Initial phospholipid-dependent Irgb6 targeting to Toxoplasma gondii vacuoles mediates host defense

Youngae Lee1,3, Hiroshi Yamada5, Ariel Pradipta1, Ji Su Ma1,3, Masaki Okamoto1, Hikaru Nagaoka6, Eizo Takashima6, Daron M Standley2,4, Miwa Sasai1,3, Kohji Takei5, Masahiro Yamamoto1,3

Toxoplasma gondii is an obligate intracellular protozoan parasite capable of infecting warm-blooded animals by ingestion. The organism enters host cells and resides in the cytoplasm in a membrane-bound parasitophorous vacuole (PV). Inducing an interferon response enables IFN-γ–inducible immunity-related GTPase (IRG protein) to accumulate on the PV and to restrict parasite growth. However, little is known about the mechanisms by which IRG proteins recognize and destroy T. gondii PV. We characterized the role of IRG protein Irgb6 in the cell-autonomous response against T. gondii PV. We show that Irgb6 is capable of binding a specific phospholipid on the PV membrane. Furthermore, the absence of Irgb6 causes reduced targeting of other effector IRG proteins to the PV. This suggests that Irgb6 has a role as a pioneer in the process by which multiple IRG proteins access the PV. Irgb6-deficient mice are highly susceptible to infection by a strain of T. gondii avirulent in wild-type mice.

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Introduction

Healthy mammalian hosts activate immune responses against pathogenic infections. The innate immune response first induces IL-12 production by antigen-presenting cells, such as macrophages and dendritic cells. This is carried out via recognition of pathogen-derived components by microbe pattern recognition receptors such as toll-like receptors (Hunter & Remington, 1995; Yarovinsky & Sher, 2006). IL-12 subsequently stimulates the antipathogen type 1 immune response, wherein naïve CD4+ or CD8+ T cells become antigen-specific Th1 cells and cytotoxic T cells, respectively, with the help of antigen-presenting cells. Th1 cells, cytotoxic T lymphocytes, and natural killer cells produce IFN-γ to activate the various cell-autonomous programs targeting vacuolar pathogens (Suzuki et al, 1988; Gazzinelli et al, 1991). One of the IFN-γ–induced cell-autonomous programs is associated with IFN-γ–inducible GTPases, such as p47 immunity-related GTPases (IRGs) and p65 guanylate-binding proteins (GBPs) (Kim et al, 2012). Immunity-related GTPases and GBPs belong to the dynamin GTPase superfamily (Martens & Howard, 2006; Pawlowski, 2010; Kim et al, 2012) and can target wide ranges of bacterial, fungal, and protozoan vacuolar pathogens (Coers et al, 2008; Al-Zeer et al, 2009; Ferreira-da-Silva Mda et al, 2014; Kurikose & Kanneganti, 2017; Santos & Broz, 2018). In mice, the IRG protein family consists of three regulator IRG proteins (Irgm1, Irgm2, and Irgm3) and over 20 effector IRG proteins and decoys (Bekpen et al, 2005; Muller & Howard, 2016). There are four effector IRG proteins known to be expressed in mice: Irga6, Irgb6, Irgb10, and Irgd (Martens & Howard, 2006). Regulator IRG proteins harboring GX4GXS in the first nucleotide-binding motif (G1) are mainly associated with host endomembranes, such as the Golgi apparatus and ER (Bekpen et al, 2005; Hunn et al, 2011). Effector IRG proteins possess a universally conserved GX4GKS sequence in the G1 motif, enabling binding to both GTP and GDP (Taylor et al, 1996; Uthaiah et al, 2003; Bekpen et al, 2005; Hunn et al, 2008). The GTPase activity has been demonstrated for Irga6 and Irgm3 (Taylor et al, 1996; Uthaiah et al, 2003; Hunn et al, 2008). Regulator IRG proteins can maintain effector IRG proteins in an inactive GDP-bound state, potentially preventing the latter from inappropriate activation on host cell membrane–bound vesicular systems. In their absence, effector IRG proteins likely form GTP–GDP aggregates and are unable to interact with the Toxoplasma gondii parasitophorous vacuole (PV) (Martens et al, 2004; Hunn et al, 2008; Hunn & Howard, 2010; Coers, 2013; Haldar et al, 2013). There are 11 members in the mouse GBP family, all of which have the conserved GTP binding motifs (Kresse et al, 2008). Guanylate-binding protein mutants lacking GTPase activity are incapable of accumulating at T. gondii PV membrane (PVM) (Degrandi et al, 2013; Ohshima et al, 2015). When these IFN-inducible GTPases are recruited to the PVM, it becomes vesiculated and disrupted, resulting in death of the vacuolar pathogen (Martens et al, 2005; Ling et al, 2006; Degrandi et al, 2007; Virreira Winter et al, 2011; Yamamoto

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et al, 2012; Selleck et al, 2013). Thus, GTPase activity–dependent IRG and GBP accumulation is well established as important for cell-autonomous immunity to vacuolar pathogens.

The mechanism by which IRG proteins access T. gondii PV from the cytosolic compartments can be passive. This process depends on diffusion from the cytoplasmic pools rather than active transport involving toll-like receptor–mediated signaling pathways or microtubule networks (Khaminets et al, 2010). Although IRG proteins are localized on the PVM within a few minutes of T. gondii infection (Hunn et al, 2008; Khaminets et al, 2010), little is known about the mechanism by which IRG proteins recognize and destroy the PVM thus far. This process is important for IFN-γ–induced cell-autonomous immunity. Among the effector IRG proteins, Irgb6 and Irgb10 are loaded first and most efficiently onto T. gondii PVM (Khaminets et al, 2010).

Here, we aimed to determine the role of Irgb6 in the cell-autonomous response against T. gondii, mediating ubiquitination and disruption of T. gondii PVM.

Results

Irgb6 contributes to IFN-γ–induced cell-autonomous resistance to T. gondii

Several studies using gene-deficient mice (Taylor et al, 2000, 2007; Liesenfeld et al, 2011) have shown that Irgm1 (also called LRG-47), Irgm3 (IGTP), Irga6 (IIGP, IIGP1), and Irgd (IRG-47) have critical roles in the anti–T. gondii response. Earlier studies have shown that Irgb6 and Irgb10 proteins function as pioneers, loading first to the T. gondii vacuoles (Khaminets et al, 2010). However, there are no reports clearly indicating the presence of the Irgb4 gene (Bekpen et al, 2005) (Fig S2B).

Together, these results show that Irgb6 is required for the IFN-γ–induced cell-autonomous response in vitro.

Irgb6 is required for recruitment of ubiquitin and other IFN-inducible GTPases to T. gondii PVM

We next examined whether Irgb6 is required for normal loading of other effector IRG proteins and GBPs onto the T. gondii PVM using immunoﬂuorescence staining (Fig 2A–E). Compared with wild-type cells, Irgb6-deficient MEFs displayed signiﬁcantly reduced accumulation of Irga6, Gbp1, Gbp2, Gbp1-5, ubiquitin, and p62 (Fig 2D). Altogether, these results show that Irgb6 is required for the IFN-γ–induced anti–T. gondii cell-autonomous response in vitro.

Other effector IRG proteins are not involved in Irgb6 and ubiquitin loading onto T. gondii vacuoles

In response to T. gondii infection, Irgb6 and Irgb10 have been shown to be initially localized on the PVM within a few minutes of infection.
Figure 1. Irgb6 significantly contributes to IFN-γ–induced cell-autonomous T. gondii killing.

(A) Schematic representation of the gene-targeting strategy for mouse Irgb6* and Irgb6 locus by Cas9-mediated genome editing. (B) Western blot analysis of the indicated protein expressions in WT and Irgb6 KO MEFs after IFN-γ stimulation or not. (C, D) Survival rate of T. gondii in the presence of IFN-γ stimulation relative to that in the non–IFN-γ–treated control by luciferase analysis at 24 h postinfection. The graphs show the mean ± SEM in four independent experiments. Two-tailed t tests were used: **P < 0.001 versus WT, Irgb10 KO MEFs or Irgm1/m3 DKO MEFs.

(E, F) Electron microscope images of T. gondii–infected WT (E) and Irgb6 KO (F) MEFs at 4 h postinfection in the presence of IFN-γ stimulation. The images are representative of three independent experiments. Red arrowheads indicate membrane blebbing. Scale bars, 1 μm.
As shown in Fig 2, Irgb6 is required for normal levels of ubiquitin and other effector IRG protein binding to the *T. gondii* PVM. However, Irga6 deficiency (Fig 3A and B) and Irgb10 deficiency (Fig 3C and D) did not affect Irgb6 and ubiquitin localization on the PVM compared with wild-type MEFs. This suggests that Irgb6 and ubiquitin initially have the ability to bind to...
Figure 3. Regulator IRG proteins, but not other effector IRG proteins, are required for loading of Irgb6 and ubiquitin on *T. gondii* PVM.

(A, B) Confocal microscope images (left) and the graphs (right) represent the localization of Irgb6 (A) and ubiquitin (B) (red) to *T. gondii* vacuoles (green), and DAPI (blue) at 4 h postinfection in IFN-γ-treated WT and Irga6 KO MEFs. (C, D) Confocal microscope images (left) and the graphs (right) represent the localization of Irgb6 (C) and ubiquitin (D) (red) to *T. gondii* vacuoles (green), and DAPI (blue) at 4 h postinfection in IFN-γ-treated WT and Irgb10 KO MEFs. (E, F) Confocal microscope images (left) and the graphs (right) represent the localization of Irgb6 (E) and ubiquitin (F) (red) to *T. gondii* vacuoles (green), and DAPI (blue) at 4 h postinfection in IFN-γ-treated WT and Irgm1/m3 DKO MEFs. All graphs show the mean ± SEM in three independent experiments. All images are representative of three independent experiments. White arrowheads indicate colocalization. Scale bars, 5 μm. ND, not detected; NS, not significant.
*T. gondii* PVM independent of Irga6 and Irgb10. Furthermore, we investigated whether Irgb10 was capable of compensating for Irgb6 deficiency. Immunofluorescence staining identified no significant difference in Irgb6, Gbp1, Gbp2, Gbp1-5, ubiquitin, and p62 recruitment on *T. gondii* PVM when observing Irgb6 single- and Irgb6/Irgb10 double-deficient MEFs (Fig S3A). This implies that only Irgb6 is required for the recruitment of other effector IRG proteins, GBP, and ubiquitin.

**Regulator IRG proteins are required for Irgb6 and ubiquitin recruitment on *T. gondii* vacuoles**

During *T. gondii* infection, regulator IRG proteins such as Irgm1 and Irgm3 have been shown to be involved in proper localization of active form, GTP-bound effector IRG proteins on the PVM. These proteins distinguish the membranes of cellular organelles from "non-self" vacuoles (Coers et al, 2008; Hunn et al, 2008; Haldar et al, 2013). Recruitment of Irgb10 and Gbp2 on the *T. gondii* PVM is completely unsuccessful in Irgm1/m3 double-deficient MEFs (Haldar et al, 2013). Our previous findings also showed that IFN-γ-induced ubiquitin and p62 localization on the *T. gondii* PVM is not detectable in Irgm1/m3 double-deficient MEFs (Lee et al, 2015). Thus, we examined the contribution of regulator IRG proteins to controlling Irgb6 localization on the *T. gondii* PVM using Irgm1/m3 double-deficient MEFs. A previous study has reported that regulator IRG proteins are required for the localization of Irgb6 on the PVM in a different system (Hunn et al, 2008). As expected, IFN-γ-induced localization of Irgb6 and ubiquitin on *T. gondii* PVM was completely unsuccessful in Irgm1/m3 double-deficient MEFs (Fig 3E and F).

**GTPase activity is required for Irgb6 localization at *T. gondii* PVM**

Irgb6 binds to PISP and phosphatidylycerine (PS), which are both detected at *T. gondii* PVM

The IRG Irgb6 belongs to the dynamin GTPase superfamily. These GTPases play an important role in the fission of clathrin–coated pits from the plasma membrane during endocytosis (Martens & Howard, 2006; Antonny et al, 2016). It has been determined that dynamin binding to acidic phospholipids is essential for dynamin-mediated membrane recognition (Salim et al, 1996; Lemmon & Ferguson, 2000). We hypothesized that acidic phospholipid binding would play a role in initial Irgb6-mediated *T. gondii* PVM recognition. We, therefore, performed a protein–lipid overlay assay using a Histagged recombinant Irgb6 protein (Fig 4A), which revealed that Irgb6 mainly bound to monophosphorylated phosphoinositides (Pis) such as P13P, PI4P, PISP, or PS. Among these, the strongest interaction was observed with PISP (Fig 4A). We next tested whether *T. gondii* PVM is composed of these phospholipids. Indeed, immunofluorescence staining revealed that anti-PS or anti-P1P2, which recognize P13P, PI4P, PISP, P(3,5)P3, P(3,4,5)P3, and PA (Fig S5A), stained *T. gondii* PVM (Fig 4B and C). This implies that PS or P1s may localize at *T. gondii* PVM. To visualize cellular P13P, PI4P, PISP, P(3,5)P3, and PS, we used HA-tagged p40-PX, OSBP, ING2-PHD, Btk-PH, and the MFG-E8–C2 domain, as previously described (Kanai et al, 2001; Hanayama et al, 2002; Levine & Munro, 2002; Gozani et al, 2003; Ebner et al, 2017). Notably, HA-tagged ING2-PHD and the MFG–E8–C2 domain were detected on the PVM, whereas HA-tagged p40-PX, OSBP, and Btk–PH were not (Fig 4D). Quantification analysis of immunofluorescence staining showed that about 34.0 or 9.7% of *T. gondii* vacuoles are HA-tagged ING2-PHD–positive or HA-tagged MFG–E8–C2 domain–positive, respectively. However, HA-tagged p40-PX, OSBP, and Btk–PH were detected much less frequently on the PVM (Fig 4E), suggesting that PISP and PS can be recognized by Irgb6 at the PVM.

**Basic amino acid residues in C-terminal α-helices are important for both Irgb6 loading onto *T. gondii* PVM and phospholipid binding**

We next sought to identify the Irgb6 regions involved in phospholipid–binding–dependent *T. gondii* PVM recognition. The C-terminal α–K helix of regulatory IRG proteins seems to be crucial for the specificity of intracellular organelle targeting (Martens et al, 2004; Martens & Howard, 2006; Henry et al, 2014) or for targeting to the mycobacterial phagosome (Tiwari et al, 2009). A recent study showed that the C-terminal α–helical domains (especially αF and αK) of Irgb10 were predicted to be required for Irgb10 antimicrobial action involving intracellular bacterial membrane targeting (Man et al, 2016). Irgb6 is predicted to possess two C-terminal α–helical domains (αF and αK) based on the crystal structure analysis of Irga6, which provides the general structure for mouse IRG proteins (Fig 5A) (Ghosh et al, 2004). We, therefore, focused on two C-terminal α–helical domains (αF and αK) in Irgb6. When we deleted the αF and/or αK domain from WT Irgb6 and expressed them in Irgb6-deficient MEFs (Fig 5B and C), the mutants lacking α–helices were not loaded onto the PVM (Fig 5B and C). This suggests that the α–helical regions are important for Irgb6 targeting to *T. gondii* PVM. Given that both of the α–helices are amphipathic (Fig 5D) (Man et al,
Figure 4. Irgb6 recognizes PI5P and PS at T. gondii PVM.

(A) The representative image from three independent experiments of PIP strips showing the binding of His-tagged recombinant Irgb6 protein to PI3P, PI4P, PI5P, and PS.

(B) Confocal microscope images from two independent experiments of the localization of PS (green) with T. gondii (red), and DAPI (blue) in WT MEFs. Scale bars, 5 μm.

(C) Confocal microscope images from three independent experiments of the localization of PIs (green) with T. gondii (red), and DAPI (blue) in WT MEFs. Scale bars, 5 μm.

(D, E) Confocal microscope images (D) and the graphs (E) from three independent experiments represent the localization of the indicated HA-tagged probes recognizing each specific PIs (green) with T. gondii (red), and DAPI (blue) in WT MEFs. Scale bars, 5 μm. White arrowheads indicate colocalization.
Figure 5. Basic amino acids in the C-terminal α-helices of Irgb6 are required for recognition of phospholipids and immune responses against T. gondii.

(A) Schematic representation of WT Irgb6 and C-terminal α-helices-deleted mutants of Irgb6. (B, C) The ratio (B) and representative images (C) of colocalization of FLAG-tagged WT Irgb6 or FLAG-tagged C-terminal deletion mutants Irgb6 (red) to T. gondii vacuoles (green) by confocal microscopy analysis at 4 h postinfection in Irgb6 KO MEFs reconstituted with the indicated proteins in the presence of IFN-γ stimulation. Scale bars, 5 μm.

(D) The helical wheel projection of the two α-helical regions of Irgb6 and alignment according to its amphiphilic properties. Yellow indicates nonpolar amino acids, purple indicates acidic amino acids, blue indicates polar amino acids, and green indicates basic amino acids.

(E, F) The ratio (E) and representative images (F) of colocalization of FLAG-tagged WT Irgb6 or FLAG-tagged the point mutants Irgb6 (red) to T. gondii vacuoles (green) by confocal microscopy analysis at 4 h postinfection in Irgb6 KO MEFs reconstituted with the indicated proteins in the presence of IFN-γ stimulation. Scale bars, 5 μm.

(H) Western blotting analysis of His-tagged WT Irgb6 and K275A/R371A Irgb6.
and that PI5P and PS are negatively charged at physiological pH, we hypothesized that basic amino acid residues such as K275 and R371 (both of which are positively charged at physiological pH) are involved in phospholipid-mediated T. gondii PVM targeting. To test this, we generated Irgb6 mutants in which various basic amino acid residues around the α-helical regions were replaced with alanine. We assessed whether the Irgb6 mutants were recruited to the PVM via ectopic expression in IFN-γ-stimulated Irgb6-deficient MEFs (Figs 5E and F, and S5B). We found that K275A and R371A accumulation on T. gondii PVM was severely impaired (Fig 5E). This did not occur for the K233A, K266A, K268A, or K395A Irgb6 mutants (Fig S5B), suggesting that K275 and R371 residues are specifically involved in Irgb6 loading on the PVM. Furthermore, recruitment was completely abolished in the Irgb6 mutant in which both K275 and R371 residues were replaced with alanine residues (K275A/R371A) (Fig 5E and F).

Next, we performed a protein–lipid overlay assay using the K275/ R371 Irgb6 mutant protein. We found that the mutant was not able to bind to phospholipids compared with WT Irgb6 (Fig 5G). The signals were detected simultaneously to ensure equal exposure times. The efficiency of recombinant protein purification was probed by Coomassie blue staining, and both proteins were detected as a single bright band in each well of the gel corresponding to the expected product sizes (Fig 5H). Ectopic K275A/R371A expression in the Irgb6-deficient MEFs consistently failed to rescue IFN-γ–induced parasite killing activity (Fig 5I).

K275 and R371 residues were mapped onto the in silico Irgb6 structural model generated using the crystallized Igr6 structure (Ghosh et al, 2004). These basic amino acids were predicted to be on the very edge of the α-helices, located on the opposite side of the GTPase domain and facing outside of the possible membrane-targeting region (Fig 5I). Collectively, these results show that basic amino acid residues in the C-terminal α-helices of Irgb6 are important for initial targeting of phospholipid-mediated Irgb6 to T. gondii PVM and cell-autonomous immunity.

Irgb6 provides in vivo host protection from T. gondii infection

The physiological relevance of Irgb6 in the host defense response against pathogens is unknown. We, therefore, investigated the role of Irgb6 in the host defense against T. gondii infection in vivo. To do this, we infected WT and Irgb6-deficient mice with luciferase-expressing type II Pru T. gondii and monitored their infection susceptibility profiles. Although the luciferase signals were comparably emitted from both WT and Irgb6-deficient mice on day 3 postinfection, the signals from the Irgb6-deficient mice on days 5 and 6 postinfection were greatly enhanced in comparison with those from the WT mice (Fig 6A and B). On day 5 postinfection, parasite numbers were counted in the peritoneal fluids or tissues collected from the infected animals. It was found that Irgb6-deficient mice had higher parasite numbers than WT mice (Fig 6C). Moreover, the proinflammatory cytokine (e.g., IL-6, IL-12, and IFN-γ) levels in the peritoneal fluids from the T. gondii–infected Irgb6-deficient mice were enhanced compared with those in WT mice (Fig 6D). Furthermore, all Irgb6-deficient mice infected with T. gondii died within 9 d of infection, whereas all WT mice survived (Fig 6E). Thus, these results show that robustly increased parasite burden in Irgb6-deficient mice causes much higher proinflammatory responses and enhanced host susceptibility. This suggests that Irgb6 has a crucial role in initial binding on the PVM during the host defense response against T. gondii infection.

Discussion

Soon after T. gondii invades IFN-γ–stimulated mouse cells, the parasite PVM becomes coated with multiple effector IRG proteins and is eventually disrupted (Martens et al, 2005). The loading process of effector IRG proteins is highly ordered, and Irgb6 and Irgb10 are reported to be the first in this process (Khaminets et al, 2010).

Here, we showed that Irgb6, but not Irgb10, plays a major role in anti-T. gondii cell-autonomous immunity. Irgb6-deficient cells were severely defective in the recruitment of ubiquitin and other IFN-γ–inducible GTPase effectors onto the T. gondii PVM. However, Irga6 or Irgb10 deficiency did not affect Irgb6 and ubiquitin coating on the PVM, indicating that Irgb6 has a larger role among effector IRG proteins. IFN-γ–dependent ubiquitination on T. gondii PVM is well established, ultimately resulting in parasite growth restriction via several mechanisms. These include mediating parasite vacuole–lysosome fusion and recruiting autophagy adaptors in human cells (Haldar et al, 2015; Selleck et al, 2015; Clough et al, 2016). Furthermore, the binding of ubiquitin and p62 to the PVM mediates parasite antigen–specific CD8+ T cell activation in mice (Lee et al, 2015). However, the PVM substrate is not clear. Our previous study also showed that ubiquitin localization on the PVM is normal in MEFs deficient in chromosome 3 GBP family or Igr6a, excluding the possibility that these are PVM substrates for ubiquitin (Lee et al, 2015). Interestingly, we found that ubiquitin localization on T. gondii PVM is dependent on Irgb6, but not Irgb10. These data, therefore, indicate the predominant role of Irgb6 in IFN-γ–induced cell-autonomous resistance against avirulent T. gondii type II. This occurs via enhancing the targeting of ubiquitin and other effector proteins to the PVM. Future study is required to clarify whether the Irgb6 is ubiquitinated on the PVM after targeting, or whether other proteins on the PVM are ubiquitinated after Irgb6-dependent PVM disruption.
We demonstrated that Irgb6 binding to the PVM is completely dependent on regulator IRG proteins such as Irgm1 and Irgm3. Furthermore, the universally conserved phosphate-binding loop GKS motif in the Irgb6 GTPase domain is essential for targeting to T. gondii vacuoles. Reconstituting Irgb6 S70N and K69A mutants in Irgb6-deficient cells revealed the essential role of GTPase activity when Irgb6 accumulates on T. gondii PVM, as shown by a previous study (Hunn et al, 2008).

We have also shown that the phospholipid binding activity of Irgb6 is required for PVM targeting. Recombinant Irgb6 protein was able to bind PS and PIs such as PI3P, PI4P, and PI5P in vitro. Cellular PI5P and PS also appeared to specifically accumulate at the PVM via unknown mechanisms. We found that K275 and R371 in the C-terminal α-helical domains were the critical amino acid residues in Irgb6 for phospholipid binding and T. gondii PVM loading. Interactions between amphipathic α-helices of antimicrobial peptides and lipids are involved in targeting antimicrobial peptides to microbial membranes composed of negatively charged lipids (Dathe & Wieprecht, 1999; Shai, 1999; Mihajlovic & Lazaridis, 2010; Zhang & Gallo, 2016). Considering this, Irgb6 likely accumulates on the T. gondii PVM via an electrostatic interaction between K275/R371 (basic or positively charged at physiological pH) and PI5P/PS (acidic or negatively charged). Among the PIs, the role of PI5P in mammalian cells remains poorly understood (De Craene et al, 2017). Cellular PI5P levels are extremely low in many cell types relative to some other PIs (1–2% of PI4P) (Grainger et al, 2012; Viaud et al, 2014). In host cells, PI5P...
can be altered by Shigella flexneri virulence factor IpgD. This converts P(4,5)P2 to PIP3, leading to direct actin remodeling and modification of the microtubule machinery (Niebuhr et al, 2002; Ramel et al, 2011; Boal et al, 2016; Hasegawa et al, 2017). Despite its presence in low levels, PIP3 seems to be present in the plasma membrane (Sarkes & Rameh, 2010) and is phosphorylated by type II PIP4K to become P(4,5)P3 (Rameh et al, 1997). This is hydrolyzed into the Ca2+-mobilizing messenger inositol trisphosphate and the protein kinase C activator diacylglycerol at the plasma membrane (Berridge & Irvine, 1984). PIP3 also seems to be present in intracellular membrane compartments, including endosomes, the ER, and the Golgi apparatus (Sarkes & Rameh, 2010). Phosphatidylinosine is synthesized in the ER and delivered to the plasma membrane by vesicular trafficking via the Golgi apparatus (Fairn et al, 2011). Although Irgb6 possibly recognizes these endomembranes, Irgm1 and Irgm3 may keep Irgb6 in a GDP-bound state, and hence an inactive form, to protect these organelles. Cytoplasmic endomembranous organelles such as the Golgi apparatus, mitochondria, ER, and endolysosomes are coated with regulator IRG proteins such as Irgm1 and Irgm3 (Hunn et al, 2008; Springer et al, 2013). Nevertheless, the protective mechanism preventing Irgb6 loading at the plasma membrane remains unclear. When T. gondii actively invades host cells, the moving junction, which forms at the host cell entry site, serves as a molecular sieve for host proteins. These protect the parasite PVM from lysosomal degradation (Mordue et al, 1999). The molecular sieve formed by moving junction formation during T. gondii invasion may alter PIP3 levels at the host plasma membrane, possibly increasing PIP3 levels on the PVM and triggering Irgb6 loading. In terms of phospholipids and the IRG protein, Irgm1 was previously shown to interact with P(3,4)P2 and P(3,4,5)P3 and is important for localization to bacteria-containing phagosomes (Tiwari et al, 2009). However, whether Irgm1 binding to phospholipids is important for IFN-γ–mediated immunity to T. gondii remains unclear. Whether Irgm1 localizes to bacteria-containing phagosomes is currently disputed (Springer et al, 2013).

Finally, our findings demonstrated the protective role of Irgb6 in vivo against T. gondii infection. Irgb6 deficiency causes markedly increased parasite burden in the tissues and increased inflammatory responses. This suggests that upon Irgb6-deficiency, failure of PVM disruption during early T. gondii infection eventually leads to host death. During in vivo infection, T. gondii can spread to other parts of the body via interaction with immune cells, such as neutrophils and inflammatory monocytes (Lambert et al, 2006; Bierly et al, 2008; Unno et al, 2008; Dunay et al, 2010; Coombes et al, 2013). Consistently, we found that Irgb6-deficient mice displayed significantly higher numbers of T. gondii–infected inflammatory monocytes and neutrophils in the peritoneal cavity compared with wild-type mice (data not shown).

In summary, our studies discuss the fact that not much is known about the mechanisms by which IRG proteins recognize and destroy T. gondii PV and proceed to investigate this in the context of Irgb6. Our findings highlight an active host defense program relying on the initial binding of phospholipid-dependent Irgb6 to T. gondii PVM. The initial binding of Irgb6 on T. gondii vacuoles is likely to involve PIP3 and PS and is a crucial feature of ubiquitination, PVM destruction, and synergistical T. gondii killing activity. Our studies cover topics involving Irgb6 gene knockout, host–pathogen interactions, and immune responses. This study, therefore, appropriately highlights its contribution to existing research. It will be necessary to examine whether other IFN-inducible GTPases targeting bacterial and fungal membranes recognize the same or different phospholipids and induce the membrane remodeling. This could further reveal the biological significance of IFN-inducible GTPase-dependent cell-autonomous immunity.

Materials and Methods

Reagents

Antibodies against Irgb6 (TGTP; sc-11079), GBP1-5 (sc-166960), Irgm1 (LRG-47; sc-11075), Irgm2 (GTPi; sc-11088), Irgm3 (IGTP; sc-136317), and His-probe (H-3; sc-8036) were purchased from Santa Cruz Biotechnology, Inc. Antibodies against FLAG M2 (F3165) and β-actin (A1978) were purchased from Sigma-Aldrich. Mouse monoclonal and rabbit polyclonal anti-HA antibodies were obtained from BioLegend and Sigma-Aldrich, respectively. Rabbit polyclonal anti-GBP2 and mouse monoclonal anti-p62 (PM045) antibodies were obtained from Proteintech and MBL International, respectively. Anti-ubiquitin mouse monoclonal antibody (FK2; MFK-004) was obtained from Nippon Biotest Laboratories. Mouse monoclonal anti-Irga6 (10D7) and rabbit polyclonal anti-Irgb10 antibodies were provided by Dr JC Howard (Instituto Gulbenkian de Ciência). Rabbit polyclonal anti-GBP1 antibody was provided by Dr EM Frickel (Francis Crick Institute). Rabbit polyclonal anti-GRA7 antibody was provided by Dr JC Boothroyd (Stanford University School of Medicine). Mouse monoclonal anti-GRA2 and rabbit polyclonal anti-Gap43 antibodies were provided by Dr D Soldati-Favre (University of Geneva). Anti-PiP2 antibody (2C11) was purchased from Abcam. Anti-PS antibody (IH6) was purchased from Millipore. Recombinant mouse IFN-γ was purchased from PeproTech.

Mice

All animal experiments were approved by the Animal Research Committee of the Research Institute for Microbial Diseases (Osaka University, Osaka, Japan). Irgb6-deficient C57BL/6 mice and their wild-type counterparts were maintained under specific pathogen-free conditions and used for experimental study at 8–10 wks. Irgb10-deficient mice were constructed as previously described (Man et al, 2016).

Cell culture

Primary MEFs were maintained in DMEM (Nacalai Tesque) supplemented with 10% heat-inactivated FBS (JRH Bioscience), 100 U/ml penicillin (Nacalai Tesque), and 100 μg/ml streptomycin (Nacalai Tesque). Bone marrow–derived macrophages were generated by cultivating BM progenitors isolated from the BM in complete medium, containing 10% L-cell conditioned medium, for 6–7 d. The complete medium consisted of 10% heat-inactivated FBS, 10 mM Heps (Sigma-Aldrich), 100 U/ml penicillin, 100 μg/ml streptomycin, 50 μg/ml gentamicin (Sigma-Aldrich), 10 μg/ml polymyxin B (Sigma-Aldrich), 1 mM sodium pyruvate (Sigma-Aldrich), 50 μM 2-mercaptoethanol...
Generation of MEFs derived from Irgb6-deficient, Irgb10-deficient, or Irgb6/b10 double-deficient mice by Cas9/CasPR genome editing

The insert fragment of Irgb6 gRNA was amplified using KOD FX NEO (Toyobo) and the following primers: Irgb6_gRNA1_F, Irgb6_gRNA1_R, Irgb6_gRNA2_F, and Irgb6_gRNA2_R. The fragment for gRNA was inserted into the gRNA-cloning vector (Plasmid 41824) using Gibson Assembly mix (New England Biolab) to generate gRNA-expressing plasmids. The T7 promoter was added to the gRNA template using KOD FX NEO and Irgb6_T7gRNA_F and gRNA_common_R primers. The T7-IRgb6 gRNA PCR product was gel purified and used as the subsequent generation of gRNA. MEGASHortscript T7 (Life Technologies) was used for gRNA generation. Cas9 mRNA was generated by in vitro transcription using mMESSAGE mMACHINE T7 ULTRA kit (Life Technologies) was used for gRNA generation. Cas9 mRNA was generated by in vitro transcription using mMESSAGE mMACHINE T7 ULTRA kit (Life Technologies). The template was amplified by PCR using pEF6-hCas9-Puro and T7Cas9_IVT_F and Cas9_R primers (Ohshima et al., 2014), followed by gel purification. Irgb10 gRNA was previously prepared (Man et al., 2016). The synthesized gRNA and Cas9 mRNA were purified using a MEGAclear kit (Life Technologies) and eluted in RNase-free water (Nacalai Tesque). To obtain Irgb6-deficient, Irgb10-deficient, or Irgb6/b10 double-deficient MEFs, C57BL/6 female mice (6 wks old) were superovulated and mated to C57BL/6 stud males. Fertilized one-cell stage embryos were collected from oviducts and injected into the pronucleus or cytoplasm with Cas9 mRNA (100 ng/μl) and Irgb6 and/or Irgb10 gRNA (50 ng/μl). The injected live embryos were transferred into oviducts of pseudopregnant ICR females at 0.5 dpc. D13.5 embryos were collected to generate primary MEFs. The Irgb6 and Irgb10 protein expression by the resulting MEFs was analyzed by Western blot using antibodies against Irgb6 or Irgb10, respectively. Irgb6-deficient MEFs were constructed as previously described (Liesenfeld et al., 2011). The male pup harboring the mutation was mated to C57BL/6 female mice and tested for germ line transmission. Heterozygous mice were intercrossed to generate homozygous Irgb6-deficient mice for in vitro and in vivo assays. The deleted region in Irgb6 genomic DNA was verified by sequencing analysis. The complete coding region of the Irgb5-b4 tandem decoy cDNA in Irgb6-deficient MEFs was confirmed by sequencing analysis.

**Type II T. gondii parasites**

Parental PruΔHX and luciferase-expressing PruΔHX were maintained in Vero cells by passaging every 3 d in RPMI1640. This was supplemented with 2% heat-inactivated FBS, 100 U/ml penicillin, and 100 μg/ml streptomycin.

Cloning and recombinant expression

The cDNA regions of interest corresponding to the wild-type control and indicating Irgb6 point or deletion mutations (GenBank accession no. NM_001145164) were synthesized from mRNA from the spleen of C57BL/6 mice. The cDNA used to generate the HA-tagged PHD of p40, OSBP, ING2, or Btk were obtained from Addgene. The cDNA used to generate the HA-tagged MFG-E8-C2 were kindly provided by Dr S Nagata (Osaka University). The Irgb6 mutants were generated using primers (Table S1). The PCR products were then ligated into the retroviral pMRX expression or pEU-E01-MCS vector (CellFree Sciences) for retroviral infection or recombinant protein expression, respectively. The sequence of all constructs was confirmed by DNA sequencing.

Generation of recombinant Irgb6 using cell-free system

His-tagged recombinant Irgb6 WT and K275A/R371A were expressed with the wheat germ cell-free system (CellFree Sciences) and purified with a Nickel-Sepharose six fast flow (GE Healthcare), as previously reported (Tsuboi et al., 2010). Protein purity was evaluated by SDS-PAGE and Coomassie brilliant blue staining.

**Immunofluorescence study**

Cells were cultured on glass coverslips in the presence or absence of IFN-γ (10 ng/ml) for 24 h. Cells were infected with type II T. gondii (MOI of 5 or 1) for 4 h and fixed in PBS containing 3.7% paraformaldehyde for 10 min at room temperature. Cells were permeabilized with PBS containing 0.1% Triton X-100 or 0.002% digitonin for 5 min, followed by blocking with 8% FBS in PBS for 1 h at room temperature. Cells were subsequently incubated with the indicated primary antibodies for 1 h at 37°C, followed by incubation with Alexa 488,- Alexa 594-, or Alexa 647-conjugated secondary antibodies (molecular probes) and DAPI for 1 h at 37°C in the dark. Finally, coverslips were mounted onto glass slides with PermaFluor (Thermo Fisher Scientific) and analyzed using confocal laser microscopy (FV1200 IX-8; Olympus). All images are shown at a 1,000× magnification. For PIP2 and PS staining, the cells were fixed in PBS containing 3.7% paraformaldehyde and 0.2% glutaraldehyde for 3 h at 4°C. After washing with PBS containing 50 mM NH4Cl, cells were incubated in PBS containing 0.5% saponin, 8% FBS, and 50 mM NH4Cl for 3 h at 4°C. The cells were then incubated overnight with primary antibodies in PBS containing 0.1% saponin and 8% FBS at 4°C. T. gondii ΔHX and luciferase-expressing PruΔHX were assayed by DNA sequencing. The complete coding region of the Irgb6 T7gRNA_F and gRNA_common_R primers. The T7-IRgb6 gRNA PCR product was gel purified and used as the subsequent generation of gRNA. MEGASHortscript T7 (Life Technologies) was used for gRNA generation. Cas9 mRNA was generated by in vitro transcription using mMESSAGE mMACHINE T7 ULTRA kit (Life Technologies). The template was amplified by PCR using pEF6-hCas9-Puro and T7Cas9_IVT_F and Cas9_R primers (Ohshima et al., 2014), followed by gel purification. Irgb10 gRNA was previously prepared (Man et al., 2016). The synthesized gRNA and Cas9 mRNA were purified using a MEGAclear kit (Life Technologies) and eluted in RNase-free water (Nacalai Tesque). To obtain Irgb6-deficient, Irgb10-deficient, or Irgb6/b10 double-deficient MEFs, C57BL/6 female mice (6 wks old) were superovulated and mated to C57BL/6 stud males. Fertilized one-cell stage embryos were collected from oviducts and injected into the pronucleus or cytoplasm with Cas9 mRNA (100 ng/μl) and Irgb6 and/or Irgb10 gRNA (50 ng/μl). The injected live embryos were transferred into oviducts of pseudopregnant ICR females at 0.5 dpc. D13.5 embryos were collected to generate primary MEFs. The Irgb6 and Irgb10 protein expression by the resulting MEFs was analyzed by Western blot using antibodies against Irgb6 or Irgb10, respectively. Irgb6-deficient MEFs were constructed as previously described (Liesenfeld et al., 2011). The male pup harboring the mutation was mated to C57BL/6 female mice and tested for germ line transmission. Heterozygous mice were intercrossed to generate homozygous Irgb6-deficient mice for in vitro and in vivo assays. The deleted region in Irgb6 genomic DNA was verified by sequencing analysis. The complete coding region of the Irgb5-b4 tandem decoy cDNA in Irgb6-deficient MEFs was confirmed by sequencing analysis.

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Transmission electron microscopy

Primary MEFs treated with 10 ng/ml IFN-γ for 24 h were infected with type II T. gondii (MOI of 5) for 4 h. After washing with PBS, the cells were fixed overnight with 25% glutaraldehyde in 0.1 M phosphate buffer at 4°C. The cells were post-fixed with 1% OsO4 in the same buffer for 1 h at 4°C, dehydrated in a graded series of ethanol, and embedded in Quetol 812 (Nissin EM). Silver sections were cut with an ultramicrotome, stained with lead citrate and uranyl acetate, and observed with an H-7650 electron microscope (Hitachi).

Western blot analysis

Cells were washed with PBS and lysed with 1× TNE buffer (20 mM Tris–HCl, 150 mM NaCl, 1 mM EDTA, and 1% NP-40) containing protease inhibitor cocktail (Nacalai Tesque). A total protein was
loaded onto and separated on a 10% or 15% SDS–PAGE and transferred to a polyvinyl difluoride membrane. The membrane was blocked with 5% skim milk in PBS/Tween 20 (0.2%) and probed overnight with the indicated primary antibodies at 4°C. After washing with PBS/Tween, the blot was probed with HRP-conjugated secondary antibodies for 1 h at 25°C and visualized by Luminata Forte Western HRP substrate (Millipore).

**Protein–lipid overlay assay**

A protein–lipid overlay assay was performed with 0.5 μg/ml of His-tagged recombinant proteins using PIP strips (Echelon Biosciences), as per the manufacturer’s instructions. Lipid binding was immunodetected with a combination of anti-His tag mouse monoclonal antibody and HRP-conjugated rabbit-anti-mouse IgG, followed by Luminata Forte Western HRP substrate. In detail, the PIP strip membrane (P-6001; Echelon Biosciences Inc.) was blocked with TBS-T (0.1% Triton X-100) + 3% BSA (fatty acid free) for 1 h at 25°C with gentle agitation. Next, we discarded blocking buffer and added 0.5 μg/ml purified recombinant protein Irgb6-His tagged in 1.5 ml TBS-T + 3% BSA and incubated overnight at 4°C. Next, we discarded the protein solution and washed with TBS-T five times with gentle agitation for 5 min each. Next, we discarded wash buffer and added 1.5 μl anti–His–HRP antibody in 1.5 ml TBS-T + 3% BSA and incubated the membrane for 1 h at 25°C, with gentle agitation. Next, we discarded the antibody solution and washed with TBS-T six times with gentle agitation for 10 min each. Next, we discarded wash buffer and detected the bound protein by ECL development followed by ImageQuant LAS 4000 (GE Healthcare). The displayed images in Figs 4 and 5 were used from ImageQuant LAS 4000.

**Mice survival and in vivo measurement of parasites by imaging**

Mice were intraperitoneally infected with PruΔHX T. gondii tachyzoites expressing luciferase (1 × 10⁶ in 200 μl PBS per mouse), and the survival of the mice was monitored for 15 d postinfection. For in vivo imaging of the parasites, mice were intraperitoneally injected with 3 mg D-luciferin in 200 μl PBS (Promega) on days 3, 5, and 6 postinfection. After inhalational anesthesia with isoflurane, abdominal photon emission was assessed during 60-s exposure by an in vivo imaging system (IVIS-Spectrum; Xenogen), followed by analysis with Living Image software (Xenogen).

**Luciferase assay**

To measure the number of T. gondii, cells were left untreated or treated with IFN-γ (10 ng/ml) for 24 h. Following this, cells were infected with luciferase-expressing PruΔHX T. gondii (MOI of 1) for 24 h. The infected cells were harvested and lysed with 100 μl 1× passive lysis buffer (Promega) with sonication for 30 s. To measure the number of T. gondii in the tissues, peritoneal cavity, mesenteric lymph nodes, spleen, liver, and lungs, these sections were removed on day 5 postinfection. The mesenteric lymph nodes, spleen, liver, and lungs were homogenized and lysed in 1 ml 1× passive lysis buffer with sonication. After centrifugation at 13,000g for 10 min at 4°C, luciferase activity was measured using 5 μl of the supernatants via the dual-luciferase reporter assay system (Promega) by GLOMAX 20/20 luminometer (Promega). The in vitro data are presented as the percentage of T. gondii survival in IFN-γ–stimulated cells relative to unstimulated cells (Con). The in vivo data are presented as absolute values.

**ELISA**

The level of mouse IL-6, IL-12, and IFN-γ secretion was measured by ELISA analysis according to the manufacturer’s protocol (eBioscience).

**In silico structural modeling**

For structural modeling of Irgb6, the Irga6 structure (Protein Data Bank ID 1TQ4) was used as a template with HHpred alignments and the spanner three-dimensional rendering tool. Figures were prepared using the PyMOL molecular graphics system (version 1.7.6.0, Schrödinger, LLC).

**Statistical analysis**

Statistical analyses were performed using the t test. P-values less than 0.05 were considered statistically significant. All graphs show the mean ± SEM of three independent experiments (three biological replicates).

**Supplementary Information**

Supplementary Information is available at https://doi.org/10.26508/lsa.201900549.

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**Author Contributions**

Y Lee: conceptualization, resources, data curation, software, formal analysis, funding acquisition, validation, investigation, visualization, and writing—original draft, review, and editing.
H Yamada: data curation, visualization, and methodology.
JS Ma: data curation and formal analysis.
M Okamoto: investigation.
H Nagaoka: resources and investigation.
E Takashima: resources and investigation.
DM Standley: software, investigation, and visualization.
E Takashima: resources and investigation.
M Yamamoto: conceptualization, resources, data curation, formal analysis, supervision, funding acquisition, validation, investigation, visualization, methodology, project administration, and writing—original draft, review, and editing.

Conflict of Interest Statement

The authors declare that they have no conflict of interest.

References

Al-Zeer MA, Al-Younes HM, Braun PR, Zerrahn J, Meyer TF (2009) IFN-gamma-inducible Irgα6 mediates host resistance against Chlamydia trachomatis via autophagy. PLoS One 4: e4588. doi:10.1371/journal.pone.0004588
Antony B, Burd C, De Camilli P, Chen E, Daumke O, Efaelker K, Ford M, Frolov VA, Frost A, Hinshaw JE, et al (2016) Membrane fission by dynamin: What we know and what we need to know. EMBO J 35: 2270–2284. doi:10.15252/embj.201694613
Bekpen C, Hunn JP, Rohde C, Parvanova I, Guehthlein L, Dunn DM, Gassella E, Leptin M, Howard JC (2005) The interferon-inducible p47 (IRG) GTPases in vertebrates: Loss of the cell autonomous resistance mechanism in the human lineage. Genome Biol 6: R92. doi:10.1186/gb-2005-6-11-r92
Berridge MJ, Irvine RF (1984) Inositol trisphosphate, a novel second messenger in cellular signal transduction. Nature 312: 315–321. doi:10.1038/s312315a0
Bierly AL, Shufesky WJ, Sukhumavasi W, Morelli AE, Denkers EY (2008) Dendritic cells expressing plasmacoid marker PDCA-1 are Trojan horses during Toxoplasma gondii infection. J Immunol 181: 8485–8491. doi:10.4049/jimmunol.181.12.8485
Boal F, Puhar A, Xuereb JM, Kudzuova O, Sansonetti PJ, Payrastre B, Tronchere H (2016) PISP triggers ICAM-1 degradation in Shigella-infected cells, thus dampening immune cell recruitment. Cell Rep 14: 750–759. doi:10.1016/j.celrep.2015.12.079
Clough B, Wright JD, Pereira PM, Hirst EM, Johnston AC, Henriques R, Frickel EM, Fairn GD, Schieber NL, Ariotti N, Murphy S, Kuerschner L, Webb RI, Grinstein S, Parton RG (2011) High-resolution mapping reveals topologically distinct cellular pools of phosphatidylserine. J Cell Biol 194: 257–275. doi:10.1083/jcb.201012028
Ferreira-da-Silva Mda F, Springer-Frauenhoff HM, Bohne W, Howard JC (2014) Identification of the microsporidian Encephalitozoon cuniculi as a new target of the IFNgamma-inducible IRG resistance system. PLoS Pathog 10: e1004449. doi:10.1371/journal.ppat.1004449
Gazzinelli RT, Hakim FT, Hiemy S, Shearer GM, Sher A (1991) Synergistic role of CD4+ and CD8+ T lymphocytes in IFN-gamma production and protective immunity induced by an attenuated Toxoplasma gondii vaccine. J Immunol 146: 286–292.
Ghosh A, Utahaia R, Howard J, Herrmann C, Wolf E (2004) Crystal structure of IIGPT: A paradigm for interferon-inducible p47 resistance GTPases. Mol Cell 15: 727–739. doi:10.1016/j.molcel.2004.07.017
Gozani O, Karupyan P, Jones DR, Ivanov D, Cha J, Lugooyvskoy AA, Baird CL, Zhu H, Field SJ, Lessnick SL, et al. (2003) The PHD finger of the chromatin-associated protein ING2 functions as a nuclear phosphoinositide receptor. Cell 114: 99–111. doi:10.1016/s0092-8674(03)00460-x
Greiner DL, Tavelis C, Ryan AJ, Hinchcliffe KA (2012) The emerging role of PtdIns5P: Another signalling phosphoinositide takes its place. Biochem Soc Trans 40: 257–261. doi:10.1042/bst20110671
Haldar AK, Foltz C, Finethy R, Piro AS, Feeley EM, Pilla-Moffett DM, Komatsu M, Frickel EM, Coers J (2015) Ubiquitin systems mark pathogen-containing vacuoles as targets for host defense by guanylate binding proteins. Proc Natl Acad Sci U S A 112: E5628–E5637. doi:10.1073/pnas.1519661112
Haldar AK, Saka HA, Piro AS, Dunn JD, Henry SC, Dunn JD, Henry SC, Taylor GA, Frickel EM, Valdivia RH, Coers J (2013) IRG and GBP host resistance factors target aberrant, “non-self” vacuoles characterized by the missing of “self” IRGM proteins. PLoS Pathog 9: e1003414. doi:10.1371/journal.ppat.1003414
Hanayama R, Tanaka M, Miwa K, Shinozha A, Iwamatsu A, Nagata S (2002) Identification of a factor that links apoptotic cells to phagocytes. Nature 417: 182–187. doi:10.1038/417182a
Hasegawa J, Tanaka M, Miwa K, Shinozha A, Iwamatsu A, Nagata S (2002) Identification of a factor that links apoptotic cells to phagocytes. Nature 417: 182–187. doi:10.1038/417182a
Hasegawa J, Strunk BS, Weisman LS (2017) PI5P and PI(3,5)P2: Minor, but essential phosphoinositides. Cell Struct Funct 42: 49–60. doi:10.1074/csf.2017003
Henry SC, Schmidt EA, Fessler MB, Taylor GA (2014a) Palmitoylation of the immunity related GTPase, Irgm1: Impact on membrane localization and ability to promote mitochondrial fission. PLoS One 9: e95021. doi:10.1371/journal.pone.0095021
Dathe M, Wieprecht T (1999) Structural features of helical antimicrobial peptides: Their potential to modulate activity on model membranes and biological cells. Biochim Biophys Acta 1462: 71–87. doi:10.1016/s0005-2736(99)00201-1
De Craene JO, Bertazzi DL, Bar S, Friant S (2017) Phosphoinositides, major actors in membrane trafficking and lipid signaling pathways. Int J Mol Sci 18: E634. doi:10.3390/ijms1803634
Degrandi D, Konermann C, Beuter–Gunia C, Kresse A, Wurthner J, Kurig S, Beer S, Pfeffer K (2007) Extensive characterization of IFN-induced GTPases mGBP1 to mGBP10 involved in host defense. J Immunol 179: 7729–7740. doi:10.4049/jimmunol.179.11.7729
Degrandi D, Kravets E, Konermann C, Beuter-Gunia C, Klumpers V, Lahme S, Wischmann E, Mausberg AK, Beer-Hammer S, Pfeffer K (2013) Murine guanylate binding protein 2 (mGBP2) controls Toxoplasma gondii replication. Proc Natl Acad Sci U S A 110: 294–299. doi:10.1073/pnas.1205635110
Dunay IR, Fuchs A, Sibley LD (2010) Inflammatory monocytes but not neutrophils are necessary to control Toxoplasma gondii in mice. Infect Immun 78: 1564–1570. doi:10.1128/iai.00472-09
Eber M, Lucic I, Leonard TA, Yudushkin I (2017) PI(3,4,5)P3 engagement restricts aCt activity to cellular membranes. Mol Cell 65: 416–431.e6. doi:10.1016/j.molcel.2016.12.028
Fairn GD, Schieber NL, Ariotti N, Murphy S, Kuerschner L, Webb RI, Grinstein S, Parton RG (2011) High-resolution mapping reveals topologically distinct cellular pools of phosphatidylserine. J Cell Biol 194: 257–275. doi:10.1083/jcb.2011021028
Grainger DL, Tavelis C, Ryan AJ, Hinchcliffe KA (2012) The emerging role of PtdIns5P: Another signalling phosphoinositide takes its place. Biochem Soc Trans 40: 257–261. doi:10.1042/bst20110671
Hanayama R, Tanaka M, Miwa K, Shinozha A, Iwamatsu A, Nagata S (2002) Identification of a factor that links apoptotic cells to phagocytes. Nature 417: 182–187. doi:10.1038/417182a
Hasegawa J, Strunk BS, Weisman LS (2017) PISP and PI(3,5)P2: Minor, but essential phosphoinositides. Cell Struct Funct 42: 49–60. doi:10.1074/csf.2017003
Henry SC, Schmidt EA, Fessler MB, Taylor GA (2014a) Palmitoylation of the immunity related GTPase, Irgm1: Impact on membrane localization and ability to promote mitochondrial fission. PLoS One 9: e95021. doi:10.1371/journal.pone.0095021
Irgb6 roles in anti-T. gondii responses  Lee et al.
Irgb6 roles in anti-T. gondii responses

Lee et al.
Toxoplasma gondii growth in a strain-specific manner in IFN-gamma-activated human cells. MBio 6: e01157–e01115. doi:10.1128/mbio.01157-15

Shai Y (1999) Mechanism of the binding, insertion and destabilization of phospholipid bilayer membranes by alpha-helical antimicrobial and cell non-selective membrane-lytic peptides. Biochim Biophys Acta 1462: 55–70. doi:10.1016/s0005-2736(99)00200-x

Springer HM, Schramm M, Taylor GA, Howard JC (2013) Irgm1 (LRG-47), a regulator of cell-autonomous immunity, does not localize to mycobacterial or listerial phagosomes in IFN-gamma-induced mouse cells. J Immunol 191: 1765–1774. doi:10.4049/jimmunol.1300641

Suzuki Y, Orellana MA, Schreiber RD, Remington JS (1988) Interferon-gamma: The major mediator of resistance against Toxoplasma gondii. Science 240: 516–518. doi:10.1126/science.3128869

Taylor GA, Collazo CM, Yap GS, Nguyen K, Gregorio TA, Taylor LS, Eagleson B, Secrest L, Southon EA, Reid SW, et al (2000) Pathogen-specific loss of host resistance in mice lacking the IFN-gamma-inducible gene IGTP. Proc Natl Acad Sci U S A 97: 751–755. doi:10.1073/pnas.97.2.751

Taylor GA, Feng CG, Sher A (2007) Control of IFN-gamma-mediated host resistance to intracellular pathogens by immunity-related GTPases (p47 GTPases). Microbes Infect 9: 1644–1651. doi:10.1016/j.micinf.2007.09.004

Taylor GA, Jeffers M, Largaespada DA, Jenkins NA, Copeland NG, Vande Woude GF (1996) Identification of a novel GTPase, the inducibly expressed GTPase, that accumulates in response to interferon gamma. J Biol Chem 271: 20399–20405. doi:10.1074/jbc.271.34.20399

Tiwari S, Choi HP, Matsuzawa T, Pyapaert M, MacMicking JD (2009) Targeting of the GTPase Irgm1 to the phagosomal membrane via PtdIns(3,4)P(2) and PtdIns(3,4,5)P(3) promotes immunity to mycobacteria. Nat Immunol 10: 907–917. doi:10.1038/ni.1759

Tsuboi T, Takeo S, Sawasaki T, Torii M, Endo Y (2010) An efficient approach to the production of vaccines against the malaria parasite. Methods Mol Biol 607: 73–83. doi:10.1007/978-1-60327-331-2_8

Unno A, Suzuki K, Xuan X, Nishikawa Y, Kito K, Takashima Y (2008) Dissemination of extracellular and intracellular Toxoplasma gondii tachyzoites in the blood flow. Parasitol Int 57: 515–518. doi:10.1016/j.parint.2008.06.004

Uthaiah RC, Praefcke GJ, Howard JC, Herrmann C (2003) IIGP1, an interferon-gamma-inducible 47-kDa GTPase of the mouse, showing cooperative enzymatic activity and GTP-dependent multimerization. J Biol Chem 278: 29336–29343. doi:10.1074/jbc.m211973200

Vlaud J, Boal F, Tronchere H, Gaiti-lacovoni F, Payrastre B (2014) Phosphatidylinositol 5-phosphate: A nuclear stress lipid and a tuner of membranes and cytoskeleton dynamics. Bioessays 36: 260–272. doi:10.1002/bies.201300132

Virreira Winter S, Niedelman W, Jensen KD, Rosowski EE, Julien L, Spooner E, Caradonna K, Burleigh BA, Saeij JP, Ploegh HL, et al (2011) Determinants of GBP recruitment to Toxoplasma gondii vacuoles and the parasitic factors that control it. PLoS One 6: e24434. doi:10.1371/journal.pone.0024434

Yamamoto M, Okuyama M, Ma JS, Kimura T, Kamiyama N, Saiga H, Ohshima J, Sasai M, Kayama H, Okamoto T, et al (2012) A cluster of interferon-gamma-inducible p65 GTPases plays a critical role in host defense against Toxoplasma gondii. Immunity 37: 302–313. doi:10.1016/j.immuni.2012.06.009

Yarovinsky F, Sher A (2006) Toll-like receptor recognition of Toxoplasma gondii. Int J Parasitol 36: 255–259. doi:10.1016/j.ijpara.2005.12.003

Zhang LJ, Gallo RL (2016) Antimicrobial peptides. Curr Biol 26: R14–R19. doi:10.1016/j.cub.2015.11.017

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