Mitochondrial Targeting of the Enteropathogenic *Escherichia coli* Map Triggers Calcium Mobilization, ADAM10-MAP Kinase Signaling, and Host Cell Apoptosis

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ABSTRACT The ability of diarrheagenic bacterial pathogens, such as enteropathogenic *Escherichia coli* (EPEC), to modulate the activity of mitogen-activated protein kinases (MAPKs) and cell survival has been suggested to benefit bacterial colonization and infection. However, our understanding of the mechanisms by which EPEC modulate these functions is incomplete. In this study, we show that the EPEC type III secreted effector Map stimulates the sheddase activity of the disintegrin and metalloproteinase domain-containing protein 10 (ADAM10) and the ERK and p38 MAPK signaling cascades. Remarkably, all these activities were dependent upon the ability of Map to target host mitochondria, mainly via its mitochondrial toxicity region (MTR). Map targeting of mitochondria disrupted the mitochondrial membrane potential, causing extrusion of mitochondrial Ca\(^{2+}\) into the host cell cytoplasm. We also found that Map targeting of mitochondria is essential for triggering host cell apoptosis. Based on these findings, we propose a model whereby Map imported into mitochondria causes mitochondrial dysfunction and Ca\(^{2+}\) efflux into the host cytoplasm. Since Ca\(^{2+}\) has been reported to promote ADAM10 activation, the acute elevation of Ca\(^{2+}\) in the cytoplasm may stimulate the ADAM10 sheddase activity, resulting in the release of epidermal growth factors that stimulate the ERK signaling cascade. As p38 activity is also Ca\(^{2+}\) sensitive, elevation in cytoplasmic Ca\(^{2+}\) may independently also activate p38. We hypothesize that Map-dependent MAPK activation, combined with Map-mediated mitochondrial dysfunction, evokes mitochondrial host cell apoptosis, potentially contributing to EPEC colonization and infection of the gut.

IMPORTANCE Enteropathogenic *E. coli* (EPEC) is an important human diarrhea-causing bacterium. The pathogenic effects of EPEC largely depend upon its ability to inject a series of proteins, termed effectors, into the host cells. One such effector is the mitochondrion-associated protein (Map). Map has been shown to induce actin-rich projections (i.e., filopodia) on the infected cell surface and activate a Rho GTPase enzyme termed Cdc42. Nonetheless, although most injected Map localizes to host mitochondria, its functions in the mitochondria remain unknown. Here, we show that Map targeting of mitochondria stimulates the disruption of mitochondrial membrane potential to induce Ca\(^{2+}\) efflux into the host cytoplasm. The efflux stimulates the activity of a protein termed ADAM10, which induces activation of a mitogen-activated protein kinase cascade leading to host cell apoptosis. As apoptosis plays a central role in host-pathogen interactions,
our findings provide novel insights into the functions of mitochondrial Map in promoting the EPEC disease.

KEYWORDS ADAM10, calcium, enteropathogenic E. coli, MAP kinases, Map effector, protein effectors, apoptosis, mitochondria

Enteropathogenic Escherichia coli (EPEC) is a human-specific bacterial pathogen that infects the enterocytes of the small intestine. EPEC infection causes acute and persistent diarrhea, mainly in children worldwide (1, 2). The virulence of EPEC is primarily due to the ability of the microbe to activate a type III secretion system (T3SS) that injects dozens of effector proteins from the bacterial cytoplasm into the host cells (3). The translocated effectors intoxicate the infected cells by hijacking and subverting diverse organelles, cytoskeletal elements, and signaling processes (4, 5). Analysis of the precise mechanisms by which these effectors perform their functions is crucial for better understanding the EPEC disease and for designing improved therapeutics.

Mitogen-activated protein kinases (MAPKs) are involved in the regulation of cell proliferation, survival, differentiation, stress response, and programmed cell death (i.e., apoptosis) (6–8). We recently showed that EspH, an EPEC type III secreted effector implicated in actin cytoskeleton remodeling (9–11) and the inhibition of Rho GTPases (10, 12), also suppresses the MAPK/extracellular signal-regulated kinases 1/2 (ERK1/2) signaling pathway at longer infection times (13). Previous studies have indicated that EPEC can rapidly stimulate the MAPK/ERK1/2 signal transduction pathway and that this T3SS-dependent event may play a role in the inflammatory response and infection, but not in tight-junction barrier disruption (14–16). However, the identity and mode of action of type III secreted components that mediate ERK1/2 activation have not been explored. Here, we provide evidence that the type III secreted effector protein mitochondrion-associated protein (Map) activates the MAPK/ERK1/2 signaling pathway at an early infection phase. Map has been previously characterized to target mitochondria by a mitochondrial targeting signal (MTS) (17, 18), activate the Rho GTPase Cdc42 by a WxxxE guanine nucleotide exchange factor (GEF) family motif (19, 20), and interact with host proteins through a C-terminal TRL PDZ class I binding motif (21). Here, we show that Map stimulates the MAPK/ERK signaling pathway by activating the sheddase activity of the disintegrin and metalloproteinase domain-containing protein 10 (ADAM10). We linked these effects to the ability of Map to target mitochondria and evoke Ca²⁺ efflux from them into the host cell cytoplasm and to the induction of host apoptosis. We hypothesize that the triggering of the ADAM10-MAPK/ERK signaling by Map in an early infection event, which is counteracted by EspH in a later infection time, may play a crucial role in the induction of EPEC pathogenesis.

RESULTS

T3SS-dependent increase in pERK levels at an early infection stage. We showed previously that active phosphorylated ERK (pERK) levels are suppressed upon prolonged (90-min) infection with EPEC and that the EspH effector mediates this effect (13). Here, we examined the effect that EPEC infection at a shorter time (30 min) may have on ERK phosphorylation (pERK). HeLa (Fig. 1A) or Caco-2 (Fig. 1B) cells were infected with EPEC-wt or with the type III secretion-deficient mutant EPEC-escV for 30 min at 37°C or were left uninfected. Cells were then lysed, and the levels of pERK were evaluated by immunoblotting. The results show a significant increase in pERK levels (mainly pERK2) in response to EPEC-wt compared to EPEC-escV-infected or uninfected cells. Infection with EPEC-escV had no significant impact on pERK levels compared to uninfected cells. These data suggest that type III secreted components increase pERK levels in epithelial cells at an early infection phase.

Translocated Map induces ERK phosphorylation. Our next goal was to identify type III secreted effectors that could mediate the stimulatory effect. To this end, we examined the capacity of EPEC strains mutated at the different locus of enterocyte
effacement (LEE)-located effector genes to upregulate pERK in HeLa cells. The results identified EPEC-
map as the only mutant strain that did not demonstrate such capability (Fig. 2A). Notably, pERK levels in EPEC-
map-infected cells were even lower than those exhibited by EPEC-
escV-infected cells. This finding suggests that in the absence of Map, other bacterial components suppress ERK. We suspected that the injected EspH, a negative regulator of ERK (13), could be such a component. To address this prediction, pERK levels in HeLa cells infected with an EPEC strain doubly mutated in the espH and map genes (EPEC-
map, espH) were examined. Indeed, data presented in Fig. 2B shows that pERK levels in cells infected with the EPEC-
map, espH mutant strain were higher than in the EPEC-
map-infected cells, reaching levels comparable to those exhibited by EPEC-
escV-infected or uninfected cells. These results point to EspH as the effector that suppresses pERK in the EPEC-
map-infected cells. Complementing EPEC-
map with a Map-encoding plasmid increased pERK levels, which reached the levels observed in EPEC-wt-infected cells (Fig. 2B). A similar Map-dependent stimulatory effect, albeit somewhat lower, was observed in infected Caco-2 cells (Fig. 2C). Notably, in these experiments, Map was effectively translocated into the cells (Fig. S1A). Since the host actin cytoskeleton can be affected by bacterial targeted MAPKs (22), and since Map itself can modulate filamentous actin (F-actin) (10, 19, 23), the F-actin levels in EPEC-
map/H11001 Mapwt infection sites were compared to those of EPEC-
map-infected cells. The results showed that the F-actin clustering levels were comparable at the infection sites of the two EPEC strains (Fig. S1B). These results argue that translocated Map is the type III secreted effector that confers pERK stimulation at an early infection time and that the effect is not contributed by changes in F-actin clustering at infection sites.

Map triggers phosphorylation of MEK, p38, and the EGFR but not B-Raf or C-Raf. Next, we asked what the upstream components that activate ERK upon stimulation by Map are. We found that the immediate upstream MEK (Fig. 3A), but not C-Raf (Fig. 3B and C) or B-Raf (Fig. 3D), is hyperphosphorylated in response to EPEC-
map+Mapinfected infection. Interestingly, pEGFR (Y1068), positioned upstream of MEK, was also upregulated (Fig. 3E), suggesting that translocated Map activates the epidermal

FIG 1 EPEC infection increases pERK levels at an early infection phase and in a T3SS-dependent manner. pERK levels were examined in HeLa (A) and Caco-2 (B) cells. Cells were infected with the indicated EPEC strains (Table S1) for 30 min at 37°C or remained uninfected. Cells were then lysed, and pERK and general (g)ERK levels were determined by SDS-PAGE followed by Western blotting, as described in Materials and Methods. Representative Western blots are shown. Results are means and SE from 3 independent experiments. ***, P < 0.0005; **, P < 0.005; ns, nonsignificant (P > 0.05).
growth factor receptor (EGFR)/MEK/ERK signaling pathway, via a Raf-independent mechanism.

Map activates the Rho GTPase Cdc42 (19, 20). Cdc42 and Rac1 can act as upstream mediators of p38 and JNK MAPK activities in response to stress or inflammatory stimuli (24). Therefore, we hypothesized that Map may stimulate the phosphorylation and activation of p38. Data in Fig. 3F indeed show a Map-dependent increase in phosphorylated-p38 (p-p38) levels. pJNK was also stimulated in response to EPEC-wt, but the clear involvement of Map in the process could not be demonstrated (data not shown). In summary, our data link Map to the stimulation of pEGFR, pMEK, pERK, and p-p38 but not to pRaf signaling.

**Map mediates ADAM10 but not ADAM17 activation.** ADAM17 acts as a sheddase of epiregulin, transforming growth factor α (TGF-α), amphiregulin, and heparin-binding EGF-like growth factor (25). Since these ligands are known to potentiate the EGFR/ERK/MAPK pathway (26), we reasoned that Map might activate this pathway by stimulating ADAM17. Data in Fig. 4A suggest that this is not the case. EPEC infection activates the

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**FIG 2** The type III secreted effector Map triggers an increase in pERK levels in HeLa and Caco-2 cells. (A) Screening for effects on pERK levels by EPEC strains mutated at different LEE-borne effector genes. HeLa cells were infected with the indicated mutant strains (Table S1), and pERK levels were analyzed, as for Fig. 1. (B and C) Map<sub>wt</sub> translocation induces pERK in HeLa and CaCo-2 cells, and EspH counteracts this effect. Cells were infected with the indicated EPEC strains (Table S1), and pERK levels in the infected cells were analyzed, as for Fig. 1. Results are means and SE from 3 independent experiments. ***: P<0.0005; **: P<0.005; *: P<0.05; ns, nonsignificant (P>0.05).
ADAM17-mediated TGF-α shedding activity, but the process is T3SS and Map independent. EGF and betacellulin (BTC) share overlapping signaling properties concerning EGFR activation (27). Because the metalloprotease ADAM10 acts as the main sheddase of BTC and EGF (25), we hypothesized that Map might be capable of activating the ADAM10 sheddase activity. Results presented in Fig. 4B show that cell infection with EPEC-wt or EPEC-map/H11001 Mapwt indeed resulted in increased BTC shedding compared to EPEC-map or EPEC-escV infection or no infection. These data suggest that Map stimulates the ADAM10 sheddase activity. To further confirm this conclusion, we examined the ability of EPEC to upregulate pERK in cells whose ADAM10 expression was silenced by small interfering RNA (siRNA) (Fig. S1C). Data in Fig. 4C show that cell infection with EPEC-map/H11001 Mapwt failed to increase pERK levels in the ADAM10 siRNA-treated cells, while the stimulatory effect was apparent in the control siRNA-treated cells. Notably, Map translocation was not affected by ADAM10 siRNA treatment (Fig. S1D). These results suggest that the Map-mediated stimulation of pERK is due to its ability to activate ADAM10.

The GEF and PDZ binding motifs of Map are not involved in ADAM10-EGFR-MAPK activation. As previously noted, Map consists of an MTS, a WxxxE Rho GEF, and a C-terminal TRL class I PDZ binding motif. To investigate which of these motifs mediate ERK activation, we generated EPEC mutant strains bearing a mutation in the WxxE or GEF and PDZ binding motifs of Map are not involved in ADAM10-EGFR-MAPK activation. As previously noted, Map consists of an MTS, a WxxxE Rho GEF, and a C-terminal TRL class I PDZ binding motif. To investigate which of these motifs mediate ERK activation, we generated EPEC mutant strains bearing a mutation in the WxxE or GEF and PDZ binding motifs of Map are not involved in ADAM10-EGFR-MAPK activation. As previously noted, Map consists of an MTS, a WxxxE Rho GEF, and a C-terminal TRL class I PDZ binding motif. To investigate which of these motifs mediate ERK activation, we generated EPEC mutant strains bearing a mutation in the WxxE or GEF and PDZ binding motifs of Map are not involved in ADAM10-EGFR-MAPK activation. As previously noted, Map consists of an MTS, a WxxxE Rho GEF, and a C-terminal TRL class I PDZ binding motif. To investigate which of these motifs mediate ERK activation, we generated EPEC mutant strains bearing a mutation in the WxxE or GEF and PDZ binding motifs of Map are not involved in ADAM10-EGFR-MAPK activation. As previously noted, Map consists of an MTS, a WxxxE Rho GEF, and a C-terminal TRL class I PDZ binding motif. To investigate which of these motifs mediate ERK activation, we generated EPEC mutant strains bearing a mutation in the WxxE or GEF and PDZ binding motifs of Map are not involved in ADAM10-EGFR-MAPK activation. As previously noted, Map consists of an MTS, a WxxxE Rho GEF, and a C-terminal TRL class I PDZ binding motif. To investigate which of these motifs mediate ERK activation, we generated EPEC mutant strains bearing a mutation in the WxxE or

FIG 3 Map activates a MAPK signaling cascade upstream of ERK. HeLa cells were infected with wild-type, escV, map, or map+Mapwt EPEC strains for 30 min at 37°C. Cells were lysed and subjected to SDS-PAGE followed by Western blotting analyses and probed with the indicated anti-phosphorylated (p) and general (g) MEK (A), C-Raf (B–C), B-Raf (D), EGFR (E), and p38 (F) antibodies (Table S2). The levels of phosphorylated and total proteins were determined as described for Fig. 1. Results are means and SE from 3 independent experiments. Notably, we used three different anti-phospho-Raf antibodies to strengthen the conclusion that Map stimulates the EGFR-MAPK pathway in a Raf-independent mechanism. The anti-pRaf antibodies used to detect Raf kinase stimulation linked to the EGFR signaling were the classical anti-c-Raf phosphorylated on Ser338 and the anti-B-Raf phosphorylated on Ser445, a site in B-Raf shown to be equivalent to that in c-Raf (Ser338) (68), and anti-pc-Raf phosphorylated on Ser289, Ser296, and Ser301. These serine residues of c-Raf have been reported to be EGF induced (69). *** P < 0.0005; ** P < 0.005; ns, nonsignificant (P > 0.05).
The capacity of these strains to alter the pERK levels in infected HeLa cells was evaluated, as before. pERK levels in EPEC-map/H11001MapWxxxA and EPEC-map/H11001MapΔTRL caused significant upregulation of pERK levels compared to those in EPEC-map-infected cells (Fig. 5B). A similar effect was observed on pMEK (data not shown), pEGFR (Fig. 5C), and ADAM10 BTC sheddase activity (Fig. 5D). Compared to Mapwt, the capacity of the mutant Maps to undergo translocation into the host cells was somewhat reduced, especially in the case of Map/H9004TRL (Fig. 5E). This may explain the slightly reduced levels of pERK and pEGFR levels in cells infected with these EPEC strains compared to levels in cells infected with the Mapwt strain. However, the amounts of the translocated effectors seemed to be sufficient for inducing the indicated effects. Notably, previous studies suggested that F-actin-rich filopodia are not generated in response to translocated Map/H9004TRL (19, 21). MapWxxxA also fails to evoke Cdc42 and filopodium formation (Fig. S1E) (19). These data suggest that Map activates the ADAM10-MAPK and the Rho GTPase signaling pathways by two distinct and independent mechanisms.

Finally, in this context, ectopic expression of Map tagged in its C terminus with eGFP (Map-eGFP) (Fig. S1F) resulted in an increase in pERK levels compared to eGFP-expressing cells (Fig. S1G). The fusion protein colocalized with the mitochondrial marker Hsp60 (Fig. S1H). In contrast, ectopic expression of N-terminal mCherry-tagged
Map did not colocalize with Hsp60 (Fig. S1H) and did not affect pERK levels (Fig. S1G) compared to mCherry expression. These results suggest that the expression of Map is sufficient for stimulating the ADAM10-MAPK signaling cascade and that targeting of mitochondria by the effector may play a role in the process.

ADAM10 and MAPK activation requires mitochondrial targeting of Map. Next, we examined the importance of Map's mitochondrial targeting in the upregulation of the ADAM10-pERK axis. The N-terminal ~25-amino-acid sequences of bacterial type III effectors typically contain a signal that mediates their specific secretion via the T3SS (28). Thus, the MTS of Map, which was localized to the N-terminal 44-amino-acid segment of the protein (18), likely contains the signal that mediates its translocation via the T3SS. Therefore, the mere deletion of the MTS (MapΔMTS) would not be a reasonable approach for examining its role in signaling, as such mutation may affect the translocation of the effector into the host. To address this potential problem, the N-terminal 25-amino-acid type III secretion signal of EspH, an effector whose localization upon translocation has been reported to be confined to the bacterial infection sites and not to mitochondria (9), was fused to the N terminus of MapΔMTS (MapΔMTS-EspH1–25) (Tables S1 and S3; Fig. 6A). This modification was expected to facilitate the translocation of the MTS deleted Map mutant. Data presented in Fig. 6B indeed show that the translocation...
efficacy of the chimeric effector was comparable to that of Map\textsubscript{wt}. The translocated Map\textsubscript{ΔMTS-EspH\textsubscript{1–25}} also showed a wide distribution within the infected cells, albeit with a somewhat reduced colocalization with the mitochondrial marker Hsp60, compared to Map\textsubscript{wt} (Fig. 6C). Cell infection with EPEC-map+Map\textsubscript{ΔMTS-EspH\textsubscript{1–25}} caused diminished pERK (Fig. 6D), pMEK (data not shown), pEGFR (Fig. 6E), and ADAM10 (Fig. 6F) activity, reaching intermediate values between those attained upon infection with EPEC-map and EPEC-map+Map\textsubscript{wt}. These results signify the importance of Map targeting of mitochondria in evoking the ADAM10-EGFR-MAPK signaling. Nevertheless, the observation that the majority of the chimeric effector protein was localized to mitochondria raised the possibility that Map harbors another motif that functions in mitochondria in evoking the signaling effect. Previous studies have implicated amino acids 101 to 152 of Map (also called the mitochondrial toxicity region [MTR] [4]) in altering mitochondrial morphology (18). To explore the significance of this motif, an EPEC-map+Map\textsubscript{Δ101–152} mutant strain was generated (Tables S1 and S3; Fig. 6A). Map\textsubscript{Δ101–152} was effectively translocated into the infected cells (Fig. 6B and localized to mitochondria (Fig. S1I). Cell infection with EPEC-map+Map\textsubscript{Δ101–152} resulted in a significant reduction in pMEK (data...
not shown), pERK (Fig. 6D), pEGFR (Fig. 6E), and ADAM10 activity (Fig. 6F), reaching the minimal values exhibited upon infection with EPEC-map. These data suggest that a Map MTR motif may be involved in promoting the ADAM10-MAPK signaling pathway.

**Map promotes Ca\(^{2+}\) mobilization from mitochondria.** Studies have shown that Map can disrupt the mitochondrial membrane potential (18, 29). We confirmed this in our experimental system by showing that cell infection with EPEC-map\(^{+}\)Map\(^{+}\) resulted in a continuous reduction in the fluorescence of the mitochondrial membrane potential reporter tetramethylrhodamine ethyl ester (TMRE). Only a moderate reduction in the TMRE fluorescent signal was observed in response to EPEC-map infection (Fig. 7A; Movie S1). The disruption of mitochondrial membrane potential can lead to Ca\(^{2+}\) efflux from mitochondria (reviewed in reference 30). Using the mitochondrial Ca\(^{2+}\)-sensitive fluorescent reporter Rhod-2, we showed that cell infection with EPEC-map\(^{+}\)Map\(^{+}\) resulted in a sharp loss of mitochondrial Ca\(^{2+}\) levels, while infection with EPEC-map\(^{+}\)Map\(^{+}\)Map\(^{\Delta 101-152}\) led to a slight reduction in mitochondrial Ca\(^{2+}\) compared to EPEC-map. Cell infection with EPEC-map\(^{+}\)Map\(^{\Delta MTS}\)-EspH\(_{1-25}\) resulted in an intermediate effect (Fig. 7B; Movie S2). The efflux of mitochondrial Ca\(^{2+}\) may cause a parallel increase

**FIG 7** Map and EspF effects on mitochondrial membrane potential disruption and Ca\(^{2+}\) release into the cytoplasm. HeLa cells were labeled with the mitochondrial membrane potential indicator TMRE (A), the mitochondrial Ca\(^{2+}\) reporter Rhod-2 (B), or the fluorescent cytoplasmic Ca\(^{2+}\) label Fluo-8 (C), as described in Materials and Methods. Cells were then washed, infected with the indicated EPEC strains, and subjected to time-lapse confocal imaging (Movies S1 to S3). Representative images are shown. The fluorescence level of each marker was quantified at each time point and normalized to the fluorescence levels determined before bacterial exposure (0 min). Results are means ± SE for 30 cells recorded in 3 independent experiments. Bar = 2 μm.
in cytoplasmic Ca\(^{2+}\). Indeed, cytosolic Ca\(^{2+}\) imaging by the Fluo-8 probe showed a parallel increase in cytoplasmic Ca\(^{2+}\) levels that was essentially spread throughout the cytoplasm in response to EPEC-map\(^{+}\)Map\(^{wt}\) but not to EPEC-map or EPEC-map\(+\)Map\(^{A101-152}\) infection. Infection with EPEC-map\(+\)Map\(^{AMTS}\)EspH\(_{1-25}\) resulted in a slight increase in cytoplasmic Ca\(^{2+}\) (Fig. 7C; Movie S3). Taken together, these data suggest that mitochondrially targeted Map disrupts the mitochondrial membrane potential, leading to Ca\(^{2+}\) efflux from the mitochondria and Ca\(^{2+}\) influx into the infected cell cytoplasm. The MTR motif of MAP seems to play a major role in this effect.

**Mitochondrial membrane potential disruption by CCCP induces Ca\(^{2+}\) extrusion and ADAM10/pERK activation.** Previous studies suggest that Ca\(^{2+}\) can trigger ADAM10 (31–35) as well as MEK, ERK, and p38 activity (36–38). Based on these studies, we hypothesized that enrichment of the host cytoplasm with Ca\(^{2+}\) might prompt ADAM10 and MAPK activation. To tackle this hypothesis, HeLa cells were treated with the protonophore carbonyl cyanide \(m\)-chlorophenylhydrazone (CCCP), which results in the dissipation of the mitochondrial membrane potential, allowing the mitochondria to release accumulated Ca\(^{2+}\) and preventing its further accumulation in this organelle (39). Our results show that CCCP treatment resulted in a significant loss of mitochondrial Ca\(^{2+}\) (Fig. 8A; Movie S4), and a parallel increase in cytosolic Ca\(^{2+}\) (Fig. 8B; Movie S5). The activity of ADAM10 (Fig. 8C) and the levels of pERK (Fig. 8D) significantly increased in response to HeLa cell treatment with CCCP. These data, combined with data presented in Fig. 5 and 6, support the hypothesis that ADAM10 and the downstream ERK cascade are activated by Map-mediated Ca\(^{2+}\) extrusion from mitochondria and the consequent Ca\(^{2+}\) increase in the host cell cytoplasm.

**Map induces apoptosis.** Activation of the ERK cascade is linked to cell survival and death (40, 41). Mitochondrial dysfunction, membrane potential perturbation, fragmentation (fission), and redistribution have all been shown to trigger cell death (30). Data have also linked Ca\(^{2+}\) signaling, mitochondria, and various forms of cell death (42, 43). Translocated Map seems to impact Ca\(^{2+}\) compartmentalization (Fig. 7B and C), and active ERK (Fig. 6D), EGFR (Fig. 6E), and ADAM10 (Fig. 6F) levels in a mitochondrial targeting-dependent manner. We, therefore, postulated that these Map-mitochondria dependent processes could promote host cell death. Data in Fig. 9 unambiguously show that cell infection with EPEC-map\(+\)Map\(^{wt}\) resulted in a significant increase in the appearance of apoptotic cells compared to cells infected with EPEC-map or uninfected cells. This phenomenon was dependent on the ability of Map to target mitochondria, as apoptotic-cell levels in EPEC-map\(+\)Map\(^{AMTS}\)EspH\(_{1-25}\) and EPEC-map\(+\)Map\(^{A101-152}\) infected cells were significantly reduced compared to those in EPEC-Map\(^{wt}\)-infected cells.

**DISCUSSION**

Map was shown to target host cell mitochondria, in vitro and in vivo (17, 18, 29). Map has also been suggested to play a significant role in bacterial colonization and the induction of the diarrheal effect (21, 29, 44, 45). Despite its importance as a virulent factor and the fact that most Map localizes at steady state in mitochondria, the functional significance of Map-mitochondrion interactions is not known. Here, we demonstrate, for the first time, that early in infection, mitochondrial targeting of Map, particularly via its MTR motif, plays a role in activating the sheddase activity of ADAM10 (Fig. 5F and 7E). This activity may, in turn, induce the pEGFR (Fig. 3E, 5C, and 6E) and possibly another EGFR family member, ErbB4, because BTC has been shown to activate both the EGFR and ErbB4 (46). The activation of the EGFR may then elicit the MEK/ERK/MAPK signaling cascade (Fig. 3A, 5B, and 6D). Our data also suggest that Map can activate p38 (Fig. 3F). Previous studies suggested that EPEC and similarly, enterohemorrhagic E. coli (47) rapidly induce T3SS-dependent ERK, p38, and JNK in infected cells (15, 16). Here, we propose that Map is the effector that activates ERK and p38. All in all, these effects of Map do not depend on the ability of the effector to activate Cdc42 and induce actin-rich filopodia, because the mutant Map\(^{V000A}\), which fails to induce Cdc42 (Fig. S1E), retains the capacity to induce the ADAM10-MAPK signaling (Fig. 5). Thus,
early upon translocation, Map localizes to infection sites, but not to mitochondria, where it exerts Rho GEF signaling to remodel the host actin cytoskeleton. In parallel, the effector fraction targeting mitochondria evokes the ADAM10-MAPK signaling, independently.

We also show that Map targeting of mitochondria mediates the disruption of the mitochondrial membrane potential and Ca\(^{2+}\) release from mitochondria into the host cytoplasm (Fig. 7). Evidence suggests that the activity of ADAM10, MEK, ERK, and p38 is regulated by Ca\(^{2+}\) (31–33, 37, 48, 49). Thus, a Map-mediated increase in cytoplasmic Ca\(^{2+}\) could have contributed to the activation of these enzymes. This hypothesis may also be supported by the observation that disruption of mitochondrial membrane potential by the protonophore CCCP increases cytoplasmic Ca\(^{2+}\) levels and the ADAM10 and ERK activities (Fig. 8). A proposed model for the action of Map is presented in Fig. 10.

FIG 8 Disruption of mitochondrial membrane potential by the protonophore CCCP causes mitochondrial Ca\(^{2+}\) extrusion and an increase in cytoplasmic Ca\(^{2+}\) levels leading to ADAM10 and pERK activation. (A and B) Monitoring mitochondrial and cytoplasmic Ca\(^{2+}\) reporters, as described in Materials and Methods. Cells were then exposed to DMSO or the mitochondrial uncoupler CCCP and immediately subjected to time-lapse confocal imaging (Movies S4 and S5), as described in Materials and Methods. Representative images are shown in the upper panels and quantified fluorescence levels in the lower panels. The fluorescence level of each marker was quantified at each time point and normalized to the fluorescence levels determined before the addition of DMSO or CCCP. Results are means ± SE for 10 cells analyzed in 3 independent experiments. Bar = 2 μm. (C and D) Effects of CCCP on host ADAM10 sheddase activity and pERK levels. HeLa cells were treated with either DMSO or CCCP and then subjected to the ADAM10 sheddase assay or pERK/gERK analyses. Results are means and SE from 3 independent experiments. **, \(P < 0.005\).
How does the activation of ADAM10-ERK and p38 relate to EPEC pathogenesis? MAPKs can act as a double-edged sword in promoting cell death and survival (50). In most cases, ERK signaling promotes cell survival, e.g., by activating prosurvival BCL2 proteins and repressing prodeath proteins (BAD, BIM, BMF, and PUMA). However, active ERK, under certain conditions, such as nutrient deprivation, can also drive the expression of the prodeath proteins to control autophagy or apoptosis (reviewed in references 51 and 52). Similarly, previous studies suggested that activation of p38 is required for inducing the mitochondrial death pathway (53–55). Hence, it is possible that by targeting mitochondria, Map induces host cell deprivation, which, combined with ERK and p38 activation, triggers host cell apoptosis (Fig. 9).

Similar to Map, the type III secreted effector EspF targets host mitochondria via an N-terminal targeting signal to cause mitochondrial dysfunction and cell death (56–58). However, in our screen, EspF was not identified as an effector that stimulates pERK (Fig. 2A). This concern was investigated by generating new EPEC strains in which EspF was expressed in the background of EPEC-map, EPEC-map + MapΔer, EPEC-map + MapMTS-EspH1-25, and EPEC-map + MapΔ101-152-infected cells were 8%, 24%, 11%, 16%, and 10%, respectively. The fraction of apoptotic cells in EPEC-espF-infected cells, which was greater than 50% and ~6-fold higher than the fraction of apoptotic cells in uninfected cells (inset), served as a positive control for these experiments, as previous studies have shown that an EPEC-espF mutant increases dramatically the levels of apoptotic and necrotic cells (70). The fractions of cells that were dually labeled with annexin V and PI were negligible (~6%) in all experiments. Results are means and SE from 3 independent experiments. ***, P < 0.0005; *, P < 0.05; ns, nonsignificant (P > 0.05).

FIG 9 Map induces host cell apoptosis. HeLa cells infected with the indicated EPEC strains were subjected to a flow cytometry-based apoptosis assay, as described in Materials and Methods. The fraction of EPEC-exposed apoptotic cells (i.e., annexin V-positive and PI-positive cells) was normalized to the fraction of apoptotic cells detected in uninfected cells. The fractions of these cells in uninfected cells and in EPEC-map-, EPEC-map + MapΔer-, EPEC-map + MapMTS-EspH1-25-, and EPEC-map + MapΔ101-152-infected cells were 8%, 24%, 11%, 16%, and 10%, respectively. The fraction of apoptotic cells in EPEC-espF-infected cells, which was greater than 50% and ~6-fold higher than the fraction of apoptotic cells in uninfected cells (inset), served as a positive control for these experiments, as previous studies have shown that an EPEC-espF mutant increases dramatically the levels of apoptotic and necrotic cells (70). The fractions of cells that were dually labeled with annexin V and PI were negligible (~6%) in all experiments. Results are means and SE from 3 independent experiments. ***, P < 0.0005; *, P < 0.05; ns, nonsignificant (P > 0.05).
translocated and efficiently targets mitochondria, where it initiates the cascade of events that leads to pERK activation. In contrast, infection with EPEC-map allows EspF translocation, which only partially targets mitochondria and thereby exhibits a limited capacity to induce pERK. Additionally, Map's MTR motif, which we show here to be critical for promoting ADAM10-MAPK signaling, does not exist in EspF (18). Nevertheless, at a later infection time, EspF may reach the host mitochondria more effectively to promote cell death that is superimposed on the process that has been started by Map. By targeting mitochondria, Map induces an apoptotic effect early upon infection. However, EspH may counteract this effect at a later infection time (13). These opposing effects may delay the enforcement of the host cell death and therefore be required for establishing bacterial colonization and successful infection. The early Map-dependent effects could be physiologically important, because in vivo infection studies suggested that Map expression is suppressed at an early colonization phase, and when Map expression is reduced, EPEC colonization is also significantly reduced (44). Thus, the downregulation of Map could be another mechanism that terminates its effects. Many intestinal microbial pathogens modulate mitochondrial cell death to achieve successful infection (60). Hence, mitochondrial targeting, e.g., by Map, may represent a universal event that plays a role in establishing bacterial pathogens' colonization of the gut. The mechanisms underlying these effects are presently unknown. One intriguing hypothesis is that by causing mitochondrial dysfunction, pathogens build a niche where they succeed in competing with nearby commensal bacteria (microbiota) whose viability depends upon functional mitochondria of the host (61).

The precise mechanism by which Map evokes ADAM10 activation through mitochondrial targeting is still not well understood. Recent studies have shown that membrane externalized phosphatidylserine is required for activating the ADAM10 sheddase activity (62). Hence, an interesting hypothesis would be that the induction of mitochondrial cell death by Map is caused by phosphatidylserine externalization. This process is shown here to take place because the detection of cell apoptosis utilizes the annexin V reporter, which binds specifically to externalized phosphatidylserine (Fig. 9).
Finally, most recent findings may signify the importance of ERK in EPEC infection by showing that the effector NleD selectively cleaves several MAPKs but not ERK (63), probably leaving it intact for activation by Map.

**MATERIALS AND METHODS**

**Bacterial strains, antibodies, primers, plasmids, and site-directed mutagenesis.** Bacterial strains, antibiotics, primers, and plasmids used in this study are listed in Tables S1 to S4, respectively. For constructing the EPEC-espH::map double mutant strain (Table S1), λ Red recombinase system (64) was used to mutate map in EPEC-espH::Kan strains. The upstream and downstream recombination sequences were PCR amplified from the genomic DNA of EPEC-wt using primer pairs 1371-4197 and 1495-4198, respectively (Table S4). Primers 1354 and 1355 (Table S4) were used to amplify a chloramphenicol cassette from pKD3 (Table S3). DNA of the map::Cam allele was made by isothermal assembly of these three PCR fragments (65), followed by electroporation into the EPEC espH::Kan strain containing pKD46 (Table S3) containing a Red genes (γ, β, and exo). The desired mutants were selected, and pKD46 was cured at 42°C. The mutation was verified using sequencing and PCR with flanking primers. The Map mutants MapΔMTS, MapΔWxxxA, which were subcloned into the pSA10 plasmid, were constructed using the 5' -phosphorylated primer pairs MTS delta F-MTS delta R and WxxxA F-WxxxA R, respectively (Table S4). Mutations were introduced by a site-directed mutagenesis method using Platinum Super II DNA polymerase, based on protocol B for deletions in the manufacturer's protocol (https://assets.thermofisher.com/TFS-Assets/LSG/manuals/MAN0014883_Platinum_SuperFi_PCR _MM_UG.pdf) (Thermo Fisher). The MapΔMTS, MapΔWxxxA-EspH1–25, and MapΔ101–152 mutants were prepared using the standard Gibson assembly method (65), using the Gibson assembly master mix (number E2611; NEB) and the respective linearization primers and the gblock (Table S4). The wild-type and mutated Map constructs were electroporated into EPEC-map, and Amp' EPEC-map::map::Kan strains were isolated (Table S1). The Map map-eGFP plasmid (Table S3) was constructed by the Gibson assembly method using mCherry-MAPΔ (Table S3) as the template. The primers EGFP F, EGFP R, Map EGFP F, and Map EGFP R (Table S4) were used for generating the Map-eGFP plasmid. All mutations were confirmed by sequencing. All bacterial strains used in this study were grown in Luria-Bertani medium supplemented with appropriate antibiotics (Table S1).

**Cell culture, ectopic expression of Map, and ADAM10 silencing by siRNA.** HeLa and CaCo-2BBe Tet-off cells were cultured as described elsewhere (13). Plasmid DNAs (eGFP, Map-eGFP, MTS-eGFP, mCherry, and mCherry-Map) (Table S3) were transiently transfected into HeLa cells (~60% confluence) for 15 h at 37°C, using the TransIT-X2 transfection reagent protocol (MIR 6004; Mirus, Madison, WI). For silencing ADAM10 expression in HeLa cells (~40% confluence), the TransIT-X2 transfection reagent was used to transfet the ADAM10 (50 nM; no. M-002000-00-0005; Dharmacon) or scrambled siRNA (50 nM; no. D-001206-14-05; Dharmacon) for 72 h at 37°C. ADAM10 expression was validated by Western blotting using specific antibodies (Table S2).

**Bacterial growth, activation, and infection.** The TSS of all bacterial strains used in the study was activated (3 h at 37°C and 5% CO₂) before infection, as described elsewhere (13). In EPEC-map::map-infected cells, Map expression was induced by supplementing the activation medium with 0.2 mM isopropyl-β-D-thiogalactopyranoside (IPTG; Promega, Madison, WI) during the last 30 min of activation. All infections were performed in a CO₂ incubator (37°C, 5% CO₂, 90% humidity) for 30 min, as described elsewhere (13).

**Confocal microscopy.** (i) Immunofluorescence labeling of fixed cells and confocal imaging. The procedures employed for confocal microscopy were described previously (13). Briefly, after infection, cells were fixed with 4% paraformaldehyde. F-actin was labeled with Texas Red (TR)-phalloidin, and the bacterial and HeLa cell DNA was labeled with DAPI (4',6-diamidino-2-phenylindole). Primary and fluorescence-labeled secondary antibodies used in this study are listed in Table S2. Cells were mounted and visualized using an Olympus FV-1200 laser scanning confocal microscope equipped with a 60×, 1.42-numerical-aperture oil immersion objective. The excitation wavelengths were 405 nm, 488 nm, 561 nm, and 633 nm, and the emission filter passbands were 430 to 470 nm, 505 to 525 nm, 575 to 625 nm, and 650 to 720 nm for DAPI, green, red, and far-red fluorescence, respectively. Confocal sections were acquired at z-axis intervals of 0.5 or 1 μm. The levels of F-actin at the infection sites were analyzed using Fiji (NIH) (66). A maximum-intensity projection was generated for each stack. The regions of cell-associated EPEC microcolonies were manually defined using the polygon-wire selections tool of Fiji and were termed infection sites. The average pixel intensity (PI) at the infection site (PI) was measured and expressed as normalized fluorescence from the average PI of an identically defined uninfected area located near the infecting microcolony (PLu), as follows: normalized fluorescence = (PI – PIu)/PIu. For colocalization analyses, line intensity profiles of the anti-hemagglutinin (HA) (Map) and anti-His60 (mitochondria) fluorescence were generated, as previously described (13), and are depicted in Fig. S11. Basically, a line was drawn across a randomly chosen confocal section showing Map and mitochondrial staining, and the fluorescence intensity along the line of the two different channels was generated using the intensity profile tool (plot-profile plug-in) of Fiji (NIH). Colocalized labeling was scored when the fluorescence intensity copeaked at a given place along the line (indicated by the green arrows in the intensity profile plots in Fig. S11). Data are presented as percentages of colocalizing and noncolocalizing fluorescence intensity determined in 10 such intensity profile determinations. (ii) Live-cell imaging. Cells were seeded on 8-well ibiTreat microslides (catalog no. 80826; ibidi) for 2 days (~70% confluence). Cells were washed twice with plain Dulbecco modified Eagle medium (DMEM) and then incubated with either the fluorescent mitochondrial membrane potential indicator tetramethyl-
Sheddase activity were expressed as AP activity levels measured in the supernatant (i.e., APreleased) determined by absorbance at 405nm, using a Synergy H1 microplate reader (Biotek). The levels of

where (13). Briefly, HeLa cells (70% confluence) were lysed in sample buffer (40% glycerol, 12% SDS, 0.2 M Tris-HCl [pH 6.8], and 100 mM dithiothreitol) supplemented with bromophenol blue, followed by heating (95°C for 5 min) and vortex shaking (10 min). Proteins separated by SDS-PAGE (Bio-Rad Mini-Protein Tetra system; 38 mA, 30 min) were transferred to a nitrocellulose membrane (Bio-Rad Trans-Blot Turbo; 2.5 A, 10 min). Membranes were then blocked (2.5% [wt/vol] bovine serum albumin [BSA] plus 1% [wt/vol] skim milk) for 1 h at 22°C, with agitation, washed with TBST (20 mM Tris [pH 8.0], 150 mM NaCl, and 0.1% Tween 20), and probed with the appropriate anti-phosphorylated-EGFR (pEGFR) or MAPK antibodies (Table S2). For stripping, the membranes were incubated in stripping buffer (62.5 mM Tris-HCl [pH 6.8], 100 mM 2-mercaptoethanol, 2% SDS) for 20 min at 55°C. Membranes were then washed with TBST and reprobed with anti-general-EGFR (gEGFR) or MAPK antibodies (Table S2). A Fusion FX Spectra imager (Vilber Smart Imaging, Collégien, France) was used to image the membranes. Band intensity was measured by Fiji (NIH). The band intensity of the phosphorylated protein was normalized to the band intensity of the total protein. The value obtained was further normalized to the protein level of the control experiment.

GTPase activity assay. The Rac/Cdc42 (p21) binding domain (PBD) of the human p21 activated kinase 1 (PAK1) protein fused to GST (pGEXTK-Fak1 70–117 [no. 12217; Addgene]) was expressed and coupled to glutathione Sepharose 4b beads (GST-PBD beads), as described (67). HeLa cells (70% confluence) were infected with various EPEC strains for 30 min at 37°C, then washed with cold PBS, and lysed in ice-cold lysis buffer (50 mM Tris [pH 7.6], 150 mM NaCl, 1% Triton X-100, 20 mM MgCl2) supplemented with protease inhibitors (phenylmethylsulfonyl fluoride [PMSF], leupeptin, and aprotinin). Lysates were sonicated for 10 s and centrifuged (14,000 rpm, 10 min, 4°C). The protein concentration of the lysates was determined by the bicinchoninic acid (BCA) protein assay method (catalog no. 23227; Thermo Fisher Scientific). Lysates (1 mg/ml) were incubated with 200 μg of GST-PBD beads for 30 min at 4°C and mixed by end-over-end rotation. Beads were washed three times with lysis buffer and dried with a Hamilton syringe, and associated proteins were analyzed by SDS-PAGE followed by Western blotting, as before.

Effector translocation assay. The effector translocation assay was performed as described elsewhere (13). Briefly, HeLa cells (~70% confluence) grown in 6-well plates were infected with preactivated EPEC strains for 30 min at 37°C. Cells were lysed with ice-cold lysis buffer (100 mM NaCl, 1 mM EDTA, 10 mM Tris-HCl [pH 7.4], 0.5% [vol/vol] NP-40) supplemented with protease and phosphatase inhibitors. Detergent-soluble (host cytoplasm and membranes containing the translocated protein effector) and -insoluble (bacteria and associated protein effectors) fractions were separated by centrifugation (16,000 × g, 4°C, 10 min) and analyzed for the presence of Map by Western blotting. The level of translocated effector was calculated by normalizing the levels of effector present in the NP-40-soluble fraction to the levels of the effector detected in the detergent-insoluble fraction.

Sheddase activity assay. The ectodomain shedding assay was performed essentially as described elsewhere (25). Briefly, HeLa cells grown on a 6-well plate (~60% confluence) were transfected with betacellulin-alkaline phosphatase (BTC-AP) or with TGF-α–AP. After 24 h, cells were infected with the appropriate EPEC strains for 30 min at 37°C. Medium containing bacteria was aspirated and replaced with Opti-MEM (reduced serum medium; no. 31985-047; Gibco) for 1 h at 37°C. The medium was then collected, and the cells were lysed in lysis buffer (2.5% [vol/vol] Triton X-100 in triple-distilled water [TDW]). Cell lysates were centrifuged (13,000 rpm, 5 min), and the supernatants were diluted 1:10 in a Triton–double-distilled-water (DDW) solution. Samples (100 μl) derived from media and cell lysates were placed in a 96-well plate and incubated with 4-nitrophenyl phosphate (100 μl) for 45 min at 37°C in the CO2 incubator. The levels of TGF-α–AP and BTC-AP released to the medium and in the cell lysates were determined by absorbance at 405 nm, using a Synergy H1 microplate reader (Biotek). The levels of sheddase activity were expressed as AP activity levels measured in the supernatant (i.e., APreleased) normalized to the total AP activity (i.e., APreleased + APcell). In experiments measuring sheddase activity in CCoP-treated cells, cells were first transfected with BTC-AP and then treated with CCoP (20 μM, 30 min, 37°C).

Apoptosis assay. Apoptosis assay was performed using the MCBCTO apoptosis kit (annexin V-fluorescin isothiocyanate [FITC] kit; no. 4700; MBL) per user instructions. Briefly, HeLa cells were grown in a 6-well plate (~70% confluence) and infected with the respective EPEC strains, as described in

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Materials and Methods. Cells were washed once with PBS, trypsinized, and washed once with serum-containing DMEM and once with PBS. They were suspended in 85 ul binding buffer, treated with annexin V-FITC and propidium iodide for 15 min at 22°C, and analyzed by flow cytometry (Cell Stream flow cytometer; Merck). The fractions of viable cells (annexin V negative, propidium iodide negative), early apoptotic cells (annexin V positive, propidium iodide negative), late apoptotic cells (annexin V positive, propidium iodide positive), and necrotic cells (annexin V negative, propidium iodide positive) were determined in each sample. Voltage settings for forward scatter (FSC) and side scatter (SSC) and fluorescence channels (488 and 561 nm) were kept constant for all samples. Data for 10,000 cells were recorded in each experiment.

Statistical analyses. Results are presented as means and standard errors (SE) of the means. A two-tailed Student’s t test determined statistical significance. A P value of < 0.05 indicates a statistically significant difference.

SUPPLEMENTAL MATERIAL
Supplemental material is available online only.

MOVIE S1, AVI file, 4.7 MB.
MOVIE S2, AVI file, 15.9 MB.
MOVIE S3, AVI file, 15.1 MB.
MOVIE S4, AVI file, 4.7 MB.
MOVIE S5, AVI file, 3.5 MB.
FIG S1, PDF file, 1.6 MB.

TABLE S1, DOCX file, 0.02 MB.
TABLE S2, DOCX file, 0.01 MB.
TABLE S3, DOCX file, 0.02 MB.
TABLE S4, DOCX file, 0.01 MB.

ACKNOWLEDGMENTS
We thank Carl P. Blobel and Gisela Weskamp (Weill Cornell Medicine), Rony Seger (Weizmann Institute), Israel Sekler (Ben-Gurion University), Gad Frankel (Imperial College), and David Engelberg and Joseph Orly (Hebrew University) for helpful discussions and reagents. We are also grateful to Carl Blobel, Rony Seger, and members of the Aroeti lab for critical reading of the manuscript.

This research was supported by the Israel Science Foundation (grants 1483/13 and 1671/19).

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Map-Induced MAPK Activity and Mitochondrial Cell Death


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Prepared for publication by mgbo.asm.org on behalf of the American Society for Microbiology.


