as a basement membrane mimick (Artym et al., 2006; Sakurai-Yageta et al., 2008; Fig. 6 A). Silencing of WASH diminished the proportion of cells that were able to degrade gelatin and led to a 60–80% reduction in these cells’ capacity to do so, similar to the effect of silencing various subunits of the exocyst complex, e.g., Sec6, Sec8, and Sec10 (Sakurai-Yageta et al., 2008) and Exo84 (Fig. 6 A). MT1-MMP levels were unaffected by the various siRNA treatments (Fig. S2 E). By comparison, silencing of N-WASP also led to 60% inhibition of matrix degradation, similar to the effect seen on WASH depletion (Fig. 6 A). Thus, both plasma membrane- and endosome-associated actin nucleation-promoting factors, N-WASP and WASH, are required for invadopodia activity in breast tumor cells. In addition, depletion of WASH or Exo84 led to a significant reduction in the ability of MDA-MB-231 cells to invade through a layer of Matrigel, similar to MT1-MMP knockdown (Fig. 6 B).

Degradation of cross-linked gelatin and Matrigel is representative of carcinoma cells remodeling and breaching the basement membrane. MT1-MMP is also required for dissemination of cancer cells within fibrous type I collagen (Fig. 6 C; Sabeh et al., 2004, 2009; Wolf et al., 2007), thus the question arose whether or not the WASH- and exocyst-dependent mechanism of exocytosis of MT1-MMP–containing late endosomes also operated during remodeling of collagen fibers and invasive migration by tumor cells. We analyzed the capacity of MDA-MB-231 cells to invade the collagen I–filled, cell-free central portion of a 96-well plate over a 2-d culture period. Knockdown of WASH or Exo84 resulted in a strong inhibition of the invasion capacity of the cells in type I collagen similar to the silencing of MT1-MMP (Fig. 6, C and D). In addition, we found that silencing of WASH or Exo84 resulted in a 75–85% inhibition of pericellular collagen I degradation revealed by the Col1-3/4C antibody recognizing the cleaved fragment of collagen I dependent on MT1-MMP activity as indicated by the reduction of Col1-3/4C signal on MT1-MMP depletion (Fig. 6, E and F; Wolf et al., 2007). Thus these data indicate a general requirement for the exocyst complex and WASH for matrix remodeling and for the invasive potential of breast tumor cells.

Exocytosis of MT1-MMP occurs at contact sites with type I collagen fibers through a WASH- and exocyst-dependent mechanism

We analyzed the organization and dynamics of invadopodia in cells invading through 3D type I collagen environments in relation with WASH- and exocyst-positive MT1-MMP–containing endosomes. Staining of the cells for F-actin and cleaved collagen I fibers (Col1-3/4C antibody) revealed the formation of linear accumulations of F-actin on the inner face of the plasma membrane associated with collagen fibers, coincident with regions of collagenolytic activity (Fig. 7 A). These structures have been previously reported and named linear invadopodia (Juin et al., 2012). Similarly to classical dotty-like invadopodia on gelatin (Fig. S3 C), N-WASP colocalized with F-actin in linear invadopodia (Fig. 7 B). Labeling with anti-WASH antibodies showed the typical pattern of endosomal WASH-positive puncta (Fig. 7 C), some of which were closely apposed to linear invadopodia stained for cortactin although WASH did not accumulate in linear invadopodia (Fig. 7, C [inset], D, and E). We used live-cell imaging to analyze the dynamic interactions of WASH- and cortactin-positive puncta with linear invadopodia in cells in a 3D fibrillar collagen environment. Live-cell imaging of MDA-MB-231 cells expressing DsRed-cortactin and GFP-WASH (Video 4 and Fig. 7, D and E) plated on a layer of fluorescent collagen I fibers revealed the dynamic assembly of cortactin-positive linear invadopodia along collagen fibers and their disassembly upon remodeling and relaxation of the underlying fibers (Video 4 and Fig. 7 E). In addition, cortactin- and WASH-positive endosomal puncta interacted highly dynamically with linear invadopodia (Video 4 and Fig. 7, D and E). On examination of the time-lapse sequences there was no obvious transfer of cortactin from endosomal puncta to the linear invadopodia (Fig. 7 E). Analysis of MT1-MMP distribution in fixed cells showed that cortactin puncta were associated with MT1-MMP–positive endosomes (Fig. 7, F and G, arrows) in the vicinity of linear invadopodia positive for cortactin and MT1-MMP (Fig. 7, G and H, arrowheads), in agreement with the collagenolytic activity of these structures. All together, these data suggested that collagen I fibers elicit actin assembly and the formation of N-WASP–positive collagenolytic linear invadopodia and the dynamic recruitment of WASH-cortactin–positive endosomes probably involved in MT1-MMP delivery to these structures.

Exocytic events of MT1-MMP were monitored in live cells cultured on fibrillar collagen using the MT1-MMPpHluorin construct. In cells expressing simultaneously mCherry- and pHluorin-tagged MT1-MMP, dynamic late endosomes (red) were observed in the cytoplasm, with visible green flashes corresponding
repeated exocytosis in the vicinity of the collagen fibers (Fig. S4, A and B, arrowheads). Live-cell imaging of cells expressing MT1-MMPpHluorin and mCherry-WASH showed that MT1-MMP–containing endosomes undergoing exocytosis were to the de-quenching of pHluorin fluorescence upon exocytosis (Video 5 and Fig. S4). Endosomes scanning portions of the plasma membrane in contact with collagen I fibers were visible (Video 5 and Fig. S4, A and B, t = 0 min), which underwent repeated exocytosis in the vicinity of the collagen fibers (Fig. S4, A and B, arrowheads). Live-cell imaging of cells expressing MT1-MMPpHluorin and mCherry-WASH showed that MT1-MMP–containing endosomes undergoing exocytosis were
flanked by WASH puncta (Video 6 and Fig. 8, A and B, red arrowheads) and exocytosis preferentially occurred at contact sites with fibers (Video 6 and Fig. 8, A and B, green arrowheads). Multiple exocytic events led to surface accumulation of MT1-MMP along the collagen fibers (at linear invadopodia; Fig. 8 A, white arrowheads) and fiber remodeling (Video 6). Quantitative assessment of MT1-MMPpHluorin flashes revealed a strong inhibition of MT1-MMP exocytic events upon knockdown of WASH or Exo84 (Fig. 8 C), whereas overall MT1-MMP expression was similar in the different cell populations (not depicted).

So far, our observations were compatible with a WASH/Exocyst-dependent mechanism whereby MT1-MMP–positive endosomes exocytose the protease at contact sites with collagen fibers but do not collapse within the plasma membrane bilayer. They also revealed that MT1-MMPpHluorin flashes lasted for several minutes (Fig. 8 D). In addition, we noticed that although some endosomes with de-quenched MT1-MMPpHluorin were moving along fibers and could possibly detached and reengage with other sites for multiple rounds of MT1-MMP exocytosis, some remained at the same position for several minutes (Videos 5 and 6; Fig. S4; and Fig. 8, A and B). These findings raised the issue of the duration of endosome-to-plasma membrane openings. Using an extracellular pH shifts protocol (Video 7; Perrais et al., 2004), we observed that some endosomes remained opened for several minutes as shown by the fluorescence of pHluorin in the endosomal lumen that was sensitive to the iterative shifts of extracellular pH (at least for 4 min in the example shown in Fig. 8 E and up to 15 min [not depicted]). Of note, similar long-lasting fusion pore openings have been reported during zymogen granule exocytosis in exocrine pancreatic acinar cells (up to 15 min; Perrais et al., 2004; Thorn et al., 2004; Sokac and Bement, 2006). Moreover, the exchange of low and high pH solution within the endosome could take up to 30 s, consistent with the restricted diffusion of solutions through a long tubular connection. Thus, through the functions of WASH and exocyst complex, tumor cells can engage late endocytic MT1-MMP storage compartment in the formation of tubular continuities with the...
membrane proteins in different cell types (Derivery et al., 2009; Gomez and Billadeau, 2009; Zech et al., 2011; Gomez et al., 2012; Piotrowski et al., 2013). In addition, the exocyst complex, which mediates the tethering of exocytic vesicles to the plasma membrane, is also required for the delivery of MT1-MMP to invadopodia (Sakurai-Yageta et al., 2008; Liu et al., 2009; this study). Strikingly, our study reveals that WASH- and F-actin–positive puncta on MT1-MMP endosomes are adjacent to exocyst-positive puncta on this compartment and some components of these two multiprotein complexes (WASH, Exo84, and Sec3 based on two-hybrid data) can interact with each other. The segregation of WASH domains on the cytosolic face of the endosomal membrane depends on actin polymerization, which is induced by WASH-mediated activation of the Arp2/3 complex (Derivery et al., 2012). The mechanism underlying the subdomain organization of the exocyst complex at the surface of MT1-MMP–positive endosomes is presently unknown; however, we found that exocyst components can negatively regulate WASH-dependent endosomal actin

Discussion

Our data, which are summarized in a model presented in Fig. 9, support a general mechanism of invadopodia formation whereby the functions of WASH and exocyst complex on closely apposed punctate domains on the cytosolic face of MT1-MMP–positive late endosomes ensure focal delivery of MT1-MMP for pericellular matrix degradation.

We found that loss of WASH function induces disappearance of endosomal F-actin puncta and a collapse of MT1-MMP–positive late endosomal system, which is accompanied by defects in MT1-MMP recycling and delivery to the invadopodial plasma membrane. This is in agreement with an established function of the WASH complex in the recycling route of various plasma membrane proteins in different cell types (Derivery et al., 2009; Gomez and Billadeau, 2009; Zech et al., 2011; Gomez et al., 2012; Piotrowski et al., 2013). In addition, the exocyst complex, which mediates the tethering of exocytic vesicles to the plasma membrane, is also required for the delivery of MT1-MMP to invadopodia (Sakurai-Yageta et al., 2008; Liu et al., 2009; this study). Strikingly, our study reveals that WASH- and F-actin–positive puncta on MT1-MMP endosomes are adjacent to exocyst-positive puncta on this compartment and some components of these two multiprotein complexes (WASH, Exo84, and Sec3 based on two-hybrid data) can interact with each other. The segregation of WASH domains on the cytosolic face of the endosomal membrane depends on actin polymerization, which is induced by WASH-mediated activation of the Arp2/3 complex (Derivery et al., 2012). The mechanism underlying the subdomain organization of the exocyst complex at the surface of MT1-MMP–positive endosomes is presently unknown; however, we found that exocyst components can negatively regulate WASH-dependent endosomal actin

plasma membrane at contact sites with the extracellular matrix to allow delivery of the protease to the surface and to ensure matrix remodeling required for invasive migration.
endosome organization and for MT1-MMP recycling to the surface. Therefore, in addition to its well-established function as a tethering factor for exocytic vesicles to the plasma membrane (Heider and Munson, 2012), our findings suggest an unprecedented role for the exocyst in regulating endosomal membrane dynamics.

Remarkably, we found that MT1-MMP–positive endosomes contact and establish connections with the plasma membrane in assembly. In this respect, two-hybrid data mapping of the binding site of Exo84 and Sec3 exocyst subunits to the amino-terminal region of WASH suggests a possible competition mechanism because this region of WASH is also required for association with other WASH complex components (Gomez and Billadeau, 2009; Jia et al., 2010). A consequence of exocyst’s interaction with WASH could thus be a fine regulation of actin assembly required for late

endosome organization and for MT1-MMP recycling to the surface.

Figure 7. WASH- and cortactin-positive MT1-MMP–containing endosomes are recruited in the periphery of linear invadopodia in association with collagen fibers. (A) MDA-MB-231 cells on a layer of collagen I fibers stained with anti-Col1-3/4C antibodies recognizing MMP-cleaved collagen I (red) and for F-actin (green) revealing collagenolytic linear invadopodia. Inset is a higher magnification of the boxed region. Bars, 5 µm. (B and C) Cells were plated on a layer of Alexa Fluor 647–conjugated type I collagen (blue) for 30 min and stained for N-WASP (green) and F-actin (red); B or WASH (green) and cortactin (red; C). Insets are higher magnification of boxed regions. Bars: (main and B [inset]) 5 µm; (C, inset) 1 µm. (D) MDA-MB-231 cells expressing GFP-WASH (green) and DsRed-cortactin (red) were plated on a layer of Alexa Fluor 647–conjugated type I collagen (blue) for 30 min and imaged by confocal spinning disk microscopy. A still image from a representative time-lapse series is shown (t = 25 min). Arrowheads point to WASH- and cortactin-positive puncta (associated with late endosomes) opposed to a cortactin-positive linear invadopodium aligned on collagen fibers. Insets show a higher magnification of WASH and cortactin endosomal puncta. The corresponding time-lapse series is shown in Video 4. Bars: (main) 5 µm; (inset) 1 µm. (E) Higher magnification of the boxed region in D at representative time points. A linear cortactin-positive invadopodium aligned with collagen fibers forms on the inner face of the plasma membrane (13–14 min; red arrowheads); at 60 min, this structure disassembled while collagen I fibers were remodeled. Time is in minutes. Bar, 5 µm. (F–H) Cells on Alexa Fluor 647–conjugated type I collagen (blue) stained for MT1-MMP (red) and cortactin (green). G and H are higher magnification of the boxed region in F. Arrows point to cortactin-positive puncta associated with MT1-MMP–containing endosomes in the vicinity of cortactin- and MT1-MMP–positive linear invadopodia (arrowheads). Bars, 5 µm.
Figure 8. A WASH- and Exo84-dependent exocytic mechanism of MT1-MMP involved in pericellular collagenolysis. (A and B) Live-cell imaging of MDA-MB-231 cells expressing MT1-MMPpHluorin (green) and mCherry-WASH (red) constructs, plated on a layer of collagen I for 30 min (see Video 6). The galleries show exocytosis of MT1-MMPpHluorin–positive endosomes (green arrowheads) in the vicinity of a collagen type I fiber (blue). These endosomes harbor WASH-positive patches (red arrowheads). The asterisks in B point the reacidification of a WASH- and MT1-MMPpHluorin–positive endosome. Time is represented in minutes. Bars, 2 µm. (C) MDA-MB-231 cells expressing MT1-MMPmCherry and MT1-MMPpHluorin silenced for the indicated proteins were seeded on a layer of type I collagen fibers and imaged over a 30-min time period and the frequency of MT1-MMPpHluorin exocytic events was quantified (box plots of events/cell/min). The number of cells analyzed for each cell population is indicated. *, P < 0.05; **, P < 0.01; ***, P < 0.001 (compared with siNT-treated cells). (D) Lifetime of MT1-MMPpHluorin–positive flashes (in minutes). Data show the number of pHluorin exocytic flashes per cell for the different lifetimes ± SEM from five cells out of three independent experiments. (E) MDA-MB-231 cells expressing MT1-MMPpHluorin were cultured on a layer of Cy3-labeled type I collagen fibers and subjected to external pH shifts alternating between 7.4 (white zones) and 5.5 (gray zones). The plots represent pHluorin intensity of a MT1-MMP–positive endosome that underwent exocytosis (circled with a solid line in the inset on the right) and of a region of the plasma membrane (dashed line). The corresponding time-lapse series is shown in Video 7. The gallery above the plot represents a sequence starting with external pH 7.4 corresponding to the circled MT1-MMPpHluorin–positive endosome (see inset). Bars: (inset) 5 µm; (gallery) 1 µm.

contact with the matrix (basement membrane or fibrous type I collagen) at the level of WASH–F-actin–cortactin– and exocyst-positive puncta. These connections, most likely tubular, that form between late endosomes and the plasma membrane allow exocytosis of MT1-MMP to the surface and they occur at and correspond to bona fide invadopodia. It is currently though that WASH-dependent actin assembly (He and Guo, 2009; Hertzog and Chavrier, 2011; this study), suggests a mechanism whereby WASH and exocyst may work in a coordinated manner to control the formation of endosome-to-plasma membrane connections ensuring MT1-MMP targeting to the invadopodial plasma membrane. In addition, the exocyst complex is known to control SNARE complex assembly and to influence vesicle fusion (Heider and Munson, 2012). Thus, the exocyst could regulate membrane fusion at invadopodial endosome-to-plasma membrane connections. Whether the exocyst can interact and influence the activity of the late endosome SNARE protein VAMP7, which is required for MT1-MMP
Finally, this study supports the conclusion that two pools of F-actin–cortactin coexist and cooperate to ensure invadopodia formation and function. One pool is associated with and depends on N-WASP activity, which is associated with the core invadopodia structure forming on the cytoplasmic face of the plasma membrane in association with the matrix. These N-WASP–F-actin–cortactin–positive structures include typical dotty-like invadopodia forming on the adherent plasma membrane in cells on gelatin or basement membrane (Yamaguchi et al., 2005; Artym et al., 2006; Oser et al., 2010; Yu et al., 2012) and linear F-actin–based structures assembling in association with collagen fibers in tumor cells invading through a 3D fibrous environment (Juin et al., 2012; Yu et al., 2012; this study). These different structures represent docking sites for MT1-MMP–containing endosomes and may provide a physical scaffold for plasma membrane protrusion and MT1-MMP retension to penetrate the extracellular matrix and basement membrane. The second pool, with a punctate organization, is associated with and depends on WASH activity on MT1-MMP–positive endosomes and, together with other components including the exocyst complex, contributes to an exocytic mechanism that fuels the invadopodia with the proteolytic activity necessary for pericellular matrix remodelling. The coordinated function of WASH and the exocyst complex in the fundamental biology of invadopodium-mediated proteolysis in different pathologically relevant matrix environments demonstrated here appears a crucial part of the invasive program of carcinoma cells.

Our findings also demonstrate that exocytosis of MT1-MMP does not occur through complete collapse of MT1-MMP–positive endosomes into the plasma membrane, but by a mechanism whereby late endosomes are retrieved apparently intact after fission of endosome-to-plasma membrane connections and can possibly engage in multiple exocytic events. Strikingly, some of these connections remain open for several minutes; however, transfer of MT1-MMP from the storage endosome to the invadopodial plasma membrane is tightly controlled as a large portion of the protease remains associated with the endosome. Thus, the tubular connection forming between the two membrane compartments in a WASH/exocyst-dependent manner may exert different functions; it could stabilize and restrict the extension of the fusion pore between the two membranes and it may also act as a filter to prevent free diffusion of MT1-MMP out of the endosome. Along this line, it was recently shown that N-WASP–dependent invadopodial F-actin can prevent the diffusion of MT1-MMP out of the invadopodia through some interaction of the cytoplasmic domain of the protease with the actin network (Yu et al., 2012). The same type of interactions with WASH-dependent F-actin surrounding the connecting tube would limit diffusion of MT1-MMP out of the endosome and ensure controlled delivery of MT1-MMP to the plasma membrane for controlled remodeling of the basement membrane and collagen fibers.

Finally, this study supports the conclusion that two pools of F-actin–cortactin coexist and cooperate to ensure invadopodia formation and function. One pool is associated with and depends on N-WASP activity, which is associated with the core invadopodia structure forming on the cytoplasmic face of the plasma membrane in association with the matrix. These N-WASP–F-actin–cortactin–positive structures include typical dotty-like invadopodia forming on the adherent plasma membrane in cells on gelatin or basement membrane (Yamaguchi et al., 2005; Artym et al., 2006; Oser et al., 2010; Yu et al., 2012). Mature invadopodia are responsible for penetrating and breaching the basement membrane in contact with carcinoma cells (1 and 2; Hotary et al., 2006; Rowe and Weiss, 2008). In the fibrous, collagen-rich extracellular membrane environment surrounding the tumor (3), fibers that oppose cell movement, involving recognition by yet unidentified collagen receptors, trigger the assembly of N-WASP– and F-actin–positive linear invadopodia and the recruitment and exocytosis of MT1-MMP–containing endosomes based on the conserved WASH and exocyst-dependent mechanism (inset iv).
Materials and methods

Antibodies and constructs

Antibodies used in this study are listed in Table S1. Full-length mouse WASH cDNA aminoterminal-terminally fused with GFP or mCherry was subcloned in pcDNA3.1 vector (Derivery et al., 2009). Full-length rat cortactin cDNA (provided by M.A. McNiven, Mayo Clinic, Rochester, MI) was subclone in pDsRed1-N1 and pEGFP-N1 (Takara Bio Inc.) in fusion with carboxy-terminal DHFR or GFP, respectively. Human full-length Exo84 and Sec3 were subcloned in pCDM6 (DHFR) plasmid with aminoterminal GFP or carboxy-terminal HA tags.

Yeast two-hybrid screening

The coding sequence for fulllength human Sec3 (GenBank accession no. gi:7023219) and Exo84 (GenBank accession no. gi:58331103) was PCR amplified and cloned into pB27 as carboxy-terminal fusions to LexA. The constructs were checked by sequencing the entire insert and used as baits to screen at saturation a highly complex human placenta domain library constructed into pPl6, pB27 and pPl6 derive from the original pBD116 (Vojet and Hollenberg, 1995) and pGADGH (Bartel et al., 1993) plasmids, respectively. For Sec3, 96 million clones (10-fold the complexity of the library) were screened using a mating approach with Y187 [matα] and L404G4α [matα] yeast strains as previously described (Fromont-Racine et al., 1997). 353 His+ colonies were selected on a medium lacking tryptophan, leucine, and histidine, supplemented with 2 mM 3-aminotriazole. For Exo84, 64 million clones were screened and 164 His+ colonies were selected using the same protocol. The prey fragments of the positive clones were amplified by PCR and sequenced at their 5′ and 3′ junctions. The resulting sequences were used to identify the corresponding interacting proteins in Genbank using a fully automated procedure. The protein interactions from this publication have been submitted to the IMEx consortium through IntAct (Kerrien et al., 2012) and assigned the identifier IMT1363.

Cell culture, DNA transfection, stable cell lines, and RNAi

The human breast adenocarcinoma cell line MDA-MB-231 (ATCC; American Type Culture Collection) was maintained in L-15 culture medium (Sigma-Aldrich) with 2 mM glutamine and 15% FCS at 37°C in 10% CO2. MDA-MB-231 cells stably expressing MT1-MMPmCherry (MT1-MMPmCh) were cultured in the same medium supplemented with 0.5 mg/ml G418 (Sakurai-Yageta et al., 2008). MDA-MB-231 cells stably expressing MT1-MMPPhluorin (Uzarraga et al., 2009), with the GFP-derivative phluorin tag (super-epichromatic variant [Miesenböck, 2012]) inserted in the extracellular domain, were generated by lentiviral transduction. For transient transfection, MDA-MB-231 cells were transfected with plasmid constructs [1 μg] using Lipofectamine 2000 (Invitrogen) or Nucleofector (Lonza) according to the manufacturers' instructions. Cells were analyzed 48 h after transfection. BT-549 (HTB-122; American Type Culture Collection) human breast cancer cells were cultured in RPMI medium with 2 mM glutamine and 10% FCS at 37°C in 5% CO2. siRNA transfection was performed using 50 nM siRNA with Lullaby reagent (Olink Bioscience). In brief, secondary antibodies were incubated in a preheated humidity chamber for 1 h at 37°C. Ligation was performed with a ligation-containing ligation solution for 30 min at 37°C. Then, the amplification step was performed with a polymerase-containing amplification solution for 1 h and 40 min at 37°C. Finally, coverslips were incubated with C3-conjugated IgGs to detect Duolink signal. Coverslips were analyzed by wide-field microscopy in WASH-Exo84 or WASH-cortactin. Duolink signal staining was manually counted for each cell and compared with control conditions in which one of the primary antibodies was omitted.

Immunoprecipitation

For expression of HA-tagged Exo84 and Sec3 exocyst complex subunits, HeLa cells were transiently transfected using FuGENE (Roche) according to the manufacturer's instructions. 48 h after transfection, cells were lysed in lysis buffer (50 mM Tris-HCl, pH 7.4, 150 mM NaCl, 1% Triton X-100, 1 mM EDTA, and complete Mini Protease Inhibitor Cocktail; Roche), and then centrifuged at 13,000 rpm for 10 min at 4°C. Supernatants (1–2 mg of total protein) were incubated with 1–2 μg of anti-WASH antibodies or control irrelevant IgGs (anti-GST) for 1 h at 4°C, and then protein G-Sepharose 4 Fast Flow (50 μl; GE Healthcare) was added and the incubation was continued for a further 2 h at 4°C. Beads were washed four times in lysis buffer and bound proteins were eluted in SDS sample buffer, separated by SDS-PAGE, and detected by immunoblotting with indicated antibodies using ECL Western Blotting Detection Reagents (GE Healthcare).

Fluorescent gelatin degradation assay

MDA-MB-231 cells were incubated for 5 h in FITC-conjugated crosslinked gelatin (Invitrogen) as described previously (Sakurai-Yageta et al., 2008), and then fixed and stained for F-actin and cortactin. Cells were imaged with the 63x objective of a wide-field microscope (DM6000 B/M; Leica) equipped with a CCD CoolSnap HQ2 camera (Roper Scientific) and steered by Metamorph (Molecular Devices). For quantification of degradation, the total area of degraded matrix in one field (black pixels) measured with the Threshold command of MetaMorph was divided by the number of phalloidin-labeled cells in the field to define a degradation index, which was normalized to the degradation index of control siNT-treated cells set to 100 as described previously (Sakurai-Yageta et al., 2008).

Indirect immunofluorescence microscopy, 3D deconvolution, and image analysis

MDA-MB-231 cells were cultured on gelatin-coated coverslips. Cells were extracted with 0.5% Triton X-100 in 4% paraformaldehyde and stained for immunofluorescence microscopy. Images were acquired with a wide-field microscope (Eclipse 90i; Nikon) using a 100x Plan Apo VC 1.4 oil objective and a highly sensitive cooled interline charge-coupled device (CCD) camera (CoolSnap HQ2; Roper Scientific). A z-dimension series of images was taken every 0.2 μm by means of a piezoelectric motor (Physik Instrumente) and the images were deconvoluted (Gibas, 2005). For quantification of the effect of Exo84 knockdown on WASH-cortactin distribution, a perinuclear region was drawn with the Region Tool of MetaMorph 7 (Molecular Devices) and the integrated intensity was measured with the Region Measurement command. For quantification of the effect of Exo84 knockdown on WASH-cortactin domains, individual WASH and cortactin subdomains were circled with MetaMorph 7 and the integrated intensity was measured with the Region Measurement command.

WASH, Exo84, and cortactin spot detection and codistribution analysis

To detect endosomes, an automatic Otus threshold is applied to the Gaussian-filtered MT1-MMP-positive endosome image (σ = 1.5 pixels). Statistics about each endosome are then saved. For each endosome, WASH and Exo84 (or WASH and cortactin) spots are searched for in a neighboring of x pixels (manually set by user) in their respective channel. Their number and position are saved per endosome (see the method in Text S2 in online supplemental material). From the position of WASH and Exo84 (or WASH and cortactin) spots around each endosome, each WASH spot is paired with its closest Exo84 (or cortactin) spot neighbor, optimized over all spots around this endosome. This allowed measuring of the distribution of distance between WASH-Exo84 (or WASH-cortactin) spots (see the codistribution analysis Zip file S3 in online supplemental material).

Duolink in situ proximity ligation assay

For Duolink assay (Söderberg et al., 2006), cells grown on coverslips coated with cross-linked gelatin were permeabilized with 0.5% Triton X-100 in 4% paraformaldehyde, fixed in 4% paraformaldehyde, and then incubated with primary antibodies: rabbit polyclonal anti-WASH, goat polyclonal anti-Exo84, and mouse monoclonal anti-cortactin. Secondary antibodies tagged with short DNA oligonucleotides were added. Hybridization, ligation, amplification, and detection were realized according to the manufacturer's protocol (Olink Bioscience). In brief, secondary antibodies were incubated in a preheated humidity chamber for 1 h at 37°C. Ligation was performed with a ligation-containing ligation solution for 30 min at 37°C. Then, the amplification step was performed with a polymerase-containing amplification solution for 1 h and 40 min at 37°C. Finally, coverslips were incubated with Cy3-conjugated IgGs to detect Duolink signal. Coverslips were analyzed by wide-field microscopy and WASH-Exo84 or WASH-cortactin. Duolink signal staining was manually counted for each cell and compared with control conditions in which one of the primary antibodies was omitted.

Live-cell spinning disc confocal microscopy and FITC-dextran pulse experiment

MDA-MB-231 cells stably expressing MT1-MMPmCh were plated on glass-bottom dishes (MatTek Corporation) coated with cross-linked gelatin and kept in a humidified atmosphere at 37°C and 1% CO2. Movement of MT1-MMPmCh (or transiently expressing Rab7-GFP or VAMP7-GFP) were plated in glass-bottom dishes coated with Alexa Fluor 594–conjugated cross-linked gelatin (or unlabeled gelatin) and a z-stack of images was acquired before adding dextran. FITC- or Alexa Fluor 647–conjugated dextran (10,000 MW) was added to the medium to 2 mg/ml final concentration (from a 20-rug/ml stock solution). After 1 min at 37°C, cells were washed and z-stack sequences were acquired.
TIRFM
MDA-MB-231 cells transfected with GFP- and mCherry-tagged proteins were plated on glass-bottom dishes coated with cross-linked unlabeled gelatin as previously described. Simultaneous dual-color TIRFM sequences were acquired on an inverted microscope (TE2000) equipped with a 100× TIRF objective (1.47 NA), a TIRF arm, an image splitter (DV; Roper Scientific) installed in front of the CCD camera and a temperature controller. GFP and m-Cherry were excited with 491- and 560-nm lasers, respectively (100 mW; Roper Scientific), both controlled for power by an acousto-optical tunable filter. Fluorescent emissions were selected with bandpass and longpass filters (Chroma Technology Corp.) and captured by a QuantEM EMCCD camera (Roper Scientific). The system was driven by Metamorph. To measure the subplasma membrane density of MT1-MMP endosomes, cells stably expressing MT1-MMPmCh and silenced for the protein of interest were plated on a layer of unlabeled gelatin and imaged by TIRFM for an exposure time of 100 ms and with 1 pixel corresponding to 160 nm. The density of MT1-MMPmCh structures was evaluated by thresholding images using ImageJ software. The threshold was set at 1.3× the cytoplasmic background. The size of the cell footprint was not affected by the siRNAs used.

High-resolution 3D structured illumination microscopy (SIM)
Image acquisition was performed in the 3D SIM mode with a N-SIM microscope (Nikon) equipped with a 100× 1.49 NA oil immersion objective, laser illumination (488 nm at 200 mW and 561 nm at 100 mW) and an EM CCD camera (DU98V; Andor Technology). Image reconstruction was performed using the NIS-Elements software (Nikon; based on Gustafsson et al. [2008]).

Matrigel and collagen I invasion assays
The Transwell Matrigel invasion assay has been previously described [Steffen et al., 2008]. Invasive migration in 3D type I collagen was assessed using the Oris Cell Invasion Assay (Platypus Technologies). H2B-EGFP–expressing MDA-MB-231 cells were reverse transfected with Lulaby reagent with MT1-MMP, WASH, or Exo84 siRNAs (50 nM). 24 h after transfection, cells were trypsinized and counted and 40,000 cells/well were seeded in the presence of Oris Cell Seeding Stoppers to restrict seeding to the outer annular regions of the wells in a 96-well plate previously coated with a bottom layer of acid-extracted rat tail tendon type I collagen (2.2 mg/ml; BD). Removal of the stoppers reveals a 2-mm diameter unseeded region in the center of each well, into which seeded cells may invade during a 48-h time period, once a collagen I overlay has been applied. H2B-EGFP nuclei images were acquired from each well at the beginning (T0) and 48 h after invasion (T2) using a cooled CCD camera (HQ2) mounted on a microscope (TE2000) equipped with a 4× objective CFI Plan Fluor 0.13 NA, 17.1 WD. Image analysis was performed using Metamorph software. The index of invasion was determined by thresholding the area occupied by H2B-EGFP nuclei in the detection zone of each well after 48 h of invasion. This index is defined by the total area occupied by H2B-EGFP nuclei in unseeded area at the end of the assay (T2) by subtracting the area occupied by H2B-EGFP nuclei at the beginning of the experiment (T0). This area of invasion was normalized and results from three independent experiments are represented.

Live-cell imaging of cells on a layer of fibrous type I collagen
Glass bottom dishes (MatTek Corporation) were layered with 100 µl of a solution of type I collagen mixed with Alexa Fluor 647–conjugated type I collagen (10% final) at a final concentration of 2.2 mg/ml (Sahai et al., 2004). After letting for 3 min at 37°C, the collagen layer was washed gently in PBS and 1 ml of the cell suspension in L15 medium with 15% FCS (1.5–2.5 × 10^5 cells/ml) was added. Cells were incubated for 30 min at 37°C in 1% CO2 before live-cell imaging using multi-color spinning disk confocal microscopy.

Quantification of pericellular collagenolysis
Cells treated with siRNAs against MT1-MMP, WASH, Exo84, or non-targeting siRNA for 48 h (Table S2) were trypsinized and resuspended in 0.2 ml of 2.2 mg/ml collagen I solution (2.5 × 10^5 cells/ml) loaded on a glass coverslip. After letting for 30 min at 37°C, complete medium was added and collagen-embedded cells were incubated for 24 h at 37°C in 1% CO2. After fixation in 4% paraformaldehyde in PBS at 37°C for 30 min, samples were incubated with anti-Col1/3/4/C antibodies (2.5 µg/ml) for 2 h at 4°C, washed extensively with PBS, and counterstained with Cy3-conjugated anti-rabbit IgG antibodies, DAPI, and Alexa Fluor 488–phalloidin to see the cell shape. Image acquisition was performed with an A1R confocal microscope (Nikon) with a 40× oil objective. Quantification of degradation spots was performed with a homemade Image macro (Zip file S1 in the online supplemental material) images were preprocessed by a laplacian of Gaussian filter (Sage et al., 2005), with variance reflecting the expected spot size. The spot detection then consists in finding the local minima; sorting them in ascending order of intensity, applying a flood-fill algorithm to each of them using a fixed noise tolerance value set up for all experiments at 10,000, and discarding higher minima whose fill regions touch those of lower minima. Detected spots are then counted and saved for visual verification. No manual correction was done. Degradation index is the number of degradation spots divided by the number of cells present in the field, normalized to the degradation index of control cells to 100.

MT1-MMP/PHuorin exocytosis on collagen type I fibers
MDA-MB-231 cells stably coexpressing MT1-MMP-Cherry and MT1-MMP/PHuorin constructs were treated with nontargeting, WASH, or Exo84 siRNA for 72 h and plated atop of a drop of polymerized type I collagen. Cells were imaged by confocal spinning disk microscopy. The number of exocytic events of MT1-MMP/PHuorin (i.e., GFP flashes) was measured per minute and per cell. As a control we measured the mean fluorescence intensity of MT1-MMP-Cherry for each cell.

Extracellular pH shifts experiment
MDA-MB-231 cells stably expressing MT1-MMP/PHuorin were plated on polymerized type I collagen conjugated with Cy3. Cells were imaged on a microscope (IX71; Olympus) with a 100×, 1.49 NA objective, a frame rate of 0.5 Hz with a QuantEM EMCCD camera. Cells were bathed in a solution containing 140 mM NaCl, 2 mM KCl, 2 mM MgCl2, 2 mM CaCl2, 10 mM Hepes, and 5 mM d-glucose, and adjusted to pH 7.4 and 315 mosm. For the solution at pH 5.5, Hepes was replaced by MES. Solution exchange was performed as described previously (Perrais et al., 2004).

Statistical analyses
Statistical analyses were performed using analysis of variance or Student’s t test in Prism software.

Online supplemental material
Fig. S1 shows additional data on markers associated with MT1-MMP–positive endosomes in MDA-MB-231 and BT-549 cells. Fig. S2 shows levels of proteins in cells targeted by different siRNAs. Fig. S3 shows additional staining of the basal and invadopodial markers. Fig. S4 shows a gallery of images taken from a time-lapse sequence of cells expressing MT1-MMPmCh and MT1-MMP/PHuorin. Tables S1 and S2 list antibodies and show their closest neighbored cortactin spots (or Exo84 spots) for each event of MT1-MMP–containing endosomes occurring near collagen fibers. Table S3 shows the fluorescence index of collagen. Text file S2 is the macro used to detect endosomes in MDA-MB-231 and BT-549 cells.

Degradation index
The threshold was set at 1.3× the cytoplasmic background. The size of the cell footprint was not affected by the siRNAs used.
References


