



# WWOX-Mediated Degradation of AMOTp130 Negatively Affects Egress of Filovirus VP40 Virus-Like Particles

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ABSTRACT Ebola virus (EBOV) and Marburg virus (MARV) continue to emerge and cause severe hemorrhagic disease in humans. A comprehensive understanding of the filovirus-host interplay will be crucial for identifying and developing antiviral strategies. The filoviral VP40 matrix protein drives virion assembly and egress, in part by recruiting specific WW domain-containing host interactors via its conserved PPxY late (L) domain motif to positively regulate virus egress and spread. In contrast to these positive regulators of virus budding, a growing list of WW domain-containing interactors that negatively regulate virus egress and spread have been identified, including BAG3, YAP/TAZ, and WWOX. In addition to host WW domain regulators of virus budding, host PPxY-containing proteins also contribute to regulating this late stage of filovirus replication. For example, angiomotin (AMOT) is a multi-PPxY-containing host protein that functionally interacts with many of the same WW domaincontaining proteins that regulate virus egress and spread. In this report, we demonstrate that host WWOX, which negatively regulates egress of VP40 virus-like particles (VLPs) and recombinant vesicular stomatitis virus (VSV) M40 virus, interacts with and suppresses the expression of AMOT. We found that WWOX disrupts AMOT's scaffoldlike tubular distribution and reduces AMOT localization at the plasma membrane via lysosomal degradation. In sum, our findings reveal an indirect and novel mechanism by which modular PPxY-WW domain interactions between AMOT and WWOX regulate PPxY-mediated egress of filovirus VP40 VLPs. A better understanding of this modular network and competitive nature of protein-protein interactions will help to identify new antiviral targets and therapeutic strategies.

**IMPORTANCE** Filoviruses (Ebola virus [EBOV] and Marburg virus [MARV]) are zoonotic, emerging pathogens that cause outbreaks of severe hemorrhagic fever in humans. A fundamental understanding of the virus-host interface is critical for understanding the biology of these viruses and for developing future strategies for therapeutic intervention. Here, we reveal a novel mechanism by which host proteins WWOX and AMOTp130 interact with each other and with the filovirus matrix protein VP40 to regulate VP40-mediated egress of virus-like particles (VLPs). Our results highlight the biological impact of competitive interplay of modular virus-host interactions on both the virus life cycle and the host cell.

**KEYWORDS** Marburg, WWOX, AMOT, VLP budding, VP40, filovirus, PPxY motif, L domain, angiomotin, WW domain, Ebola virus

Ebola virus (EBOV) and Marburg virus (MARV) are emerging pathogens that can cause severe hemorrhagic disease in humans, and these emerging pathogens remain a global public health threat that warrants urgent development of antiviral therapeutics (1–3). Toward this end, a more in-depth understanding of the interplay between the host and EBOV/MARV proteins would be beneficial both for our understanding of the fundamental molecular mechanisms of the virus life cycle and for identifying new targets and strategies for antiviral development. **Editor** Bryan R. G. Williams, Hudson Institute of Medical Research

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Our focus is on the filovirus VP40 protein, which is the major structural protein that drives virion assembly and egress. Expression of VP40 alone is sufficient for the formation and egress of virus-like particles (VLPs) that mimic the morphology of authentic virus particles (4–10). In our investigations of the VP40-host interactome, we identified a series of specific host interactors that either positively or negatively regulated the budding process (11–17). These host proteins contain one or more modular WW domains that interacted with the conserved PPxY late (L) domain motif at the N termini of both EBOV VP40 (eVP40) and MARV VP40 (mVP40). The PPxY L domain plays a key role in promoting efficient release of VLPs and live virus, in part, by hijacking or recruiting host factors that enhance VLP or virus egress from the plasma membrane (18). For example, the VP40 PPxY motif interacts with several members of the HECT family of E3 ubiquitin ligases, such as Nedd4, Itch, WWP1, and Smurf2, which facilitate monoubiquitination of VP40 and subsequent engagement and relocalization of the host ESCRT machinery to the site of virus budding at the plasma membrane to enhance virus-cell separation and release of VLPs or infectious virus (8, 11, 16, 17). In contrast to these positive regulators of virus budding, we have identified more recently a growing list of novel WW domain interactors that negatively regulate egress, such as host proteins BAG3, YAP/TAZ, and WWOX (12, 13, 15). Notably, these negative regulators of budding are multifunctional proteins that regulate diverse pathways/processes in the cell, including apoptosis/autophagy, transcription, cytoskeletal dynamics, cell migration/ morphology, and tight junction (TJ) formation (19-24). Moreover, regulation of these diverse pathways by WW domain-containing BAG3, YAP/TAZ, and WWOX is achieved, in part, via their interactions with host PPxY-containing proteins, such as angiomotin (AMOT) (25-30). Notably, AMOT contains three N-terminal PPxY motifs, two of which are identical to that in eVP40. Thus, we speculated that the modular mimicry of the viral and host PPxY motifs could lead to competitive interactions with host WW domaincontaining proteins, resulting in meaningful biological consequences for both the virus and the host. Indeed, we recently demonstrated that expression of AMOT had a positive influence on egress of both VP40 VLPs and live EBOV and MARV by counteracting the negative effects of YAP and WWOX (12, 13, 31).

In this report, we investigated further the interplay among AMOT, WWOX, and eVP40/mVP40 to understand the molecular mechanism by which these host proteins and modular interactions regulate budding. Our results revealed that WWOX interacts with and regulates expression and intracellular distribution of AMOT and that this interplay between AMOT and WWOX contributes mechanistically to the efficiency of VP40 VLP egress. Specifically, we found that expression of WWOX reduced the levels of AMOT in the cytoplasm and at the plasma membrane, as well as modified the intracellular localization of AMOT, and that these changes in AMOT expression and localization correlated with a decrease in VP40 VLP egress. Moreover, expression of WWOX led to lysosomal degradation of AMOT and a concomitant decrease in VP40 VLP egress. In sum, these data highlight a novel, complex network of modular PPxY-WW domain interactions that may impact both the host and the late stages of filovirus egress and spread.

# RESULTS

**WWOX** interacts with and reduces AMOTp130 expression levels. We demonstrated previously that expression of full-length, endogenous AMOT (AMOTp130) was important for the efficient egress of both VP40 VLPs and for live infectious EBOV and MARV, whereas expression of WWOX inhibited egress of VP40 VLPs from HEK293T cells. In light of the contrasting roles for PPxY-containing AMOTp130 and WW domain-containing WWOX in regulating VP40-mediated egress, we first sought to determine whether WWOX and AMOTp130 interact. Toward this end, we transfected HEK293T cells with expression plasmids for WWOX (myc tagged) and AMOTp130 (hemagglutinin [HA] tagged) and used an immunoprecipitation (IP)/Western blot (WB) approach to demonstrate a strong interaction between the two exogenously expressed proteins



**FIG 1** WWOX interacts with AMOT and reduces its expression levels in a dose-dependent manner. (A) Extracts from HEK293T cells transfected with myctagged WWOX and HA-tagged AMOT were immunoprecipitated (IP) with either normal mouse IgG or anti-myc antibody. AMOT and WWOX were detected in the precipitates by Western blotting (WB). Expression controls for AMOT, WWOX, and  $\beta$ -actin are shown in the left panel. (B) Extracts from HEK293T cells transfected with WWOX alone were IP with either normal mouse IgG or anti-myc antibody. Endogenous AMOT and WWOX were detected in the precipitates by WB. (C) HEK293T cells were transfected with AMOTp130 plus vector alone (-) or increasing amounts of WWOX. The indicated proteins were detected by WB, and AMOT expression levels were quantified using NIH ImageJ, as shown in parentheses. The amounts of AMOT in control cells (Ian 1) were set at 100%. The relative levels of AMOT are shown in parentheses. (D) Quantification of the AMOT levels from three independent experiments. Statistical significance was analyzed by a one-way analysis of variance (ANOVA). ns, not significant; \*\*, P < 0.01; \*\*\*\*, P < 0.0001. (E) HEK293T cells were transfected with AMOTp80 plus vector alone (-) or increasing amounts of WWOX. The indicated proteins were detected by WB, and AMOTp80 expression levels were quantified.

(Fig. 1A). In addition, we used the same approach to demonstrate that exogenously expressed WWOX interacted with endogenous AMOTp130 (Fig. 1B).

Next, we asked whether transfecting HEK293T cells with increasing amounts of WWOX plasmid would affect expression levels of AMOTp130 (Fig. 1C). Intriguingly, we observed a dose-dependent decrease in the levels of AMOTp130 as the amount of transfected WWOX plasmid was increased (Fig. 1C and D). Notably, WWOX expression did not reduce the level of exogenously expressed AMOTp80 (Fig. 1E)—an N-terminally deleted isoform of AMOTp130 that lacks all of the PPxY motifs. Together, these data show that WWOX interacts with AMOTp130 and that increased expression of WWOX leads to a dose-dependent decrease in AMOTp130 levels in HEK293T cells.

The PPxY-WW domain interplay is important for the AMOTp130-WWOX interaction and reduced expression of AMOTp130. Here, we sought to more precisely identify the regions of AMOTp130 and WWOX that are critical for mediating this interaction. Briefly, HEK293T cells were transfected with wild-type (WT) WWOX and WT AMOTp130 (Fig. 2A), single PPxY (PY) motif mutant PY1, PY2, or PY3 (Fig. 2B), or with the triple-PY-motif mutant PY123 (Fig. 2C), and an IP/Western blot assay was utilized to detect an interaction. Not surprisingly, the AMOTp130-PY123 triple mutant was unable to bind to WWOX and was undetectable on the gel (Fig. 2C, lane 2). In contrast, the



**FIG 2** WWOX interacts with and suppresses AMOT expression in a PY motif-dependent manner. (A to C) Extracts from HEK293T cells transfected with myctagged WWOX and HA-tagged AMOT WT (A), Flag-tagged AMOT PY1, PY2, and PY3 (B), or PY123 (C) were immunoprecipitated with either normal mouse IgG or anti-myc antibody. WWOX, AMOT WT, and PY motif mutants were detected in the precipitates by WB. Expression controls for AMOT WT and PY mutants WWOX and  $\beta$ -actin are shown in the left panels. \* indicates the IgG heavy chain. (D and E) HEK293T cells were transfected with AMOT WT or PY123 (D), PY1, PY2, and PY3 (E) mutants plus vector alone (-) or increasing amounts of WWOX, the indicated proteins were detected by WB, and AMOT levels were quantified using NIH ImageJ as shown in parentheses. (F) Quantification of the AMOT levels in panels D and E from three independent experiments. Statistical significance was analyzed by a one-way ANOVA. ns, not significant; \*, P < 0.05; \*\*, P < 0.01; \*\*\*\*, P < 0.001. (G) Schematic diagram of AMOTp130 with key domains highlighted. CC, coiled-coil domain; PDZ, PSD-95/DIg1/ZO-1 domain.

single PY mutants of AMOTp130 showed various degrees of binding to WWOX (Fig. 2B). We observed that mutant PY1 was essentially unable to interact with WWOX (Fig. 2B, lane 4), whereas both mutants PY2 and PY3 showed some degree of binding to WWOX, albeit significantly less than WT AMOTp130 (compare Fig. 2B, lanes 5 and 6, with Fig. 2A, lane 2). These data suggest that all three PY motifs of AMOTp130 participate in binding to WWOX, with PY motif 1 being the most important for an efficient interaction.

Next, we wanted to ask whether increased expression of WWOX would lead to reduced levels of expression of the PY mutants of AMOTp130, as we observed for WT AMOTp130 (Fig. 1C and D). Briefly, HEK293T cells were cotransfected with the indicated combinations of plasmids, and protein levels were quantified by Western blotting (Fig. 2D to F). While increased expression of WWOX once again resulted in reduced levels of WT AMOTp130 (Fig. 2D, lanes 1 to 3), increased expression of WWOX had no effect on expression of the AMOTp130-PY123 triple mutant in repeated experiments (Fig. 2D, lanes 4 to 6, and Fig. 2F). This finding strongly suggests that the PPxY-mediated interaction between AMOTp130 and WWOX is essential for the observed reduction in AMOTp130 expression. For the individual PY mutants, we found that increased expression of WWOX also had no significant effect on expression levels of PY1 (Fig. 2E, lanes 4 to 6, and Fig. 2F), which correlates well with our finding that PY motif 1 is likely most critical for mediating a strong interaction with WWOX (Fig. 2B). Increased expression of WWOX had a modest effect on reducing the levels of expression of mutants PY2 and PY3 in repeated experiments (Fig. 2E, lanes 7 to 12, and Fig. 2F).

Next, we wanted to interrogate the WW domains of WWOX for their role in interacting with AMOTp130 and in reducing expression levels of AMOTp130 in HEK293T cells. We focused our analysis of WW domain 1 (WW1) of WWOX, since this domain was first identified as the interacting domain with the VP40 PPxY motif and since WW domain 2



**FIG 3** WW domain 1 of WWOX interacts with AMOT. Extracts from HEK293T cells transfected with myc-tagged WT or the indicated mutants of WWOX and HA-tagged AMOT were subjected to IP. WWOX and AMOT in the precipitates (A) and cell extracts (B) were subjected to Western blotting. \* indicates the IgG heavy chain. (C) Quantification of the AMOT protein levels in panel B from three independent experiments. Statistical significance was analyzed by a one-way ANOVA. \*\*, P < 0.01; \*\*\*\*, P < 0.0001. (D) Schematic diagram of WWOX with key domains highlighted. Y33, W44, and P47 are three crucial amino acids in WW1 domain that mediate binding to the PY motif. NLS, nuclear localization signal; SDR, short-chain dehydrogenase/reductase domain.

(WW2) of WWOX is atypical in its amino acid sequence (32). Briefly, we generated a series of WW1 mutations, including the Y33R single-point mutant, W44A P47A doublepoint mutant, Y33R W44A P47A triple-point mutant, and  $\Delta$ WW deletion mutant. We observed that each of the WW1 domain mutants of WWOX were reduced significantly in their ability to interact with AMOTp130 as determined by IP/Western analysis (Fig. 3A). In addition, we observed approximate 2.5- and 5-fold reductions in the expression level of AMOTp130 in the presence of the W44A P47A double mutant and Y33R single mutant, respectively, compared to the control (Fig. 3B, lanes 3 and 4, and Fig. 3C). Coexpression of the Y33R W44A P47A and  $\Delta$ WW mutants resulted in a <2-fold reduction in AMOTp130 expression compared to the control (Fig. 3B, lanes 5 and 6, and Fig. 3C). Thus, the more subtle mutations (Y33R and W44A P47A) resulted in a more significant decrease in expression of AMOTp130 than that observed with the more severe mutations (Y33R W44A P47A and  $\Delta$ WW) of WWOX. Taken together, these results suggest that the PPxY-WW domain interplay between AMOTp130 and WWOX is critical for their ability to physically interact and plays a contributing role in the mechanism by which WWOX reduces the levels of AMOTp130 accumulation in HEK293T cells.

WWOX alters the intracellular localization pattern of AMOTp130. We have shown previously that AMOTp130 displays a tubular, filamentous pattern of distribution in the cytoplasm of HEK293T cells as determined by confocal microscopy (31). Here, we sought to determine whether coexpression of WWOX would alter this intracellular distribution of AMOTp130, in addition to its observed role in reducing the level of AMOTp130 expression. Briefly, HEK293T cells were transfected with AMOTp130, WWOX, or both AMOTp130 and WWOX, and cells were imaged using confocal microscopy (Fig. 4). We observed the expected tubular pattern of AMOTp130 (green) throughout the cytoplasm when expressed alone, and we observed an overall diffuse cytoplasmic pattern with some punctate staining for WWOX (red) when expressed alone (Fig. 4). Interestingly, cells coexpressing AMOTp130 and WWOX revealed a dramatic change in the distribution pattern for AMOTp130 from a tubular pattern to a more punctate and perinuclear localized pattern (Fig. 4). Indeed, AMOTp130 appeared to become sequestered in small puncta/ vesicles with a minimal amount of colocalization (Fig. 4, inset) in cells expressing WWOX. In sum, we observed a profound change in the pattern of distribution for AMOTp130 in the absence versus the presence of WWOX, which likely correlates with the observed reduction in expression levels of AMOTp130 described above.

**Increased expression of WWOX reduces AMOT expression and VP40 VLP egress.** Next, we wanted to determine whether increased expression of WWOX would simultaneously reduce expression of AMOTp130 and inhibit egress of VP40 VLPs.



**FIG 4** WWOX alters the intracellular distribution of AMOT. HEK293T cells were transfected with HA-tagged AMOTp130 (green) and myc-tagged WWOX (red) alone or cotransfected with both. Cells were then visualized via immunofluorescence staining. Scale bars, 10  $\mu$ m.

HEK293T cells were transfected with constant amounts of AMOTp130 and eVP40 or mVP40 plasmids, along with increasing amounts of WWOX, and the levels of the indicated proteins were quantified in cell extracts and VLPs by Western blotting (Fig. 5). Interestingly, we observed that WWOX selectively and significantly reduced the levels of AMOTp130 in cell extracts in a dose-dependent manner without any effect on the levels of eVP40 (Fig. 5A and B) or mVP40 (Fig. 5C and D) in the same cell extracts. In contrast, the production of eVP40 (Fig. 5A and B) and mVP40 (Fig. 5C and D) VLPs was significantly decreased in a dose-dependent manner with increasing expression of WWOX (Fig. 5A and C, VLPs, compare lanes 2 to 6).

To further confirm that WWOX-mediated reduction of AMOTp130 is specific and selective, we utilized another AMOTp130 interactor: HECT E3 ubiquitin ligase ltch. Itch contains four WW domains and is known to interact with and ubiquitinate AMOTp130 (33). We first confirmed the interaction of AMOTp130 and Itch by IP/Western analysis (Fig. 5E). Next, we demonstrated that unlike increasing amounts of WWOX, increasing amounts of Itch did not alter the levels of AMOTp130 in cell extracts (Fig. 5F), nor did Itch expression result in a decrease in either eVP40 or mVP40 VLP egress (Fig. 5G and H). Taken together, these results indicate that the observed WWOX-mediated reduction of AMOTp130 expression is selective and likely not a general phenomenon for all AMOTp130 interactors.

**WWOX and AMOT affect the intracellular localization of VP40.** Next, we used confocal microscopy to visualize any changes in the spatial distribution of VP40 in the absence or presence of WWOX and AMOTp130. Briefly, eVP40 and AMOT were expressed in HEK293T cells in the absence (Fig. 6, top row) or presence (Fig. 6, middle and bottom rows) of WWOX, and representative confocal images are shown. As expected, eVP40 was present throughout the cytoplasm, with light accumulation at the plasma membrane, and AMOT was distributed as tubular bundles throughout the cells (Fig. 6, top row, arrow). Conversely, VP40 appeared to accumulate more heavily at



**FIG 5** WWOX suppresses AMOT expression and inhibits VP40 VLP egress. (A and C) HEK293T cells were transfected with a constant amount of eVP40 (A), mVP40 (C), and AMOT plus vector alone (-) or increasing amounts of WWOX. The indicated proteins were detected in cell extracts and VLPs by WB. The cellular levels of AMOT, eVP40, mVP40, and VP40 in VLPs were quantified using NIH ImageJ. The amounts of eVP40 (A, cells, lane 1), mVP40 (C, cells, lane 1), and AMOT (A and C, cells, lane 2) in control cells were set at 100%. Also, eVP40 (A, VLPs, lane 1) and mVP40 (C, VLPs, lane 1) VLP production from control cells was set at 100%. Numbers in parentheses represent relative protein levels and VLP budding efficiency compared to the control. (B and D) Quantification of the indicated cellular protein levels and relative budding efficiency of eVP40 (B) and mVP40 (D) VLPs from three independent experiments. Statistical significance was analyzed by a one-way ANOVA. ns, not significant; \*\*, P < 0.01; \*\*\*, P < 0.001; \*\*\*\*, P < 0.001. (E) Extracts from HEK293T cells transfected with myc-tagged Itch and HA-tagged AMOTp130 were IP with either normal mouse IgG or anti-myc antibody. AMOTp130 and Itch were detected in the precipitates by WB. (F) HEK293T cells were transfected with AMOTp130 plus vector alone (-) or increasing amounts of Itch. The indicated proteins were detected by WB, and AMOTp130 expression levels were quantified. (G and H) HEK293T cells were transfected with a constant amount of eVP40 (G), mVP40 (H), and AMOTp130 plus vector alone (-) or increasing amounts of Itch. The indicated proteins were detected in cell extracts and NLPs by WB, and VP40 in VLPs was quantified using NIH ImageJ (numbers in parentheses).



FIG 6 Expression of WWOX and AMOT affects localization of VP40. HEK293T cells were transfected with GFP-eVP40 (green) and HA-tagged AMOTp130 (red) alone or with myc-tagged WWOX (white) and visualized by confocal microscopy. Scale bars, 10 μm.

the cell periphery and could be detected more readily in the nucleus in cells coexpressing both AMOT and WWOX (Fig. 6, middle and bottom rows, triangles). Notably, AMOT no longer displayed the tubular pattern in these cells, but rather was more punctate and disperse. These data show that exogenous expression of WWOX resulted in redistribution of both AMOT and VP40. It is tempting to speculate that WWOX disrupts the normal cytoskeletal association of AMOT, leading to its degradation in punctate vesicles. Moreover, WWOX may "drag" a portion of VP40 into the nucleus and the remainder of VP40 accumulates at the plasma membrane where it remains tethered and unable to bud efficiently due in part to the disruption of AMOT tubular distribution.

Inhibition of VP40 VLP egress by WWOX is dependent on expression of AMOTp130. To further test our hypothesis that disruption of AMOTp130 expression by WWOX leads to a decrease in VP40 VLP egress, we used shCtrl and shAMOT knock-down cell lines in our VLP budding assay (Fig. 7). Briefly, shCtrl and shAMOT cells were transfected with a constant amount of eVP40 (Fig. 7A and B) or mVP40 (Fig. 7C and D) in the absence or presence of increasing amounts of WWOX, and cell lysates and VLPs were harvested for analysis by Western blotting. We observed a significant decrease in both eVP40 (10-fold) and mVP40 (50-fold) VLP egress in the shCtrl cells in the presence of WWOX (Fig. 7A and C, compare lanes 1 to 3). As expected from prior results, budding of both eVP40 and mVP40 VLPs is significantly reduced in shAMOT cells compared to that in shCtrl cells (Fig. 7A and C, compare lanes 1 and 4). Notably, expression of WWOX did not significantly decrease budding of either eVP40 (<2-fold) or mVP40 (2-fold) in the shAMOT cells (Fig. 7A and C, compare lanes 4 to 6). In sum, these results support our hypothesis that inhibition of VP40 VLP egress by WWOX is due, in part, to WWOX's physical interaction with, and functional disruption of, AMOTp130.

**WWOX induces lysosomal degradation of AMOTp130.** The ubiquitin-proteasome system and the lysosomal pathway are the two major pathways involved in protein degradation and turnover in eukaryotic cells. As a multifunctional scaffolding protein, AMOTp130 expression and stability are tightly regulated by other host proteins/



**FIG 7** Expression of AMOT is required for WWOX-mediated inhibition of VP40 VLP egress. (A and C) shCtrl and shAMOT cells were transfected with a constant amount of eVP40 (A) or mVP40 (C) with vector alone (-) or increasing amounts of WWOX. The indicated proteins were detected in cell extracts and VLPs by WB. VP40 levels in VLPs were quantified using NIH ImageJ software as shown in parentheses. (B and D) Quantification of the relative budding efficiency of eVP40 (B) or mVP40 (D) VLPs from three independent experiments (n = 3). WWOX-negative samples were normalized independently for control and shAMOT conditions (B and D). Statistical significance was analyzed by a one-way ANOVA. ns, not significant; \*, P < 0.05; \*\*\*, P < 0.001; \*\*\*\*, P < 0.0001.

pathways, including LATS1/2 kinases and E3 ubiquitin ligases such as Nedd4 and Itch (25, 33–39). Thus, it was of interest to determine more precisely how WWOX expression may induce degradation of AMOTp130. Since AMOTp130 appeared to be sequestrated in punctate vesicles in the presence of WWOX (Fig. 4 and 6), we hypothesize that AMOTp130 may be subjected to degradation by the lysosomal pathway. To test our hypothesis, we utilized confocal microscopy and incorporated lysosomal marker protein LAMP1, to determine whether AMOTp130 localizes to lysosomes in the presence of WWOX (Fig. 8A). We observed the expected tubular pattern of AMOTp130 in the absence of WWOX, and under these conditions, AMOTp130 did not colocalize with LAMP1 (Fig. 8A, top row). In contrast, we observed clear colocalization of AMOTp130 in vesicles containing LAMP1 and the loss of the tubular pattern in the presence of WWOX (Fig. 8A, bottom row, arrows and enlarged view).

To further illustrate the temporal and spatial distribution dynamics of AMOTp130



**FIG 8** WWOX induces lysosomal degradation of AMOT. (A) HEK293T cells were transfected with the indicated combinations of plasmids, including lysosome marker mCherry-LAMP1 (red), HA-tagged AMOT (green), and myc-tagged WWOX (white). Cells were visualized using confocal microscopy. Scale bars, 10  $\mu$ m. The enlarged view and arrows highlight colocalization of AMOT and LAMP1 in lysosomes. (B) HEK293T cells were transfected with YFP-AMOT and mCherry-LAMP1 plus vector alone (Control) or WWOX. Live cells were observed via spinning disk confocal microscopy, beginning at 12 h posttransfection. Representative images showing the localization of AMOTp130 in control and WWOX-expressing cells at each hour during observation. Scale bars, 10  $\mu$ m.

and LAMP1 in the absence or presence of WWOX, we conducted time-lapse confocal microscopy using live HEK293T cells (Fig. 8B). Briefly, yellow fluorescent protein (YFP)-AMOTp130 and mCherry-LAMP1 fusion proteins were coexpressed in HEK293T cells with vector alone or WWOX. At 12 h posttransfection, cells were subjected to live cell spinning disk confocal microscopy for 5 h, and images were taken every 10 min. Representative images at each hour time point are shown highlighting the changes in localization of AMOTp130 in the absence or presence of WWOX (Fig. 8B). We observed that the tubular pattern of AMOTp130 did not change significantly over time in cells lacking WWOX (Fig. 8B, top panels). In contrast, the tubular distribution pattern of AMOTp130 was altered to a more punctate pattern over time in the presence of WWOX (Fig. 8B, bottom panels). Taken together, these results suggest that the mechanism by which WWOX reduces expression of AMOTp130 involves lysosomal-mediated degradation.

Pharmacological inhibition of lysosome function restores expression of AMOTp130 and rescues VP40 VLP budding in the presence of WWOX. If WWOX-mediated degradation of AMOTp130 occurs via the lysosomal pathway, then we reasoned that treating cells with the lysosomal inhibitor chloroquine (CQ) (40, 41) should restore expression of AMOTp130 and rescue VP40 VLP egress in the presence of WWOX. To assess restoration of AMOTp130 expression, HEK293T cells were either mock treated or treated with CQ and transfected with the indicated combination of AMOTp130 and WWOX plasmids (Fig. 9A and B). As expected, AMOTp130 levels were significantly reduced in mocktreated cells in the presence of WWOX (Fig. 9A, lanes 1 to 3, and Fig. 9B); however,



**FIG 9** Pharmacological inhibition of lysosome function restores AMOT expression and rescues VP40 VLP budding. (A) HEK293T cells were transfected with a constant amount of AMOTp130 plus vector alone (-) or increasing amounts of WWOX. After transfection, cells were treated with (lanes 4 to 6) or without (lanes 1 to 3) the lysosomal inhibitor chloroquine (CQ) (50  $\mu$ M) for 16 h. The indicated proteins were detected in cell extracts by WB. AMOT levels were quantified using NIH ImageJ as shown in parentheses. (B) Quantification of AMOT (A) from three independent experiments (n = 3). Statistical significance was analyzed by a one-way ANOVA. ns, not significant; \*\*\*, P < 0.001; \*\*\*\*, P < 0.001. (C) HEK293T cells were transfected with a constant amount of eVP40 and AMOT plus vector alone (-) or increasing amounts of WWOX. Cells were treated with (lanes 5 and 6) or without (lanes 3 and 4) CQ (50  $\mu$ M) for 16 h. The indicated proteins were detected in cell extracts and VLPs (Continued on next page)

there was no significant change in AMOTp130 levels in cells treated with CQ in the presence of WWOX (Fig. 9A, lanes 4 to 6, and Fig. 9B). When we coexpressed eVP40 under the same conditions, we observed almost a complete rescue of eVP40 VLP budding back to WT levels in cells treated with CQ in the presence of WWOX (Fig. 9C, lanes 5 and 6, and Fig. 9D) compared to mock-treated controls (Fig. 9C, lanes 3 and 4, and Fig. 9D). These findings suggest that the rescue of eVP40 VLP egress in CQ-treated cells is likely due, in part, to the restoration of AMOTp130 back to WT levels.

Finally, we asked whether the ubiquitin-proteasome system may also be involved in WWOX-mediated degradation of AMOTp130 using a pharmacological approach. Briefly, HEK293T cells were treated with dimethyl sulfoxide (DMSO) or the proteasome inhibitor MG132 (39), and cells were transfected with the indicated combinations of plasmids (Fig. 9E). In contrast to our findings following treatment with CQ, treatment of cells with MG132 did not lead to restoration of AMOTp130 levels in the presence of WWOX (Fig. 9E and F). In sum, these results suggest that WWOX represses AMOT by induction of its lysosomal degradation and thus modulates AMOT and eventually leads to the inhibition of the VLPs' egress.

# DISCUSSION

EBOV and MARV VP40 matrix protein utilize L domain motifs (PPxY, PTAP, and YxxL) to recruit specific host proteins to facilitate virus egress and dissemination (5, 8, 9, 42, 43). In addition to the recruitment of host proteins that positively regulate budding (e.g., Tsg101, Nedd4, Itch, WWP1, and Smurf2) (11, 16, 17, 42, 44), the VP40 PPxY L domain also engages with host proteins that negatively regulate VP40-mediated egress (e.g., BAG3, YAP/TAZ, and WWOX) (12, 13, 15). Thus, the modular and competitive nature of the PPxY-WW domain interplay likely impacts both host and virus functions (18). Notably, host PPxY-WW domain interactions regulate diverse signaling networks and major cellular processes, such as the Hippo pathway and cell division/migration.

Here, we describe how the physical and functional interaction between host AMOTp130 and host WWOX affects VP40 VLP budding (Fig. 10). AMOTp130 is a key multi-PPxY-containing host protein that engages host WW domain-containing proteins that both positively and negatively impact viral budding. For example, AMOTp130 interacts with the YAP, BAG3, and WWOX in a PPxY-WW domain-dependent manner to function as a "master regulator" of several physiologically relevant pathways/processes, including transcription (Hippo pathway), apoptosis, cytoskeletal dynamics, and tight junction (TJ) integrity (19, 23, 25, 26, 28, 45–48). In addition, AMOT stability and turnover are tightly regulated via PPxY-WW domain interactions with Nedd4 E3 ubiquitin ligase family members (33, 35, 36, 39). While AMOT was previously shown to regulate assembly and egress of non-PPxY-containing viruses (49–51), we recently revealed a role for endogenous AMOT in positively regulating egress of PPxY-containing EBOV and MARV (12, 13, 31).

We showed that WWOX reduced expression and modulated intracellular distribution of AMOTp130 in a PPxY-WW domain-dependent manner. Indeed, we observed a dose-dependent reduction in expression of PPxY-containing AMOTp130, but not PPxYlacking AMOTp80, in the presence of WWOX (Fig. 1). Moreover, expression of a mutant AMOTp130 containing mutations in all three PPxY motifs was not affected by exogenous expression of WWOX (Fig. 2), and a WW domain mutant of WWOX did not reduce the levels of AMOTp130 compared to that of WT WWOX (Fig. 3). Our data suggest that PY1 (LPTY) of AMOTp130 may be most important for binding strongly to WWOX, with

## FIG 9 Legend (Continued)

by WB. The yields of eVP40 VLPs were quantified using NIH ImageJ software as shown in parentheses. (D) Quantification of the relative budding efficiency of eVP40 VLPs under the indicated conditions from three independent experiments (n = 3). Statistical significance was analyzed by a one-way ANOVA. ns, not significant; \*, P < 0.05; \*\*\*\*, P < 0.0001. (E) HEK293T cells were transfected with a constant amount of AMOT plus vector alone (-) or increasing amounts of WWOX. Cells were treated with (lanes 4 to 6) or without (lanes 1 to 3) proteasomal inhibitor MG132 (10  $\mu$ M) for 8 h before harvesting. The indicated proteins were detected in cell extracts by WB. (F) Quantification of AMOT (E) from three independent experiments (n = 3). Statistical significance was analyzed by a one-way ANOVA. ns, not significant; \*\*\*, P < 0.001; \*\*\*\*, P < 0.001.



**FIG 10** Working model of PPxY-WW domain interactions among AMOT, WWOX, and VP40. (Left) AMOTp130 facilitates VP40 VLP egress via its ability to bind actin and regulate its dynamics at the plasma membrane. (Right) Exogenously expressed WWOX interacts with both AMOTp130 and VP40 in a PPxY-WW domain-dependent manner. These interactions lead to reduced levels of VP40 at the plasma membrane, enhanced nuclear localization of VP40 (red arrow), and lysosomal-mediated degradation of AMOTp130 (black arrow), all of which results in a decrease of VLP egress.

PY2 and PY3 binding with lower affinity. Interestingly, a similar order of binding affinity was identified for the AMOTp130 PY motifs (PY1 > PY2 > PY3) binding to WW domain 3 of Nedd4L as a mechanism to stimulate release of HIV-1 (49). Thus, the more noncanonical PY1 motif of AMOTp130 may play the most important role in regulating (either positively or negatively) both PPxY-containing and PPxY-lacking viruses (49–51).

Under conditions of exogenous expression of WWOX, we observed that budding of both eVP40 and mVP40 VLPs was significantly reduced (Fig. 4). We hypothesized that one possible mechanism could be that WWOX was negatively regulating budding of VP40 VLPs indirectly, by reducing expression of AMOTp130 and preventing it from facilitating egress of VLPs from the plasma membrane as reported previously (13, 31). In support of this hypothesis, we found that budding of both eVP40 and mVP40 VLPs was not significantly reduced in shAMOT cells in the presence of WWOX, but was significantly reduced in shCtrl cells in the presence of WWOX (Fig. 7). These findings suggest that WWOX's negative effect on VP40 VLP egress is the result of a novel, indirect mechanism of action that requires expression of, and likely an interaction with, endogenous AMOTp130.

We sought to further understand the mechanism by which WWOX reduced expression of AMOTp130. Toward this end, our results suggest that WWOX mediates reduction of AMOTp130 levels via the lysosomal degradation pathway, as judged by confocal microscopy and the use of pharmacological inhibitors (Fig. 8 and 9). Notably, WWOX was unable to reduce expression of AMOTp130 in cells treated with chloroquine compared to controls. These conditions also resulted in the restoration of budding of VP40 VLPs to near WT levels in cells expressing WWOX and treated with chloroquine. In contrast to the results with chloroquine, WWOX retained the ability to reduce the levels of AMOTp130 in cells treated with MG132, suggesting that WWOX-mediated degradation of AMOTp130 was not occurring via the ubiquitin/proteasome pathway.

In addition to the indirect mechanism of inhibition of VLP budding described above, our data also suggest that WWOX can inhibit egress of VP40 VLPs via a direct PPxY-WW domain interaction that leads to reduced levels of VP40 at the plasma membrane and increased levels of VP40 detected in the nucleus (Fig. 10) (12). Indeed, we observed enhanced nuclear localization of VP40 in cells expressing WWOX. Since WWOX contains a nuclear localization signal and normally shuttles in and out of the nucleus (52), one possibility is that WWOX may sequester or drag some VP40 into the nucleus as a result

of a direct PPxY-WW domain interaction, leading to a subsequent decrease in VLP budding. Interestingly, nuclear localization of eVP40 has been reported previously (53-55); however, a functional role for nuclear eVP40 has not been described. Investigations into a potential role for eVP40 in the nucleus may be warranted due in part to the identification of nuclear transcriptional regulators such as WWOX and YAP/TAZ as specific host interactors. It is tempting to speculate that host proteins such as WWOX and YAP/TAZ may interact with and translocate eVP40 into the nucleus, where these virus-host complexes may then affect transcription of WWOX and/or YAP/TAZ-responsive genes to generate a cellular environment beneficial for virus replication, budding, and/or disease progression. Alternatively, since WWOX is known to directly bind to multiple transcriptional activators, such as p38, p73, AP-2y, ErBb4, c-Jun, and RUNX2, in a PPxY-WW domain-dependent manner (22, 23, 56-60), it will be of interest to determine whether there is any competitive interplay among these host proteins and PPxY-containing VP40 proteins in VP40-expressing or virus-infected cells that may result in biological consequences having an impact on both the virus and host. Such studies may also provide novel insights into the development of new host-oriented antivirals that target these modular virus-host interactions.

#### **MATERIALS AND METHODS**

Cells, antibodies, and plasmids. HEK293T-based shCtrl and shAMOT cells (kindly provided by J. Kissil, Scripps Research, FL) and HEK293T cells were maintained in Dulbecco's modified Eagle's medium (DMEM) (Corning) supplemented with 10% fetal bovine serum (FBS) (Gibco), penicillin (100 U/mL)/streptomycin (100 µg/mL) (Invitrogen). Cells were grown at 37°C in a humidified 5% CO, incubator. The primary antibodies used in this study include mouse anti-myc (Millipore), mouse anti-HA antibody (Sigma), mouse anti-Flag (Sigma), mouse anti-AMOT (Santa Cruz), rabbit anti-eVP40 (IBT), and mouse anti- $\beta$ -actin (Proteintech). The plasmids encoding eVP40 and green fluorescent protein (GFP)-eVP40 were described previously (8, 42). Flag-tagged mVP40 was kindly provided by S. Becker (Institut für Virologie, Marburg, Germany). The Flag-tagged AMOT PY123, PY1, PY2, and PY3 mutants were kindly provided by J. Kissil (Scripps Research Institute, FL). The YFP-tagged AMOTp130 was kindly provided by K.-L. Guan (University of California, San Diego). The mCherry-tagged LAMP1 was a gift from Amy Palmer (Addgene plasmid 45147). The myc-tagged Itch plasmid was kindly provided by G. Melino (Leicester University, United Kingdom). The myc-tagged WWOX plasmid was kindly provided by R. I. Aqeilan (Jerusalem, Israel). The WWOX mutants were generated via the QuikChange method, and the primers used are as follows: Y33R1 (5' GGTGTGATTGGCGTAGCGAACCCAGCCGTCCTTG 3'), Y33R2 (5' CAAGGACGGCTGGGTT CGCTACGCCAATCACACC 3'), W44AP47A1 (5' TTTCTTTTTCCAGTTTTTGCATGTTCCGCCTGAGTCTTCTCCTC GGTG 3'), W44AP47A2 (5' CACCGAGGAGAAGACTCAGGCGGAACATGCAAAAACTGGAAAAAGAAA 3'),  $\Delta$ WW1, (5' CATCCACAGTAAACGCGTCCTCACTGTCCGTG 3'), and  $\Delta$ WW2 (5' CACGGACAGTGAGGACGCGT TTACTGTGGATG 3').

**Immunoprecipitation assay.** HEK293T cells seeded in 6-well plates were transfected with the indicated plasmid combinations using Lipofectamine reagent (Invitrogen). At 24 h posttransfection, cells were harvested and lysed, and the cell extracts were subjected to Western blotting (WB) and coimmunoprecipitation (co-IP). The protein complexes were precipitated by either mouse IgG or anti-myc antibody. First, the cell extracts were incubated with antisera overnight at 4°C with continuous rotation, then the protein A/G agarose beads (Santa Cruz) were added to the mixtures and incubated for 5 h with continuous rotation. After incubation, beads were collected via centrifugation and washed 5 times. The input cell extracts and immunoprecipitates were then detected by WB with appropriate antisera as indicated.

Western blotting and VLP budding assays. HEK293T cells were transfected with 0.25  $\mu$ g AMOTp130 or -p80 plus increasing amounts (0.1, 0.25, 0.5, and 1.0  $\mu$ g) of WWOX plasmids, or cells were transfected with 0.25  $\mu$ g AMOTp130 WT or PY123, PY1, PY2, and PY3 mutants as well as with increasing amounts (0.25 and 0.5  $\mu$ g) of WWOX plasmids. The total amount of transfected DNA was equivalent in all samples. Cell extracts were harvested at 24 h posttransfection and then subjected to SDS-PAGE and WB analyses.

HEK293T cells were transfected with 0.25  $\mu$ g AMOT and 0.5  $\mu$ g WT or mutant WWOX. Cell extracts were subjected to WB and IP analyses. For VLP budding and WWOX titration experiments, HEK293T cells were transfected with 0.2  $\mu$ g of eVP40 or Flag-tagged mVP40 plus 0.25  $\mu$ g AMOTp130 and increasing amounts (0.1, 0.25, 0.5, and 1.0  $\mu$ g) of WWOX plasmids. The eVP40 and mVP40 in VLPs and the indicated proteins in cell extracts were detected by WB.

**Indirect immunofluorescence assay.** HEK293T cells were transfected with the indicated plasmid combinations. At 24 h posttransfection, cells were washed with cold phosphate-buffered saline (PBS) and fixed with 4% formaldehyde for 20 min at room temperature, then permeabilized with 0.2% Triton X-100. After being washed 3 times with PBS, cells were blocked for 1 h and then incubated with rabbit anti-myc (WWOX) or mouse anti-HA (AMOT) antisera. Next, cells were stained with Alexa Fluor 488, 594, or 647 goat anti-mouse/rabbit secondary antibodies (Life Technologies). The GFP-eVP40 and mCherry-LAMP1 were visualized via fluorescent tag. Cells were mounted with ProLong glass antifade mountant

with Hoechst 33342 (Life Technologies). Microscopy was performed using a Leica SP5 FLIM inverted confocal microscope. Serial optical planes of focus were taken, and the collected images were merged into one image by using the Leica microsystems (LAS AF) software.

VLP budding assay in HEK293T shCtrl and shAMOT cells. HEK293T shCtrl and shAMOT cells were transfected with 0.2  $\mu$ g of eVP40 or mVP40 plus vector or 0.25 or 0.5  $\mu$ g of WWOX. VP40 VLPs and eVP40, mVP40, WWOX, and endogenous AMOTp130 in cell extracts were detected by WB using appropriate antisera.

**Live-cell imaging and time-lapse microscopy.** HEK293T cells were seeded on chambered cover glasses and transfected with YFP-AMOT (0.25  $\mu$ g) and mCherry-LAMP1 (0.25  $\mu$ g) plus vector or WWOX (0.5  $\mu$ g). Live cells were observed at 12 h posttransfection using a Leica DMI4000 microscope with Yokagawa CSU-X1 spinning disk confocal attachment. Images were taken every 10 min over a 5-h window of observation.

**Pharmacological inhibition of lysosome or proteosome functions.** HEK293T cells were transfected with 0.25  $\mu$ g AMOTp130 plus vector or 0.25 or 0.5  $\mu$ g WWOX plasmids. For lysosomal inhibition, cells were untreated or treated with CQ (50  $\mu$ M) for 16 h. For proteasomal inhibition, cells were treated with DMSO or MG132 (10  $\mu$ M) at 8 h before harvest. The indicated proteins were detected via WB. For VLP budding, HEK293T cells were transfected with 0.2  $\mu$ g eVP40 alone or with 0.25  $\mu$ g AMOTp130 and 0.25 or 0.5  $\mu$ g WWOX plasmids. At 6 h posttransfection, cells were untreated or treated with CQ (50  $\mu$ M) for 16 h. Then cell extracts and VLPs were harvested and subjected to WB analysis.

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