PtdIns(4,5)P$_2$ and PtdIns3P coordinate to regulate phagosomal sealing for apoptotic cell clearance

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Phagocytosis requires phosphoinositides (PIs) as both signaling molecules and localization cues. How PIs coordinate to control phagosomal sealing and the accompanying switch of organelle identity is unclear. In this study, we followed dynamic changes in PIs during apoptotic cell clearance in *Caenorhabditis elegans*. We found that phosphatidylinositol-4,5-bisphosphate (PtdIns(4,5)P$_2$) and phosphatidylinositol-3-phosphate (PtdIns3P), which accumulate transiently on unsealed and fully sealed phagosomes, respectively, are both involved in phagosome closure. We identified PtdIns3P phosphatase MTM-1 as an effector of PtdIns(4,5)P$_2$ to promote phagosomal sealing. MTM-1 coordinates with the class II PI3 kinase PIKI-1 to control PtdIns3P levels on unsealed phagosomes. The SNX9 family protein LST-4 is required for sealing, and its association with unsealed phagosomes is regulated by PtdIns(4,5)P$_2$, PIKI-1, and MTM-1. Loss of LST-4 or its retention on phagosomes disrupts sealing and suppresses PtdIns3P accumulation, indicating close coupling of the two events. Our findings support a coincidence detection mechanism by which phagosomal sealing is regulated and coupled with conversion from PtdIns(4,5)P$_2$ enrichment on unsealed phagosomes to PtdIns3P enrichment on fully sealed phagosomes.

**Introduction**

Phosphoinositides (PIs) are concentrated on different organelles to direct signaling and membrane trafficking events. For example, phosphatidylinositol-4,5-bisphosphate (PtdIns(4,5)P$_2$) is enriched in the inner leaflet of plasma membranes and is involved in almost all cellular events occurring at the cell surface, whereas phosphatidylinositol-3-phosphate (PtdIns3P) is concentrated on early endosomes and is a key determinant of endosome maturation (Di Paolo and De Camilli, 2006; Bohdanowicz and Grinstein, 2013). The differential distribution of PIs is determined by PI kinases and phosphatases that catalyze local PI synthesis and turnover. PIs mediate signaling or trafficking events by affecting activities of integral membrane proteins or recruiting cytosolic effectors to specific membrane compartments (Di Paolo and De Camilli, 2006; Balla, 2013). PI–protein interactions mediated by electrostatic interaction or by protein modules are of low affinity, and stable protein–membrane association requires additional lipid, protein, or geometry determinants within the membrane (Carlton and Cullen, 2005; Di Paolo and De Camilli, 2006). For instance, EEA1 binds both PtdIns3P and phosphatidylyserine through distinct sites in the FYVE domain, suggesting that it can be enriched on membranes containing both lipids (Kutateladze et al., 2004; Carlton and Cullen, 2005). EEA1 also binds RAB-5 and is recruited to early endosomes through both lipid and RAB-5 binding (Lawe et al., 2000). Moreover, SNX1 contains both a PtdIns3P-binding PX domain and a curvature-sensing BAR (Bin–Amphiphysin–Rvs) domain and localizes to high-curvature membrane tubules emanating from endosomes, indicating that membrane geometry can also be a localization cue (Carlton et al., 2004; Carlton and Cullen, 2005). Thus, multiple components of a given membrane serve as the coincidence detection code to define organelle identity and direct membrane trafficking by controlling the spatial and temporal recruitment and release of effectors. Nascent transport vesicles must change identity through interconversion of PIs before fusing with the next membrane. How PI conversion is controlled and coordinated with fusion and/or fission events remains poorly understood.

Eukaryotic cells engulf and eliminate large particles like microorganisms and apoptotic cells through phagocytosis, which involves many typical features of vesicle biogenesis and maturation (Flannagan et al., 2012). In *Caenorhabditis elegans* hermaphrodites, 131 somatic cells and about half of the germ cells die through apoptosis, and the resulting cell corpses are phagocytosed and cleared by neighboring cells (Pinto and Hengartner, 2012). Apoptotic cells are recognized and engulfed through evolutionarily conserved pathways, leading to cytoskeleton reorganization and formation of a...
In C. elegans, apoptotic cell removal requires both DYN-1, the C. elegans dynamin, and LST-4, an SNX9 family protein, which are thought to function early in phagosome maturation (Yu et al., 2006; Kinchen et al., 2008; Almendinger et al., 2011; Chen et al., 2013). LST-4 interacts with DYN-1 and promotes phagosomal association of the latter (Lu et al., 2010; Chen et al., 2013). However, it is unclear whether LST-4 and DYN-1 regulate early maturation events or are involved in the sealing process.

Here, we identify a coincidence detection mechanism that regulates phagosomal sealing through LST-4 and couples sealing with the switch of membrane identity from PtdIns(4,5)P_2-enriched unsealed phagosomes to PtdIns3P-enriched fully sealed phagosomes.

### Results

**PtdIns(4,5)P_2 and PtdIns3P accumulate sequentially on apoptotic cell–containing phagosomes**

To examine how PIs change during apoptotic cell clearance in C. elegans, we expressed genetically coded PI sensors, including PLCδ1-PH (PtdIns(4,5)P_2), 2xFYVE (PtdIns3P), TAPP1 (PtdIns(3,4)P_2), BTK1-PH (PtdIns(3,4,5)P_3), and AKT1-PH (PtdIns(3,4)P_2 and PtdIns(3,4,5)P_3), which do not affect apoptotic cell clearance (Fig. S1, A–Q; Dowler et al., 2000; Lemon, 2008). In embryos, PLCδ1-PH and 2xFYVE mainly labeled plasma membranes and intracellular vesicles, respectively, which is consistent with enrichment of PtdIns(4,5)P_2 and PtdIns3P on cell membranes and endosomes (Fig. S1, A–B’).

We followed cell corpse clearance by four-dimensional microscopy analyses. PLCδ1-PH labeled extending pseudopods, which gradually surrounded cell corpses, a process that took ∼2.6 min (Fig. 1A). PLCδ1-PH stayed on phagosomes for 2.8 min, and its release was followed immediately by enrichment of 2xFYVE that persisted for ∼3.8 min (Fig. 1A and Video 1). No overlapping appearance of PLCδ1-PH and 2xFYVE was observed, suggesting that PtdIns(4,5)P_2 and PtdIns3P accumulate transiently on apoptotic cell–containing phagosomes at distinct stages. Later, 2xFYVE-negative phagosomes fused with lysosomes, marked by the lysosomal membrane protein LAAT-1 (Liu et al., 2012), to form phagolysosomes where apoptotic cells are degraded (Fig. 1B and Video 2). The sequential and nonoverlapping appearance of these reporters divides the phagosome formation and maturation processes into different stages, including PLCδ1-PH/PtdIns(4,5)P_2-positive, 2xFYVE/PtdIns3P-positive, FYVE/PtdIns3P-negative, and phagolysosomal (LAAT-1-positive) stages. In gla-3 RNAi worms, which have increased germ cell apoptosis but normal cell corpse clearance (Kritikou et al., 2006), PLCδ1-PH and 2xFYVE were observed on phagosomes enclosing germ cell corpses (Fig. S1, F and G).

In contrast, no obvious phagosomal association was seen in either embryos or germline when TAPP1, BTK1-PH, or AKT1-PH was expressed (Fig. S1, C–E’ and H–O). This suggests that PtdIns(3,4)P_2 and PtdIns(3,4,5)P_3 do not accumulate significantly on apoptotic cell–containing phagosomes. Consistent with this, AGE-1 and DAF-18, which produce and hydrolyze PtdIns(3,4)P_2, respectively, or R01H10.7, an INP4A homologue that dephosphorylates PtdIns(3,4)P_2, are all dispensable for apoptotic cell clearance (Fig. S1, R–X).

**PtdIns(4,5)P_2 regulates the phagosomal dynamics of the PtdIns3P phosphatase MTM-1**

The absence of 2xFYVE from PtdIns(4,5)P_2-positive extending pseudopods and phagosomes indicates that PtdIns3P does not accumulate at this stage, which may be because of a lack of PtdIns3P production or PtdIns3P hydrolysis. To test this, we examined the dynamics of C. elegans myotubularin MTM-1, a plasma membrane–localizing PtdIns3P phosphatase involved in apoptotic cell clearance (Fig. S2, A–C; Zou et al., 2009; Neukomm et al., 2011). As overexpression of wild-type MTM-1 causes persistent cell corpses (Zou et al., 2009), we used MTM-1(c378S), a catalytically inactive MTM-1 that does not affect cell corpse removal (Fig. S2, D and E). MTM-1(c378S) colocalized very well with PLCδ1-PH on plasma membranes and apoptotic cell–containing phagosomes (Fig. 1, C–C’’). It dis-
Figure 1. PtdIns(4,5)P$_2$ regulates the phagosomal dynamics of MTM-1. (A) Time-lapse images of a cell corpse in a wild-type embryo expressing mCHERRY::PLC$_\delta$1-PH and YFP::2xFYVE. "0 min" represents the time point when PLC$_\delta$1-PH was first detected around the cell corpse. The time period until the cell corpse was surrounded by PLC$_\delta$1-PH was defined as the pseudopod extension stage. Quantification of the pseudopod extension time and phagosomal

| transgene/duration | Control | +PPK-1::GFP |
|--------------------|---------|-------------|
| mCHERRY::PLC$_\delta$1-PH | 3.5 ± 1.2 (min) (n=11) | >19.1 (min) (n=24) |
| mCHERRY::MTM-1(C378S) | 3.8 ± 1.3 (min) (n=10) | >20.8 (min) (n=11) |

Embryonic stages

5% input 50% bound protein

PC S P PtdIns(4,5)P$_2$ PLC$_\delta$1-PH SMT3 MTM-1(C378S)

Coomassie Blue

15 kD 100 kD 25 kD

SMT3

MTM-1(C378S)

SMT3

PC + PtdIns(4,5)P$_2$ PLC$_\delta$1-PH

5% input 50% bound protein

SMT3

MTM-1(C378S)

PLC$_\delta$1-PH

Coomassie Blue

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played identical dynamics to PLCδ1-PH on extending pseudopods and phagosomes (Fig. 1 D, Fig. S2 F, and Video 3). The simultaneous appearance and release of MTM-1 and PLCδ1-PH prompted us to investigate whether MTM-1 dynamics on phagosomes are regulated by PtdIns(4,5)P2 levels. We expressed PPK-1, the sole C. elegans PtdIns(4,5)-kinase, and found that it displayed identical phagosomal dynamics to PLCδ1-PH (Fig. 1 E). PPK-1 expression significantly extended phagosomal duration of PLCδ1-PH, suggesting that PPK-1 produces PtdIns(4,5)P2 on forming phagosomes (Fig. 1, E and G). Similarly, MTM-1(C378S) appeared simultaneously with PPK-1 and persisted for significantly longer on phagosomes when PPK-1 was coexpressed (Fig. 1, F and G). Inactivation of OCR1-1, the inositol 5-phosphatase that hydrolyzes PtdIns(4,5)P2, caused prolonged MTM-1(C378S) duration on phagosomes like PPK-1 expression (see Fig. 4, F and G). These data suggest that MTM-1 dynamics on phagosomes are regulated by PtdIns(4,5)P2 levels. Overexpression of PPK-1 or inactivation of ocr1-1 affected apoptotic cell clearance, causing accumulation of cell corpses, indicating the importance of maintaining PtdIns(4,5)P2 and MTM-1 dynamics in this process (Fig. 1 H; Fig. 2, P–X; and Fig. S4 G). MTM-1(C378S) bound efficiently to PtdIns(4,5)P2 and MTM-1 dynamics in this process (Fig. 1 H; Fig. 2, P–X; and Fig. S4 G). MTM-1(C378S) bound to PC liposomes containing 4% PtdIns(4,5)P2 is more abundant in pellets (P) than in supernatants (S). These independent experiments were performed. Unpaired t tests were performed. *, P < 0.05. **, P < 0.0001. (J) MTM-1(C378S) bound to PC liposomes containing 4% PtdIns(4,5)P2 is more abundant in pellets (P) than in supernatants (S). Three independent experiments were performed. Unpaired t tests were performed. *, P < 0.005. **, P < 0.0001. (K) MTM-1(C378S) bound to PC liposomes containing 4% PtdIns(4,5)P2 is more abundant in pellets (P) than in supernatants (S). Three independent experiments were performed. Unpaired t tests were performed. *, P < 0.005. **, P < 0.0001. (L) MTM-1(C378S) bound to PC liposomes containing 4% PtdIns(4,5)P2 is more abundant in pellets (P) than in supernatants (S). Three independent experiments were performed. Unpaired t tests were performed. *, P < 0.005. **, P < 0.0001.

We found that MTM-1 was enriched on PtdIns(4,5)P2-positive phagosomes and was released before the appearance of 2xFYVE, RAB-5, and RAB-7, indicating that MTM-1 acts very early during phagosome formation and/or maturation (Fig. 1 D and Fig. S2, H and I). We isolated a recessive mutation of mtm-1, qx322, which contained a C to T transition that resulted in an early stop codon after Lys99. qx322 worms contained significantly more germ cell corpses than the wild type (Fig. S2 J). This phenotype is similar to inactivation of mtm-1 by RNAi as previously reported (Neukomm et al., 2011) or gk890934, which contains two missense mutations in mtm-1 (Fig. S2 J). Expression of wild-type, but not catalytically inactive, MTM-1 significantly reduced germ cell corpses in qx322 and gk890934, indicating that phosphatase activity is important for MTM-1 function in apoptotic cell clearance (Fig. S2 K). Expression of human myotubularin MTM1, which localized to plasma membranes and phagosomal surfaces like worm MTM-1, rescued the cell corpse phenotype of mtm-1(qx322) (Fig. S2, L–N”), suggesting that human MTM1 can substitute for worm MTM-1 in removing germ cell corpses. The persistent cell corpses in mtm-1(RNAi) animals were all surrounded by Myr-GFP, a plasma membrane reporter, as in gla-3(RNAi) worms, suggesting that cell corpse recognition and initiation of engulfment are not disrupted (Fig. S2, O–Q). However, mtm-1(lf) phagosomes were not stained by LysoSensor green, indicating defects in phagosomal acidification (Fig. 2, C–D’ and O). Moreover, phagosomal association of LST-4, DYN-1, RAB-5, and RAB-7, which are recruited at early or late maturation stages, were all disrupted, and the lysosome reporter NUC-1 was also absent from phagosomes in mtm-1(RNAi) animals (Fig. 2, E–O; Guo et al., 2010). In contrast, phagosomal labeling by PLCδ1-PH was significantly increased in mtm-1(lf) worms (Fig. 2, A–B’ and O). These data indicate that apoptotic cell–containing phagosomes are arrested very early at the PtdIns(4,5)P2-positive stage in mtm-1(lf) worms. This is consistent with association of MTM-1 with PtdIns(4,5)P2-positive phagosomes and strongly supports the hypothesis that MTM-1 acts at the PtdIns(4,5)P2-positive stage to regulate phagocytosis of apoptotic cells.

Loss of mtm-1 affects phagosomal sealing

As PtdIns(4,5)P2 associates with unsealed but not sealed phagosomes in mammalian cells (Botelho et al., 2000; Scott et al., 2005; Sarantis et al., 2012), we investigated whether loss of mtm-1 affects phagosomal sealing, causing accumulation of PtdIns(4,5)P2-enriched unsealed phagosomes. We used FM4-64 staining, which was used previously to determine sealing of bacterium-containing phagosomes (Sarantis et al., 2012), and found that it stained the boundary but not the cytosol or nuclei of living germ cells, consistent with its inability to penetrate membranes (Fig. 3, A–B’). We examined two mutants, ced-1(lf), in which germ cell corpses are not recognized and engulfed (Zhou et al., 2001), and laat-1(lf), in which corpses are engulfed but not degraded in phagolysosomes (Liu et al., 2012). FM4-64 labeled corpses in ced-1(lf) but not laat-1(lf), indicating that fully internalized cell corpses are not accessible to FM4-64 (Fig. 3, C–D’ and L). About 50% of cell corpses were stained in gla-3(RNAi) worms, consistent with continued cell corpse clearance (Fig. 3, B, B’, and L). We found that FM4-64–positive, but not FM4-64–negative, cell corpses were stained by Annexin V, a phosphatidylserine-binding protein that selectively labels apoptotic cells, confirming that FM4-64 labels cell corpses that were not fully internalized (Fig. S3, A–C”, H, and I). FM4-64 staining correlated well with PLCδ1-PH labeling in gla-3(RNAi) and laat-1(qx42) worms, supporting the idea that PtdIns(4,5)P2 associates with unsealed but not fully separated phagosomes (Fig. 3, B–B”, D–D”, L, and M). The persistent cell corpses...
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in ced-1(lf) were not labeled by PLCδ₁-PH, confirming defects in cell corpse recognition (Fig. 3, C–C’’). Phagosomes in mtm-1(gk890934) and ocrl-1 RNAi worms were all labeled by FM4-64, Annexin V, and PLCδ₁-PH, indicating defects in phagosomal sealing (Fig. 3, E–F’’, L, and M; and Fig. S3, D–D’’, H, and I). Inactivation of OCRL-1 causes persistent cell corpses,
which are labeled by FM4-64, Annexin V, and PLCδ1-PH, indicating that PtdIns(4,5)P2 hydrolysis is required for sealing of apoptotic cell–containing phagosomes (Fig. 2, P–Q'; Fig. 3, L, and M; and Fig. S3, E–E''', H, and I). Consistent with the sealing defect, ocrl-1(lf) phagosomes were not acidified, and the recruitment of maturation factors (RAB-5 and RAB-7) and lysosomes (NUC-1) was blocked (Fig. 2, R–U' and X).

MTM-1 and PIKI-1 coordinate to control PtdIns3P levels on PtdIns(4,5)P2-positive phagosomes

To investigate how MTM-1 regulates phagosomal sealing, we followed the phagosomal dynamics of MTM-1(C378S) and 2xFYVE simultaneously. MTM-1(C378S) and 2xFYVE appeared sequentially on phagosomes, with MTM-1 release co-
inciding with 2xFYVE appearance (Fig. 4, A and F; Video 4; and Video 5). The coincidence of MTM-1 release and 2xFYVE appearance suggests that MTM-1 may hydrolyze PtdIns3P to repress its accumulation on PtdIns(4,5)P₂-positive phagosomes. To test this, we first determined whether increased MTM-1 on phagosomes affects 2xFYVE appearance. Overexpression of PPK-1 extended the PtdIns(4,5)P₂-positive stage, causing prolonged phagosomal association of MTM-1 (C378S) (Fig. 1, E–G). Similarly, RNAi inactivation of the inositol 5-phosphatase OCRL-1 led to sustained MTM-1(C378S) duration on phagosomes (Fig. 4 G). In both cases, the appearance of 2xFYVE on phagosomes was severely impaired, suggesting that sustained MTM-1 suppresses PtdIns3P accumulation (Fig. 4, B, D, and G). Consistent with this, expression of wild-type MTM-1 reduced phagosomal labeling by 2xFYVE (Fig. 4, C and E).

As sustained MTM-1 abrogates 2xFYVE appearance on phagosomes, we next examined whether loss of mtm-1 increases PtdIns3P accumulation. In mtm-1(lf) or mtm-1RNAi worms, almost all germ cell corpses were labeled by YFP::2xFYVE, whereas <50% of them were YFP positive in wild type (Fig. 5, A–C’ and G). Moreover, extending pseudopods, which are positive for PLCᵦ₁-PH and MTM-1(C378S) but not 2xFYVE in wild type, were labeled by YFP::2xFYVE in mtm-1(lf) (Fig. 5, H and J; Fig. S3, J–K”; and Video 6). C. elegans germ cell corpses are removed by gonadal sheath cells, which form a single layer covering germline components (Fig. 5 K). In wild-type sheath cells, 2xFYVE formed small puncta in the cytosol but was absent from the cell surface (Fig. 5 L). In mtm-1(lf), however, YFP::2xFYVE appeared on the surface of sheath cells, similar to PLCᵦ₁-PH, which labels plasma membranes (Fig. 5, K–K”, M, and N). This suggests that PtdIns3P accumulates on the cell surface in mtm-1(lf) sheath cells. Endosomal and phagosomal labeling by YFP::2xFYVE were also seen in mtm-1(lf) when images were taken at the top focal plane because of the extremely thin cytosol of the sheath cell layer (Fig. 5, M and N, arrowheads). The cell surface and increased phagosomal labeling of 2xFYVE in mtm-1(lf) can be suppressed by expressing wild-type but not catalytically inactive MTM-1, indicating that increased PtdIns3P accumulation is caused by loss of MTM-1 function (Fig. S3, L–P). Together, these data indicate that loss of MTM-1 leads to increased PtdIns3P accumulation, causing the appearance of 2xFYVE on cell surfaces, extending pseudopods, and phagosomes.

The class III and II PI3 kinases VPS-34 and PIKI-1 coordinate to regulate PtdIns3P generation and accumulation on phagosomes (Fig. S3, Q–U; and Table S1; Lu et al., 2012; Cheng et al., 2013). We found that loss of piki-1 but not vps-34 significantly suppressed 2xFYVE labeling on extending pseudopods and phagosomes in mtm-1(lf) (Fig. 5, D–J), and 2xFYVE labeling on pseudopods was further reduced in mtm-1::piki-1;vps-34RNAi worms (Fig. 5, D–J). Moreover, piki-1 mutation reduced cell surface labeling by 2xFYVE in mtm-1(lf) sheath cells (Fig. 5, S and U). Like MTM-1 and PLCᵦ₁-PH, GFP::PIKI-1 appeared on extending pseudopods and nascent phagosomes, and its release coincided with enrichment of 2xFYVE (Fig. 5 V). These data indicate that PIKI-1 functions at the same stage as MTM-1 to control PtdIns3P levels on plasma membranes, extending pseudopods, and nascent phagosomes. RNAi of vps-34 further reduced FYVE labeling on mtm-1::piki-1 pseudopods (Fig. 5 J), suggesting that VPS-34 may contribute to PtdIns3P production at this stage.

As in mtm-1(lf), persistent phagosomes in piki-1 or piki-1;vps-34RNAi worms were labeled by both FM4-64 and PLCᵦ₁-PH, indicating that phagosomal sealing is defective (Fig. 3, H–I’, L, and M; and Fig. S3, F–F”, H, and I). Thus, MTM-1 and PIKI-1, which control PtdIns3P levels at the PtdIns(4,5)P₂-positive stage, are both required for phagosomal sealing. Consistent with this, germ cell corpse numbers were similar in mtm-1(lf) and piki-1;vps-34RNAi and slightly increased in mtm-1::piki-1;vps-34RNAi triple mutants (Table S1).

The SH3-PX-BAR protein LST-4 acts at the PtdIns(4,5)P₂-positive stage to regulate phagosomal sealing

Scission of endocytic vesicles requires the large GTPase dynamin and its partner SNX9. In C. elegans, the SNX9 family protein LST-4 acts through DYN-1, the worm dynamin, to regulate the early stage of phagosome maturation (Almendinger et al., 2011; Chen et al., 2013). In lst-4 mutants, phagosomes are arrested at the early stage like in mtm-1(lf) (Chen et al., 2013). We found that GFP::LST-4 appeared on extending pseudopods and nascent phagosomes simultaneously with MTM-1(C378S), and its release coincided with, or preceded the enrichment of 2xFYVE on phagosomes (Fig. 6, A and B; Fig. S4 A; and Video 7 and Video 8). The germ cell corpses in lst-4(lf) and dyn-1(lf) mutants were all labeled by FM4-64 and PLCᵦ₁-PH, suggesting that phagosomal sealing is defective (Fig. 3, J–M, and Fig. S3, G–G”, H, and I). lst-4(lf) mutants accumulated similar numbers of germ cell corpses as in mtm-1(lf) or piki-1(ok2356);vps-34RNAi worms, and this phenotype was unchanged in mtm-1::lst-4 double or lst-4::piki-1;vps-34RNAi triple mutants (Fig. 6 E and Table S1). These data indicate that LST-4 acts at the PtdIns(4,5)P₂-positive stage in the same pathway with MTM-1 and PIKI-1 to regulate phagosomal sealing.

Phagosomal association of LST-4 is regulated by PtdIns(4,5)P₂, PIKI-1, and MTM-1

We next investigated whether LST-4 is regulated by PtdIns(4,5)P₂, PtdIns3P, or both. Overexpression of PPK-1 or inactivation of OCRL-1, which causes increased PtdIns(4,5)P₂ on phagosomes, led to extended LST-4 duration accompanied by defective 2xFYVE appearance (Fig. 2, V–X; Fig. 6 C; and Fig. S4, B–F). This suggests that the phagosomal dynamics of LST-4 are regulated by PtdIns(4,5)P₂ levels. As previously reported (Lu et al., 2012), loss of piki-1, which affects PtdIns3P generation, abolished phagosomal association of LST-4, suggesting that PtdIns3P is important for recruiting LST-4 to phagosomes (Fig. 6 D). Surprisingly, loss of mtm-1, which led to increased PtdIns3P accumulation on plasma membranes and forming phagosomes, also abrogated LST-4 recruitment (Fig. 2, K–O; Fig. 6 D; and Fig. S4 H). These data suggest that phagosomal association of LST-4 may require precisely controlled PtdIns3P, or both mechanisms are involved.

We attempted to test these hypotheses by examining the lipid-binding activity of LST-4 in vitro. LST-4 bound efficiently to PC liposomes containing 4% PtdIns(4,5)P₂, but not to PC liposomes containing 4% PtdIns3P (Fig. 7 A). When PtdIns3P was incorporated into PC liposomes at 20%, LST-4 binding became obvious (Fig. 7 A). The PI-binding property of LST-4 is similar to SNX9 (Lundmark and Carlsson, 2003; Yarar et al., 2008) and is consistent with it associating with PtdIns(4,5)P₂, but not PtdIns3P-enriched phagosomes in vivo. We performed liposome sedimentation assays using limited amounts of LST-4...
protein and a lower concentration of liposomes, a condition under which only minimal binding of LST-4 to PtdIns(4,5)P₂ liposomes was detected (Fig. 7 B). Addition of PtdIns3P but not PtdIns4P significantly increased binding of LST-4 to PtdIns(4,5)P₂ liposomes, suggesting that PtdIns3P may enhance LST-4 binding to PtdIns(4,5)P₂ (Fig. 7 B and Fig. S5 A). LST-4::GFP was diffuse in the cytoplasm of wild-type sheath cells but appeared on plasma membranes in mtm-1(lf) worms, which accumulate both PtdIns(4,5)P₂ and PtdIns3P on their cell membranes (Fig. 5, K–N; and Fig. S5, B–D). The plasma membrane

Figure 4. MTM-1 regulates PtdIns3P accumulation on phagosomes. (A) Time-lapse images of a cell corpse in a wild-type embryo expressing both mCHERRY::MTM-1(C378S) and GFP::2xFYVE. “0 min” represents the time point when MTM-1(C378S) was first detected around the cell corpse. The time period until the cell corpse was surrounded by MTM-1(C378S) was defined as the pseudopod extension stage. The pseudopod extension time and duration of MTM-1(C378S) and 2xFYVE were quantified. (B and C) Time-lapse images of cell corpses in wild-type embryos coexpressing mCHERRY::MTM-1(C378S), GFP::2xFYVE, and PPK-1 (B) or mCHERRY::MTM-1 and YFP::2xFYVE (C). (D and E) MTM-1(C378S) duration and 2xFYVE labeling with or without PPK-1 expression are quantified in D. MTM-1(C378S) and MTM-1 duration and 2xFYVE labeling are quantified in E. (F and G) Time-lapse images of germ cell corpses in gla-3RNAi (F) or ocr-1RNAi (G) animals coexpressing mCHERRY::MTM-1(C378S) and YFP::2xFYVE. The pseudopod extension time and duration of MTM-1(C378S) and 2xFYVE are shown beneath the images. In ocr-1RNAi animals, most phagosomes (75%, arrowheads) were not labeled by 2xFYVE during the time-lapse analysis, whereas all phagosomes followed in gla-3RNAi were positive for 2xFYVE. (A and D–G) “n” indicates the number of quantified phagosomes. (A–C, F, and G) Images in 20–25 z series were captured. Representative images are shown. Data are shown as mean ± SD. Bars, 5 µm.
Figure 5. MTM-1 and PIKI-1 coordinate to regulate PtdIns3P levels during phagosome formation. [A–G] DIC and fluorescent images of the gonads in the indicated strains expressing YFP::2xFYVE. White arrowheads and boxes indicate germ cell corpses labeled by YFP::2xFYVE, and yellow ones designate unlabeled corpses. Quantifications are shown in G. At least 15 animals were scored in each strain. [H and I] Time-lapse images of germ cell corpses in...
localization of LST-4 decreased in mtm-1(lf);piki-1;vps-34RNAi triple mutants, suggesting that PtdIns3P facilitates LST-4 binding to PtdIns(4,5)P₂-containing membranes (Fig. S5 E). Histagged LST-4 was successfully pulled down by GST–MTM-1 and GST–MTM-1(C378S) but not GST alone, suggesting that MTM-1 interacts with LST-4 and the interaction is independent of MTM-1 phosphatase activity (Fig. 7 C). The PX-BAR region, but not the SH3 domain of LST-4, was coprecipitated with FLAG-tagged MTM-1, indicating that MTM-1 interacts with LST-4 through the PX-BAR domain (Fig. 7 D). GFP::LST-4 FLAG-tagged MTM-1, indicating that MTM-1 interacts with LST-4 and the interaction is independent of apoptotic cell–containing phagosomes. PtdIns(4,5)P₂ association with, and requires timely depletion of, PtdIns(4,5)P₂. The inositol 5-phosphatase OCRL-1, which contributes to PtdIns(4,5)P₂ elimination in mammalian phagocytosis (Bohdanowicz et al., 2012), is important for this process. Loss of ocrl-1 leads to prolonged association of MTM-1, LST-4, and DYN-1 with unsealed phagosomes, suggesting that defective sealing may be caused by sequestration of sealing factors by PtdIns(4,5)P₂-enriched membranes. Our finding is consistent with a recent study that shows OCRL is required for dissociation of SNX9 from endocytic vesicles during uncoating (Náñez et al., 2014).

Our data indicate that PtdIns3P is produced on plasma membranes and forming phagosomes where PtdIns(4,5)P₂ is enriched, but its accumulation is suppressed by the PtdIns(4,5)P₂ effector MTM-1. Loss of MTM-1 causes PtdIns3P accumulation on plasma membranes, extending pseudopods, and nascent phagosomes and numerous unsealed PtdIns(4,5)P₂-positive phagosomes. Inactivation of the PI3 kinases PIKI-1 and VPS-34 completely suppresses PtdIns3P accumulation caused by mtm-1 mutation but also leads to defects in phagosomal sealing, indicating that PtdIns3P must be tightly controlled in this process. Consistent with this, the 2xFYVE reporter is only visible on plasma or unsealed phagosomal membranes in mtm-1(lf) but not wild-type worms. The timely depletion of PtdIns(4,5)P₂ by OCRL-1 may release MTM-1, LST-4, and DYN-1, which completes the scission process and allows subsequent accumulation of PtdIns3P on fully sealed phagosomes (Fig. 7 F). In this way, phagosomal sealing and the switch of membrane identity from unsealed (PtdIns(4,5)P₂ enriched) to sealed (PtdIns3P enriched) phagosomes are coregulated and precisely coupled. More work is needed to determine how OCRL-1 is regulated when apoptotic cell–containing phagosomes are being sealed.

**Discussion**

Sealing of apoptotic cell–containing phagosomes involves both PtdIns(4,5)P₂ and PtdIns3P

PtdIns(4,5)P₂ and PtdIns3P are key determinants of phagosome formation and maturation, respectively (Bohdanowicz and Grinstein, 2013). Here, we found that both are involved in closure of apoptotic cell–containing phagosomes. PtdIns(4,5)P₂ associates with extending pseudopods and unsealed but not sealed phagosomes. Inhibition of PtdIns(4,5)P₂ hydrolysis by ocrl-1 RNAi or its increased generation by PPK-1 overexpression causes defective clearance of apoptotic cells and accumulation of unsealed phagosomes. Thus, like mammalian phagocytosis, sealing of apoptotic cell–containing phagosomes coincides with, and requires timely depletion of, PtdIns(4,5)P₂. The inositol 5-phosphatase OCRL-1, which contributes to PtdIns(4,5)P₂ elimination in mammalian phagocytosis (Bohdanowicz et al., 2012), is important for this process. Loss of ocrl-1 leads to prolonged association of MTM-1, LST-4, and DYN-1 with unsealed phagosomes, suggesting that defective sealing may be caused by sequestration of sealing factors by PtdIns(4,5)P₂-enriched membranes. Our finding is consistent with a recent study that shows OCRL is required for dissociation of SNX9 from endocytic vesicles during uncoating (Náñez et al., 2014).

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The PtdIns(4,5)P₂ effector MTM-1 coordinates with PIKI-1 to control PtdIns3P levels on unsealed phagosomes

Previously, we and others reported that MTM-1 localizes to plasma membranes and negatively regulates cell corpse engulfment through the CED-5/12/10 module (Zou et al., 2009; Neukomm et al., 2011). Here, we found that MTM-1 associates with forming phagosomes to promote sealing. MTM-1 binds specifically to PtdIns(4,5)P₂, and its membrane association is regulated by the latter. Thus, MTM-1 acts as an effector of PtdIns(4,5)P₂ to regulate both initiation and completion of engulfment. The acceleration in the initiation step but impairment in the sealing of phagosomes caused by loss of mtm-1 suggest that PtdIns3P turnover may have distinct effects on apoptotic cell clearance at different stages, a phenomenon that is also observed in autophagy regulation (Vergne et al., 2009; Taguchi-Arashiki et al., 2010; Cebollero et al., 2012; Wu et al., 2014).

Our data indicate that PIKI-1, the class II PI3 kinase, acts with MTM-1 to control a subpool of PtdIns3P on plasma membranes and unsealed phagosomes for cell corpse removal. The class III PI3 kinase VPS-34 may play a major role in producing PtdIns3P on sealed phagosomes, as both MTM-1 and PIKI-1 are released at this stage. Human class II PI3 kinase C2α is reported to control clathrin-mediated endocytosis by generating PtdIns(3,4)P₂ (Posor et al., 2013). In our study, however, 2xFYVE but not TAPPI1, a PtdIns(3,4)P₂ reporter, is readily seen on phagosomes and abrogated by loss of PIKI-1. Loss or overexpression of R01H10.7, the C. elegans homologue of INPP4A, which converts PtdIns(3,4)P₂ to PtdIns3P, has no effect on cell corpse removal. Moreover, TAPPI1 was not observed on phagosomes in r01h10.7(fj) or r01h10.7(fj);mtm-1RNAi worms (Fig. S1, Y and Z). Thus, PtdIns(3,4)P₂ is probably not involved in apoptotic cell clearance either directly or as a precursor of PtdIns3P.

The PI3 kinases PIKI-1 and VPS-34 were reported to regulate phagosome maturation by acting sequentially to control PtdIns3P oscillation on phagosomes, a process antagonized by MTM-1 (Lu et al., 2012). By examining >100 embryonic cell corpses, however, we found that PtdIns3P oscillation is not...
Figure 6. LST-4 acts at the PtdIns(4,5)P2-positive stage, and its dynamics are regulated by PtdIns(4,5)P2, PIKI-1, and MTM-1. (A) Time-lapse images of a cell corpse in a wild-type embryo coexpressing mCHERRY::MTM-1(C378S) and LST-4::GFP. (B and C) Time-lapse images of germ cell corpses in a gla-3RNAi (B) or an ocrl-1RNAi (C) animal coexpressing LST-4::mCHERRY and YFP::2xFYVE. (D) Time-lapse images of germ cell corpses in the indicated strains.
readily observed on cell corpses other than the three designated corpses (C1, C2, and C3) on which two waves of 2xFYVE were previously observed and characterized. Thus, PtdIns3P oscillation may associate with clearance of specific corpses like C1, C2, and C3. MTM-1(C378S) and 2xFYVE appear sequentially on the C3 phagosome, and MTM-1 release coincides with 2xFYVE enrichment (Fig. S5 H and Video 9), confirming that MTM-1 acts before PtdIns3P accumulation on phagosomes. MTM-1 remains absent when 2xFYVE disappears and reappears on the C3 phagosome, indicating that MTM-1 is probably not involved in regulating PtdIns3P oscillation on the C3 phagosome (Fig. S5 H and Video 9).

The copresence of PtdIns(4,5)P2, PtdIns3P, and MTM-1 recruits LST-4 to unsealed phagosomes

LST-4 was reported as a PtdIns3P effector that promotes phagosome maturation (Lu et al., 2010). However, we found that LST-4 associates with phagosomes at the PtdIns(4,5)P2-positive stage and is released before PtdIns3P accumulates. This is consistent with the lipid-binding property of LST-4 and two studies showing that LST-4 acts earlier than RAB-5 in cell corpse removal (Almendinger et al., 2011; Chen et al., 2013). We observed a large number of unsealed phagosomes in worms lacking LST-4 or DYN-1, indicating essential roles of these proteins in the sealing process. Thus, LST-4 may act with DYN-1 to control scission of apoptotic cell–containing phagosomes as in endocytosis.

Our data strongly support the hypothesis that PtdIns(4,5)P2, PtdIns3P, and MTM-1 serve as a coincidence detection code on unsealed phagosomes to recruit LST-4 for the sealing of apoptotic cell–containing phagosomes (Fig. 7 F). LST-4 binds efficiently to PtdIns(4,5)P2 liposomes in vivo and associates with PtdIns(4,5)P2- but not PtdIns3P-enriched phagosomes in vivo, indicating an essential role of PtdIns(4,5)P2 in membrane recruitment of LST-4. Loss of the PtdIns3P-producing kinase PIKI-1 or the LST-4–interacting PtdIns3P phosphatase MTM-1 disrupts phagosomal association of LST-4, causing accumulation of unsealed phagosomes. This indicates that PtdIns3P is important for LST-4 recruitment and its levels may be tightly controlled during sealing. Expression of MTM-1(C378S), which interacts with LST-4 but lacks catalytic activity, failed to rescue the defective phagosomal association of LST-4 in mthm-1(lf);piki-1(lf) double mutants (Fig. S5, F–G‴), further supporting the role of PtdIns3P in LST-4 recruitment to phagosomes. Addition of PtdIns3P, which by itself displays limited affinity to LST-4, significantly enhances LST-4 binding to PtdIns(4,5)P2 liposomes in vitro, suggesting that PtdIns3P may facilitate or stabilize the binding of LST-4 with PtdIns(4,5)P2. Our results are consistent with a recent study that shows SNX9 is selectively recruited by coincidence of PtdIns(4,5)P2 and PtdIns3P on curved liposomal membranes (Gallop et al., 2013). Binding of SNX9 to liposomes is significantly increased by the combination of PtdIns(4,5)P2 and PtdIns3P compared with PtdIns(4,5)P2 alone and PtdIns3P alone (Gallop et al., 2013). We found that LST-4 interacts with MTM-1 through the lipid-binding PX-BAR domain but not the protein–protein-interacting SH3 domain, raising the possibility that binding of LST-4 to MTM-1 and lipids may be mutually regulated. However, more work is needed to uncover how PtdIns3P and/or MTM-1 may modulate LST-4 binding to PtdIns(4,5)P2 liposomes or phagosomes.

The PtdIns(4,5)P2–PtdIns3P–MTM-1–LST-4 regulatory module may be used in mammals

When expressed in worms, human MTM1 localizes to plasma membranes and efficiently rescues the persistent cell corpse phenotype of mthm-1(lf), suggesting evolutionarily conserved roles of myotubularin in this process. Mutations in MTM1, BIN1 (which encodes an N-BAR and SH3 domain–containing protein), and DNM2 lead to X-linked, autosomal recessive, and autosomal dominant centronuclear myopathies, respectively, and cause similar tubule defects in skeletal muscle, suggesting that they may act in a common pathway important for muscle function (Cowling et al., 2012). MTM1 interacts with BIN1 in skeletal muscle and enhances BIN1-mediated membrane tubulation dependent on both binding and phosphatase activity (Royer et al., 2013). A PI-binding motif, which is present only in the muscle isoform of BIN1, facilitates PtdIns(4,5)P2 binding and promotes membrane tubulation (Lee et al., 2002). Thus, the PtdIns(4,5)P2–PtdIns3P–MTM-1–BAR regulatory module may play a conserved role to maintain muscle function and could be relevant to pathological alterations in centronuclear myopathies.

Materials and methods

C. elegans strains

Strains of C. elegans were cultured at 20°C using standard protocols (Brenner, 1974). The N2 Bristol strain was used as the wild-type strain. The following strains were used in this work: linkage group (LG) I, mthm-1(gk890934), mthm-1(qx322), and ced-1(e1735); LG II, laat-1(qx42), age-1(hx546), and age-1(mg44); LG III, R01H10.7(ok489); LG IV, daf-18(ok480) and let-4(m2423); LG X, piki-1(ok2346) and dyn-1(ky51). age-1(mg44) was maintained as mnh1/age-1, and the homozygous progenies were examined. dyn-1(ky51) was maintained at 20°C, and L4 larvae were moved to 25°C for 36 h before germ cell corpses were examined.

mthm-1(qx322) was isolated from a genetic screen by defective clearance of residual bodies. qx322 contains a C to T transition that results in an early stop codon after Lys99 and is probably a null mutation of mthm-1. mthm-1(gk890934), which was provided by the Caenorhabditis Genetics Center, contains the amino acid substitutions V188M and R229K.

mtm-1(qx322) was out-crossed with the wild-type N2 strain six and four times, respectively, before further analysis.

The reporter strains used in this study are listed below. Transgenic animals carrying extrachromosomal arrays (qEx) were generated using standard microinjection methods, and genome-integrated arrays (qxd) were obtained by γ irradiation to achieve stable expression from arrays with low copy numbers (Evans, 2006).
Figure 7. **LST-4 interacts with PtdIns(4,5)P2, PtdIns3P, and MTM-1.** (A and B) Binding of His-tagged LST-4 to the indicated liposomes was detected by Coomassie blue staining (A) or Western blotting (B). The lipid–protein complex is recovered in the pellet (P) but not the supernatant (S). The final concentration of LST-4 and liposomes used in each experiment is indicated. Three (A) and six (B) independent experiments were performed. Data (shown as mean ± SD) were compared by one-way ANOVA with Tukey’s post-test. *, P < 0.05; **, P < 0.0001. (C) His-tagged LST-4 is pulled down by GST::MTM-1::FLAG and GST::MTM-1(C378S)::FLAG but not GST::FLAG. Purified GST fusion proteins stained by Coomassie blue (arrows) are shown underneath. (D) Full-length LST-4 and the PX-BAR region, but not the SH3 domain, are coprecipitated from HEK293T cells by MTM-1. (E) mCHERRY::MTM-1(C378S) is coprecipitated on August 22, 2017 jcb.rupress.org Downloaded from.
Quantification of cell corpses and phagosome maturation

The number of somatic cell corpses in the head region of living embryos and the number of germ cell corpses in one gonad arm at various adult ages (24 or 48 h after L4/adult molt) were scored as described previously (Gumienny et al., 1999; Wang et al., 2002). The cell corpses were identified by their raised-button–like morphology using Nomarski optics. At least 15 animals were scored at each stage in each strain. To study apoptotic cell clearance in C. elegans germlines, gla-3 RNAi was performed to increase germ cell apoptosis without affecting cell corpse removal (Kritikou et al., 2006), thus allowing examination of more phagosomes near the gonad loop. To induce gene expression by heat-shock treatment, L4 larval worms were picked to fresh nematode growth medium plates and treated at 36°C. At least 15 animals were scored in each strain.

Visualization of plasma membrane labeling by 2xFYVE in gonadal sheath cells

Adult worms (48 h after L4/adult molt) were dissected to expose their gonads in 200 µl gonad dissection buffer with or without 2 µl Alexa Fluor 488–conjugated Annexin V (Invitrogen; Wang et al., 2007). 155 µl gonad dissection buffer was removed, and 5 µl FM4-64 was added to make a final concentration of 200 ng/ml. Staining was performed at 4°C for 5 min before examination by fluorescence microscopy. At least 10 animals were scored in each strain. The fluorescence intensity of Annexin V conjugates varied general, at least 10 phagosomes from three independent animals were followed and quantified in each strain.

FM4-64, Annexin V, and LysoSensor staining

Adult worms (36 or 48 h after L4/adult molt) were dissected to expose their gonads in 200 µl gonad dissection buffer with or without 2 µl Alexa Fluor 488–conjugated Annexin V (Invitrogen; Wang et al., 2007). 155 µl gonad dissection buffer was removed, and 5 µl FM4-64 (200 µg/ml; Invitrogen) was added to make a final concentration of 20 µg/ml. Staining was performed at 4°C for 5 min before examination by fluorescence microscopy. At least 10 animals were scored in each strain. The fluorescence intensity of Annexin V conjugates varied on different apoptotic cells, which is likely the result of differences in phosphosserine levels on cell corpses that persist for different periods of time. LysoSensor staining was performed as described previously with some modifications (Guo et al., 2010). In brief, adult animals (48 h after L4/adult molt) were dissected in gonad dissection buffer with 1-µM LysoSensor green DND-189 (Invitrogen) and examined by fluorescence microscopy.

by LST-4::GFP in lysates prepared from mix-staged worms coexpressing both fusion proteins. (f) Proposed model of coregulation of phagosomal sealing and PI conversion by the PtdIns[4,5]P2–PtdIns3P–MTM-1–LST-4 module in apoptotic cell clearance. Dashed arrows indicate regulations that need further investigation. IB, immunoblotting; IP, immunoprecipitation.
In the required molar ratio was prepared in chloroform and dried under nitrogen. Lipid films were hydrated in a buffer containing 50-nM Tris-HCl, pH 7.5, and 150-mM NaCl with a 1-mM total lipid concentration. Unilamellar vesicles (100 nm) were generated using a mini extruder (Avanti Polar Lipids, Inc.). For sedimentation assays, 2.5–30 µg proteins were incubated with 4–50 µl (1 mM) liposomes in a 60-µl reaction at room temperature for 10 min before sedimentation at 60,000 rpm for 40 min in a rotor (TLA-100; Beckman Coulter). Both supernatant and pellet were resolved with SDS-PAGE and revealed by Coomassie blue staining or Western blotting. At least three independent experiments were performed for all liposome sedimentation assays.

**Liposome flotation assay**

1,2-dioleoyl-sn-glycero-3-PC alone or mixed with other phospholipids (1,2-dioleoyl-sn-glycero-3-phospho-l-serine, PtdIns3P, PtdIns4P, PtdIns(3,4)P₂, PtdIns(3,4,5)P₃; Avanti Polar Lipids, Inc.) in a molar ratio of 9:1 was prepared in chloroform and dried under nitrogen. Lipid films were hydrated in TBS buffer containing 50-mM Tris-HCl, pH 7.5, and 150-mM NaCl with a 1-mM total lipid concentration. For flotation assays, 6–18 µg proteins were incubated with 90 µl (1 mM) liposomes in a 100-µl reaction at room temperature for 15 min. A 900-µl 50% (wt/vol) sucrose solution in TBS was added to the protein-liposome mixtures to a final concentration of 45% (wt/vol). A 2.2-ml 30% (wt/vol) sucrose solution in TBS was layered on the samples, followed by 1 ml TBS layered on top. The samples were then subjected to ultracentrifugation at 45,000 rpm for 5 h in a rotor (SW 60 Ti; Beckman Coulter). The top fraction containing floating liposomes (100 µl) was recovered, and half of it (50 µl) was resolved with SDS-PAGE and revealed by Coomassie blue staining. At least three independent experiments were performed for each assay, and representative results are shown.

**RNAi**

The bacteria-feeding protocol was used in RNAi experiments. In each experiment, 3–10 L4 larvae (P0) were cultured on the RNAI plate, and F1 progeny at the L4 stage were transferred to fresh RNAI plates and aged for 24, 36, or 48 h before examination of germ cell corpses.

The DNA sequences that are targeted by the double-stranded RNAs in the RNAi experiments are as follows: **gla-3** (T02E1, 12,950–13,970 nt), **mtm-1** (Y110A7A, 35,061–37,303 nt), **ocr-1** (C16C2, 2,650–4,215 nt), and **vps-34** (B0025, 10,243–11,439 nt; Table S3).

**Phosphatase assay**

The phosphatase activity of MTM-1 was assayed using a Malachite green assay kit (Echelon Biosciences, Inc.) by measuring the liberated phosphate after the phosphatase reaction. Experiments were performed according to the manufacturer’s instructions with some modifications. Free lipid was incubated with 1,000 pmol of substrate (DiC8 PtdIns3P, DiC8 PtdIns(3,4)P₂, DiC8 PtdIns(3,4,5)P₃, Echelon Biosciences, Inc.) in a 25-µl reaction containing 1-mM total phosphate standard for 20 min at room temperature, and absorbance was measured at 620 nm. Averages of triplicate samples were used to draw the phosphate standard curve and determine the free phosphate in each reaction and control.

**GST pull-down assay**

GST::FLAG, GST::MTM-1::FLAG, or GST::MTM-1(C378S):FLAG immobilized on glutathione-agarose beads was incubated with BL21 bacterial lysates containing His-tagged LST-4 for 2 h at 4°C. The beads were washed extensively with PBS, and the bound proteins were resolved with SDS-PAGE and revealed by Western blotting with an antibody against the hexahistidine tag.

**Coimmunoprecipitation assay in worms**

Mix-staged worms expressing both LST-4::GFP and mCHERRY::MTM-1(C378S), or mCHERRY::MTM-1(C378S) alone, were lysed with a tissue grinder in worm lysis buffer containing 50-mM Tris-HCl, pH 7.5, 150-mM NaCl, 1% Triton X-100, and protease inhibitor cocktail (EDTA free; Roche). The homogenized solutions were kept on ice for 15 min and centrifuged at 14,000 rpm for 20 min at 4°C to remove the debris. 400 µl of precleared solution was incubated with 1 µl GFP antibody (polyclonal antibody raised in rabbits against recombinant GFP; C. Yang laboratory, Institute of Genetics and Developmental Biology, Chinese Academy of Sciences, Beijing, China) for 2 h at 4°C, followed by incubation with 20 µl BSA-blocked protein G–Sepharose (Invitrogen) for 2 h. After extensive washing with 50-mM Tris-HCl, pH 7.5, and 250-mM NaCl, the bound proteins were resolved with SDS-PAGE and revealed by Western blotting with an antibody against mCHERRY (mouse, HX1810; Huaxingbio, Inc.).

**Statistical analysis**

The SD was used as y axis error bars for bar charts plotted from the mean value of the data. Data derived from different genetic backgrounds were compared by Student’s two-way unpaired t test, one-way analysis of variance (ANOVA) followed by Tukey’s post-test, or two-way ANOVA followed by Bonferroni post-test, as indicated in the figure legends. Data were considered statistically different at P < 0.05, * P < 0.05; **, P < 0.0001 (0.001 in a two-way ANOVA analysis).

**Plasmid construction**

To generate Pppk-1::mCHERRY::PLCδ1::GFP::PLCδ1-PH, PLCδ1-PH was amplified from a human cDNA library using primers PXL249/PZH42 and was cloned into pPD49.26-P_ced-1_mCHERRY1 and pPD49.26-P_ppk-1::GFP, respectively, through the KpnI site. To construct Pppk-1::PPK-1 and Pppk-1::PPK-1::GFP, the full-length cDNA of pppk-1 was amplified from a C. elegans cdNA library (Invitrogen) by PHBW400/PROM5 and PHBW400/PSYC50, respectively, and ligated into pPD49.26-P_ced-1_mCHERRY1 and pPD49.26-P_ppk-1::GFP. To generate Pppk-1::mCHERRY::MTM-1, Pppk-1::PLCδ1::GFP, and Pppk-1::mCHERRY::MTM-1(C378S) were constructed by ligating the mCHERRY genomic sequence or the C378S mutant form into pPD49.26-P_ced-1_mCHERRY1 and pPD49.26-P_ppk-1::GFP. To generate Pppk-1::TAPP1::GFP, TAPP1 was amplified from a human cDNA library using primers PSYC554/56 and cloned into pPD49.26-P_ced-1.
To construct pET41b-GST::MTM-1(C378S)::FLAG, GST::FLAG was amplified using primers PWDL108/PSYC443 and ligated into pET41b through SpeI–HindIII sites. To construct pET41b-MTM-1::HIS6, the cDNA was amplified from MTM-1 genomic DNA using primers PSYC501/502 were ligated into pPD49.78 and cloned into pPD49.26-GFP3 through the KpnI site. To generate pET21b-PLC1-PH::HIS6, PLC1-PH was amplified from a plasmid containing phagosomes. Video 9 shows that MTM-1 and PtdIns(4,5)P2 display identical dynamics on extending pseudopods and forming phagosomes. Video 1 shows dynamic changes of PtdIns(4,5)P2 and PtdIns3P on apoptotic cell–containing phagosomes. Video 5 shows that LST-4 release coincides with PtdIns3P depletion precedes phagosomes formation. Video 3 shows that MTM-1 and PtdIns(4,5)P2 display identical dynamics on extending pseudopods and forming phagosomes. Video 2 shows that PtdIns3P depletion precedes phagosomes formation. Video 3 shows that MTM-1 and PtdIns(4,5)P2 display identical dynamics on extending pseudopods and forming phagosomes.

Online supplemental material

Fig. S1 shows that PtdIns(3,4)P2 and PtdIns(3,4,5)P3 do not accumulate significantly on apoptotic cell–containing phagosomes. Fig. S2 shows that MTM-1 acts at the PtdIns(4,5)P2-positive stage to regulate apoptotic cell clearance. Fig. S3 shows that MTM-1 regulates PtdIns3P levels on plasma membranes, extending pseudopods, and phagosomes. Fig. S4 shows that phagosomal association of LST-4 is regulated by PtdIns(4,5)P2–containing membranes. Video 1 shows dynamic changes of PtdIns(4,5)P2 and PtdIns3P on apoptotic cell–containing phagosomes. Video 2 shows that PtdIns3P depletion precedes phagosomes formation. Video 3 shows that MTM-1 and PtdIns(4,5)P2 display identical dynamics on extending pseudopods and forming phagosomes. Videos 4 and 5 show that MTM-1 release coincides with PtdIns3P accumulation on phagosomes containing somatic (Video 4) or germ cell corpses (Video 5). Video 6 shows that loss of MTM-1 causes PtdIns3P accumulation on extending pseudopods and forming phagosomes. Video 7 shows that LST-4 and MTM-1 display identical phagosomal dynamics. Video 8 shows that LST-4 release coincides with PtdIns3P accumulation on phagosomes. Video 9 shows that MTM-1 release coincides with PtdIns3P accumulation on the C3 phagosome.

Table S1 shows that LST-4 acts in the same pathway with MTM-1 and PIK-I to regulate apoptotic cell clearance. Table S2 lists primers that are used for plasmid construction. Table S3 lists double-stranded RNAs used for RNAi experiments. Online supplemental material is available at http://www.jcb.org/cgi/content/full/jcb.201501038/DC1.

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References

Almendinger, J., K. Doukoumetzidis, J.M. Kinchen, A. Kaech, K.S. Ravichandran, and M.O. Hengartner. 2011. A conserved role for SNX9-family members in the regulation of phagosome maturation during engulfment of apoptotic cells. PLoS ONE. 6:e18325. http://dx.doi.org/10.1371/journal.pone.0018325.

Balla, T. 2013. Phosphoinositides: tiny lipids with giant impact on cell regulation. Physiol. Rev. 93:1019–1137. http://dx.doi.org/10.1152/physrev.00028.2012.

Bohdanowicz, M., and S. Grinstein. 2013. Role of phospholipids in endocytosis, phagocytosis, and macroautophagy. Physiol. Rev. 93:69–106. http://dx.doi.org/10.1152/physrev.00002.2012.

Bohdanowicz, M., D.M. Balkin, P. De Camilli, and S. Grinstein. 2012. Recruitment of OCR1 and Inpp5B to phagosomes by Rab5 and APPL1 depletes phosphoinositides and attenuates Akt signaling. Mol. Biol. Cell. 23:176–187. http://dx.doi.org/10.1091/mbc.E11-06-0489.
Distinct roles of class I and class III phosphatidylinositol 3-kinases in phagosome formation and maturation. J. Cell Biol. 155:19–26. http://dx.doi.org/10.1083/jcb.200107069

Wang, X., C. Yang, J. Chai, Y. Shi, and D. Xue. 2002. Mechanisms of AIF-mediated apoptotic DNA degradation in Caenorhabditis elegans. Science. 298:1587–1592. http://dx.doi.org/10.1126/science.1076194

Wang, X., Y.C. Wu, V.A. Fadok, M.C. Lee, K. Gengyo-Ando, L.C. Cheng, D. Ledwich, P.K. Hsu, J.Y. Chen, B.K. Chou, et al. 2003. Cell corpse engulfment mediated by C. elegans phosphatidyserine receptor through CED-5 and CED-12. Science. 302:1563–1566. http://dx.doi.org/10.1126/science.1087641

Wang, X., J. Wang, K. Gengyo-Ando, L. Gu, C.L. Sun, C. Yang, Y. Shi, T. Kobayashi, Y. Shi, S. Mitani, et al. 2007. C. elegans mitochondrial factor WAH-1 promotes phosphatidyserine externalization in apoptotic cells through phospholipid scramblase SCRM-1. Nat. Cell Biol. 9:541–549. http://dx.doi.org/10.1038/ncb1574

Wu, Y., S. Cheng, H. Zhao, W. Zou, S. Yoshina, S. Mitani, H. Zhang, and X. Wang. 2014. PI3P phosphatase activity is required for autophagosome maturation and autolysosome formation. EMBO Rep. 15:973–981. http://dx.doi.org/10.15252/embr.201438618

Yarar, D., M.C. Surka, M.C. Leonard, and S.L. Schmid. 2008. SNX9 activities are regulated by multiple phosphoinositides through both PX and BAR domains. Traffic. 9:133–146. http://dx.doi.org/10.1111/j.1600-0854.2007.00675.x

Yu, X., S. Odera, C.H. Chuang, N. Lu, and Z. Zhou. 2006. C. elegans dynamin mediates the signaling of phagocytic receptor CED-1 for the engulfment and degradation of apoptotic cells. Dev. Cell. 10:743–757. http://dx.doi.org/10.1016/j.devcel.2006.04.007

Zhou, Z., E. Hartwig, and H.R. Horvitz. 2001. CED-1 is a transmembrane receptor that mediates cell corpse engulfment in C. elegans. Cell. 104:43–56. http://dx.doi.org/10.1016/S0092-8674(01)00190-8

Zou, W., Q. Lu, D. Zhao, W. Li, J. Mapes, Y. Xie, and X. Wang. 2009. Caenorhabditis elegans myotubularin MTM-1 negatively regulates the engulfment of apoptotic cells. PLoS Genet. 5:e1000679. http://dx.doi.org/10.1371/journal.pgen.1000679