Host Cell Vimentin Restrains Toxoplasma gondii Invasion and Phosphorylation of Vimentin is Partially Regulated by Interaction with TgROP18

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Abstract

The obligate intracellular parasite, Toxoplasma gondii, manipulates the cytoskeleton of its host cells to facilitate infection. A significant rearrangement of host cell vimentin around Toxoplasma parasitophorous vacuoles is observed during the course of infection. ROP18 (TgROP18) is a serine-threonine kinase secreted by T. gondii rhoptry and a major virulence factor; however, the mechanisms by which this kinase modulates host factors remain poorly understood. Different and dynamic patterns of vimentin solubility, phosphorylation, and expression levels were observed in host cells infected with T. gondii strain RH and RH Δrop18 strains, suggesting that TgROP18 contributes to the regulation of these dynamic patterns. Additionally, host cell vimentin was demonstrated to interact with and be phosphorylated by TgROP18. A significant increase in T. gondii infection rate was observed in vimentin knockout human brain microvessel endothelial cells (HBMEC), while vimentin knockout or knock down in host cells had no impact on parasite proliferation and egress. These results indicate that host cell vimentin can inhibit T. gondii invasion. Interestingly, western blotting of different mouse tissues indicated that the lowest vimentin expression level was present in the brain, which may explain the mechanism underlying the nervous system tropism of T. gondii, and the phenomenon of huge cyst burdens developing in the mouse brain during chronic infection.

Key words: Toxoplasma gondii; Vimentin; TgROP18; Phosphorylation; Reorganization; Tropism.

Introduction

Toxoplasma gondii, a member of the phylum Apicomplexa, is an obligate intracellular protozoan capable of infecting a wide range of hosts, including most warm-blooded animals and humans [1, 2]. Approximately one third of the world population is infected with T. gondii [3]. The majority of infections are asymptomatic, and the parasites are under the control of host immunity; however, the infection can cause severe complications and even death in immunocompromised individuals [4]. Primary T. gondii infection in pregnant women may result in abortion/stillbirth or congenital toxoplasmosis [2].

Efficient host cell invasion and egress are critical for the survival, dissemination, and transmission of intracellular pathogens. Host cell invasion by T. gondii is a rapid process, typically completed in less than a minute [5], and is also a multistep process which includes gliding motility, host cell attachment, and active penetration [6, 7]. To date, numerous researchers have reported parasitic factors involved in cell invasion [8, 9], whereas little attention has been focused on the roles of host components in T. gondii infection. Therefore, the factors involved in T. gondii infection were explored from the perspective of the host cell in this article.

Host cell invasion by T. gondii is an active process that relies on regulated secretion of adhesive proteins and gliding motility powered by the
actin–myosin motor complex [10]. This process also relies on host cell cytoskeleton reorganization, which is regulated by T. gondii during infection [11]. Disruption of host cell actin cytoskeleton integrity decreases T. gondii cell invasion [12]. Vimentin, a type III intermediate filament cytoskeletal protein, is highly conserved in all vertebrates and is dynamically expressed and undergoes a complex phosphorylation pattern during development and differentiation [13, 14]. Vimentin has critical implications for pathophysiology; for example, it is not only an organizer of a number of critical proteins involved in cell adhesion, migration, and signaling [15, 16], it also has key roles in bacterial [17] and viral [18, 19] infections, including pathogen entry and multiplication. A significant rearrangement of host cell vimentin around the Toxoplasma parasitophorous vacuoles (PVs) occurs throughout the course of infection, and this association between host cell vimentin and PVs can be observed within an hour after T. gondii invasion [20]. Moreover, the expression of vimentin in different cells or tissues is regulated by T. gondii infection. For example, vimentin expression is up-regulated in Müller cells in a mouse model of congenital ocular toxoplasmosis [21], while it is down-regulated in Toxoplasma-infected macrophages [22].

ROP18, a serine-threonine kinase secreted by T. gondii rhoptries into the PV and host cytosol has emerged as a major virulence factor [23]; however, the mechanisms by which this kinase modulates host factors to exert its pathogenic action remain poorly understood [24]. TgROP18 can target members of the mouse IFN-γ inducible large GTPases (immunity-related GTPases [IRGs]), such as Irgb6 [25, 26], and phosphorylate human ATF6β [27], to mediate its virulence during infection. Herein, we demonstrate that expression of host cell vimentin is restrained by Toxoplasma gondii invasion, and that vimentin interacts with and is phosphorylated by TgROP18.

Results

Disruption and tagging of rop18 gene by CRISPR/CAS9-induced homologous recombination, and ectopic expression of GFP in the T. gondii RH strain

The CRISPR-Cas9 system was adopted to both disrupt rop18 gene and tag rop18 gene with eGFP-FLAG to generate a knockout mutant RH Δrop18 strain and a strain endogenously expressing GFP-FLAG-tagged Rop18 (RH-ROP18-eGFP-FLAG), respectively. To disrupt rop18, we designed a guide RNA (gRNA) to target the 5′ coding region of rop18 (sgROP18), and paired this sgRNA with a PCR amplicon consisting of the DHFR* selectable marker reading frame, flanked with regions homologous to rop18 (Figure 1A; Supplementary Material). Parasites were co-transfected with the sgROP18 CRISPR plasmid and homology template, and the recombinant tachyzoites were screened with pyrimethamine and obtained by limiting dilution. Single clones were isolated and verified by PCR to confirm the disruption of rop18 and integration of the DHFR* cassette (Figure 1B). To tag endogenous rop18 in T. gondii RH, a Cas9/CRISPR gRNA targeting the C-terminus of rop18 was generated and a homologous template was prepared, as described in methods (Figure 1C), and these were cotransfected into RH tachyzoites to generate rop18 with an eGFP-FLAG-tag at its C-terminus. Recombinant clones were confirmed by PCR (using genomic DNA as template) (Figure 1D), immunofluorescence (IF) (Figure 1E), and western blotting (WB) (Figure 1F).

Figure 1. CRISPR/Cas9-mediated gene disruption and tagging of the rop18 locus. A. Schematic of the CRISPR/CAS9 strategy used to inactivate rop18 by insertion of pyrimethamine-resistant DHFR (DHFR*). B. Verification PCR demonstrating homologous integration and gene disruption in a representative clone, compared with RH parental line tachyzoites. C. Schematic illustration of the CRISPR/CAS9 strategy used to tag endogenous ROP18 at the C-terminus with eGFP-FLAG. D. Verification PCR showing correct integration of eGFP-FLAG, SAG1 3′-UTR, and DHFR*. E. IF revealing that eGFP was successfully fused to the C-terminus of endogenous ROP18. F. Demonstration that FLAG tagged ROP18 was expressed by WB with rabbit anti-DDDDK antibody.
Additionally, to generate an RH/GFP strain ectopically expressing GFP, the recombinant plasmid, pBlue-p24-eGFP-DHFR-TS, was linearized with ApaI and then transfected into parasites. Cell lines in which the plasmid was stably integrated were selected using 3μM pyrimethamine and isolated by limiting dilution (data not shown) [28].

Dynamic patterns of vimentin were affected by T. gondii infection

To investigate the effects of T. gondii infection on vimentin reorganization, the ratios of soluble to insoluble vimentin, and total to phosphorylated vimentin were evaluated at different infection time points, based on western-blot signals. The solubility of vimentin decreased in host cells in the first 6h post-infection (Figure 3A and B), and the decrease was statistically significant (one way ANOVA; Figure 3B). Vimentin phosphorylation patterns in host cells infected with T. gondii for different amounts of time were determined by Phos-tag assays and western blotting. Vimentin phosphorylation levels were highest at 6h post-infection and subsequently decreased to, and were maintained at low levels compared with uninfected cells (Figure 3C). Vimentin total protein levels also appeared to be regulated by T. gondii infection at different time points (Figure 3D). These data suggest that the dynamic regulation of vimentin, including its solubility, phosphorylation, and expression levels, are regulated by T. gondii infection.

Vimentin inhibits T. gondii invasion into HBMEC

To explore the function of host cell vimentin during T. gondii infection, the effect of vimentin on T. gondii invasion, growth, and egress were evaluated in HBMEC and HBMEC ΔVim cells. When vimentin was knocked out, the efficiency of T. gondii invasion increased significantly (Figure 4B and C), while no significant difference was found in the number of parasites per vacuole between the two groups (Figure 4D and E). T. gondii proliferation assays were also performed in HFF cells with or without vimentin knockdown, and the results were consistent with the findings of the tests in HBMEC ΔVim and wild type cells (Figure S2). To further investigate the functions of vimentin in the lytic cycle of T. gondii, parasite egress, triggered by the Ca²⁺ ionophore A23187, was assayed. No significant difference was identified in the percentage of egressed vacuoles between HBMEC and HBMEC ΔVim, infected with RH/GFP (Figure 4F and G). Together, these results suggest that host cell vimentin inhibited the invasion of T. gondii into HBMEC, but had no significant influence on T. gondii proliferation or egress.
Figure 3. Dynamics of vimentin characteristics during infection with T. gondii RH. A. Changes in vimentin solubility in Cos7 cells infected with T. gondii for different periods of time. B. Proportion of soluble and insoluble vimentin in control and T. gondii infected cells detected by WB. The percentage of soluble vimentin was determined by analysis of grey scale intensity using Image J software (n=3, *p<0.05 by t-test). The solubility of vimentin was significantly decreased during the first 6h post T. gondii infection. C. Cos7 cells were infected with T. gondii at an MOI of 3 or uninfected (Ctrl). Phosphorylation of host cell vimentin in the different groups was analyzed by a Phos-tag assay (top panel). As vimentin expression level changed with time post T. gondii infection, the amount of cell lysate protein loaded for SDS-PAGE was adjusted to maintain a consistent level of total vimentin protein. Equal loading of vimentin was confirmed by WB of the gel (bottom panel). Bands corresponding to phosphorylated and non-phosphorylated vimentin are indicated by black and white arrowheads, respectively. Similar results were obtained in another two separate experiments. Host cell vimentin phosphorylation peaked at 6h post-infection. D. Expression levels of vimentin in Cos7 cells infected with T. gondii (as described above) were detected by WB. Dynamic expression levels were observed after infection with T. gondii for different periods of time. These results suggest that the solubility, phosphorylation, and expression levels of vimentin were regulated by T. gondii infection, and that the higher phosphorylation level of vimentin may correlate with a decrease in its solubility. Statistical differences were evaluated by t-test, means ± SD combined from three independent experiments.

Transcription and translation levels of vimentin varied among mouse tissues

To investigate the expression levels of vimentin in different mouse tissues, qRT-PCR and WB were performed to determine the levels of vimentin mRNA and protein. In our study, six different organs (brain, heart, liver, spleen, lung, and kidney), were investigated. Interestingly, we found that both mRNA and protein levels of vimentin were relatively low in brain, compared with the other mouse tissues tested (Figure 5).

Host cell vimentin interacts with TgROP18

The results of Bi-molecular fluorescence complementation (BiFC) protein interaction screening in our previous study (data not published) indicated that T. gondii ROP18 interacts with host cell vimentin. Therefore, fluorescence resonance energy transfer (FRET) was performed to confirm this interaction. The results showed a significant interaction between TgROP18 and vimentin (Figure 6A and B). Moreover, co-immunoprecipitation (Co-IP) experiments further verified the association of TgROP18 and vimentin using both α-FLAG (fused with TgROP18) and α-vimentin antibodies for IP (Figure 6C). Collectively, FRET and Co-IP experiments demonstrated that host cell vimentin could bind with TgROP18.

TgROP18 affects the dynamic patterns of host cell vimentin solubility, phosphorylation, and expression

Given the interaction between host cell vimentin and TgROP18, and with the aim of exploring the effect of TgROP18 on vimentin, the solubility, phosphorylation, and expression levels of vimentin were determined in Cos7 cells infected with RH Δrop18 and RH WT. Host cell vimentin solubility was significantly different between cells infected with RH WT and RH Δrop18 at 1, 6, 24, 36, and 48h, suggesting that the solubility of vimentin was partly affected by TgROP18, in addition to other unknown parasitic factors (Figure 7A, B, and C). To further investigate the effects of TgROP18 on vimentin solubility, Cos7 cells were transfected with pcDNA3.1-ROP18-flag. Much higher vimentin solubility was observed in cells over-expressing TgROP18 than in controls (Figure 8A and B). The phosphorylation level of vimentin in Cos7 cells infected with RH Δrop18 was also evaluated using Phos-tag assays, and the results indicated that there was a trend towards up-regulation of vimentin with the extension of infection time (Figure 7D). Comparison of these results with the changes in vimentin phosphorylation levels in cells infected with RH WT for different periods of time (Figure 3), suggested that TgROP18 affects the phosphorylation of vimentin (Figure 7D). To further explore the effects...
of ROP18 on vimentin, kinase assays in vitro were performed to determine whether vimentin was phosphorylated by the \( Tg\)ROP18 kinase. The results suggested \( Tg\)ROP18 could phosphorylate host cell vimentin in vitro (Figure 7E). Intriguingly, the changes in vimentin solubility and phosphorylation levels in cells infected with RH WT (Figure 7) were much more complex than those in cells over-expressing ROP18 (Figure 8), suggesting that the dynamic patterns of vimentin solubility and phosphorylation induced by \( T. gondii \) infection were partly regulated by \( Tg\)ROP18, and may contribute to the reorganization of host cell vimentin. Furthermore, compared with the protein levels of vimentin in Cos7 cells infected with RH WT (Figure 3D), no significant changes were observed in cells infected with RH \( \Delta rop18 \) (Figure 7F), indicating that the expression levels of vimentin were not clearly affected by \( Tg\)ROP18. Additionally, over-expression of ROP18 in Cos7 cells did not have obvious effects on host cell vimentin expression (Figure 8C).

Figure 4. Function analysis of host cell vimentin on the infection of \( T. gondii \). A. Verification of vimentin knockout in HBMEC by WB. B. Effect of host cell vimentin on the invasion of \( T. gondii \) into host cells. HBMEC and HBMEC \( \Delta \)Vim cells were infected with \( T. gondii \) RH/GFP for 30min and host-cell invasion was evaluated by a two-color assay to distinguish intracellular from extracellular parasites. Data are expressed as means ± SD from three independent experiments, each performed in triplicate, and analyzed by t-test (**\( p < 0.01 \)). C. Representative fluorescence images of B. Cell nuclei were stained with DAPI and extracellular parasites were stained with mouse anti-SAG1 antibody (scale bars, 10μm). D. Effect of vimentin on the proliferation of \( T. gondii \) in host cells. \( T. gondii \) RH/GFP were used to infect HBMEC and HBMEC \( \Delta \)Vim, and invaded parasites were allowed to replicate for 22 h. The number of vacuoles containing one, two, four, or eight parasites was visualized under a fluorescence microscope (100×). Data are expressed as means ± SD combined from three independent experiments, each performed in triplicate and analyzed by two-way ANOVA. No significant difference was identified between the two comparison groups. E. Representative fluorescence images of D (scale bars, 10μm). F. Effect of vimentin on the egress of \( T. gondii \). HBMEC and HBMEC \( \Delta \)Vim were infected with RH/GFP at a MOI of 3. After infection, \( T. gondii \) were cultured in cells for 30h and then induced to egress using A23187; DMSO was used as a control. Data are expressed as means ± SD combined from three independent experiments, each performed in triplicate, and analyzed by t-test. G. Representative fluorescence images of F (scale bars, 10μm). White arrows indicate the non-released PVs, and brown arrows indicate the egressed tachyzoites from one PV.
Figure 5. Differences in vimentin mRNA and protein expression in various mouse tissues. A. Transcriptional levels of vimentin were detected by qRT-PCR. Data are expressed as means ± SD combined from three independent experiments and were analyzed by t-test. ***p≤0.01, ****p≤0.001. B. WB was performed to detect the expression of vimentin in different mouse organs. Beta-actin was used as a loading control. The lowest transcription and expression levels were observed in mouse brain, relative to the other organs.

Figure 6. Identification of the interaction between host cell vimentin and TgROP18. A. FRET was used to investigate the interaction between vimentin and ROP18. Image acquisition and parameter calculation were performed using the sensitized emission (SE) method. B. The calculated efficiency of FRET was approximately 55% (t-test, **p<0.01). C. Interaction of vimentin with ROP18 was further demonstrated by immunoprecipitation assay. HFF cells were infected with T. gondii RH-ROP18-eGFP-FLAG at a MOI of 5 for 36h and then lysed for immunoprecipitation with mAbs anti-vimentin and anti-FLAG. Starting fractions (Input) and immunoprecipitates (IP) were analyzed by WB using RBs anti-vimentin and anti-DDDDK.
Figure 7. Dynamic patterns of vimentin solubility and phosphorylation were partially affected by TgROP18. A, B. Cos7 cells were infected with T. gondii RH Δrop18. The solubility of vimentin was then detected and analyzed by WB. C. Comparison analysis of vimentin solubility in Cos7 cells infected with T. gondii RH and RH Δrop18. (t test, *p<0.05) D. Analysis of vimentin phosphorylation in Cos7 cells infected with RH Δrop18. E. TgROP18 phosphorylates host cell vimentin. Recombinant GST-ROP18 and vimentin proteins expressed and purified from E. coli were co-incubated in the presence of unlabeled ATP in vitro. Total vimentin and phosphorylated vimentin were detected with rabbit anti-vimentin and anti-phospho Ser/Thr antibodies, respectively. The phosphorylated vimentin band was only visible in the presence of TgROP18. F. Analysis of vimentin expression levels in host cells infected with RH Δrop18.

Discussion

Host cell vimentin has been demonstrated to function in facilitating the infection of some adhesive or invasive microbial pathogens, including P. multocida and African swine fever virus [29, 30]. Moreover, vimentin is identified as a novel NF-κB regulator and is involved in modulating the NF-κB signaling pathway to influence invasion of pathogens, such as E. coli K1 [31, 32]. The disassembly of vimentin caused by treatment with high concentrations of the phosphatase inhibitors, okadaic acid and calyculin A, suggests that vimentin’s assembly/disassembly is regulated by its phosphorylation and dephosphorylation [20]. The assembly and solubility of vimentin are regulated by the phosphorylation/dephosphorylation of vimentin at Ser38 and Ser55 induced by Rab7a [33]. Moreover, phosphorylation level of vimentin Ser38 was significantly upregulated in host cell infected with T. gondii at both 2h and 6h comparing to host cell without T. gondii infection [34]. Consistent with the above reports, higher vimentin expression corresponding to much lower solubility in host cell with T. gondii infection for 2h and 6h, which also suggest that vimentin phosphorylation is associated with its solubility. The reorganization of cellular vimentin is also affected by its solubility, which is, in turn, regulated by its phosphorylation [35, 36]. Progressive rearrangement of vimentin around the enlarging PVs was reported at 30min, 12h, and 18h post-infection [20], and vimentin forms a dynamic, flexible network and contributes to maintenance of the integrity of the cellular actin cytoskeleton [37]. Consistent with previous findings, a significant rearrangement of vimentin was also observed in our study in cells infected with T. gondii for different periods of time (1, 2, 6, 12, and 22h) post-infection.
from invasion to multiplication (Figure 2), and variable changes in vimentin solubility, phosphorylation, and expression levels were also observed during *T. gondii* infection at different time points (1, 2, 6, 12, 24, 36, and 48h) (Figure 3).

*TgROP18* is a key serine/threonine kinase that phosphorylates host proteins to modulate the acute virulence of *T. gondii* [25, 26, 27, 38], regulates host cell signaling pathways [39, 40], prevents host cell apoptosis [41], and alters host cell gene expression [42]. *TgROP18* can interact with, and phosphorylate host cell proteins, such as ATF6β and Irga6, to facilitate parasite infection [23, 25, 27]. Interestingly, host cell vimentin was found to interact with *TgROP18* in our previous protein interaction screening based on BiFC, and the results of this study verified those findings further using FRET and Co-IP experiments (Figure 6). In our previous phosphoproteinomics assays of *T. gondii* infected HFF cells, host cell vimentin exhibited a complex phosphorylation pattern at different infection times [34]. In this study, a dynamic phosphorylation pattern of vimentin was identified in host cells infected with *T. gondii* RH and RH Δrop18 strains using phos-Tag assays, and the results indicated that the phosphorylation pattern of vimentin was partly affected by *T. gondii* ROP18; moreover, vimentin phosphorylated by *TgROP18* was also demonstrated by our kinase assay in vitro (Figure 7).

The dynamic exchange between the assembled and disassembled pools of vimentin in *vivo* is phosphorylation dependent, and phosphorylation of this protein also affects its solubility [33, 43]. To determine whether the solubility of vimentin was affected by *TgROP18*, vimentin solubility assays were performed using Cos7 cells infected with *T. gondii* RH and RH Δrop18, and the results indicated that vimentin solubility was partly regulated by *TgROP18*, in addition to other unknown parasitic factors (Figure 7). We also found that the expression levels of vimentin were regulated by the *T. gondii* infection; however, they were not significantly affected by *T. gondii* ROP18 (Figure 3D and 7F). In summary, the dynamic patterns of vimentin phosphorylation, solubility, and expression levels were regulated by *T. gondii* infection, while its phosphorylation level and solubility were only partly affected by *TgROP18*. Hence, while vimentin has important roles in *T. gondii* infection, only its phosphorylation and solubility, but not its expression levels, are affected by *TgROP18*.

**Figure 8. Effect of overexpression of *TgROP18* on Cos7 cell vimentin solubility.** A. Verification of over-expression of ROP18 in Cos7 cells and determination of the percentage of soluble vimentin, relative to total vimentin, based on WB signals. B. Analysis of the percentage of soluble vimentin, relative to total vimentin, using Image J software. Data are expressed as means ± SD combined from three independent experiments and were analyzed by t-test. **p<0.001.** Much higher vimentin solubility was observed in cells overexpressing *TgROP18*. C. Detection of vimentin expression in Cos7 cells overexpressing *TgROP18* (transfected with pcDNA3.1-ROP18) and controls (untransfected cells (ctrl) and cells transfected with pcDNA3.1). The results indicate that the expression of vimentin was not affected by *TgROP18* overexpression.
Maintenance of host cell actin cytoskeleton integrity is important during the *T. gondii* lytic cycle, including invasion and egress. For example, host cell actin cytoskeleton integrity is essential for successful *T. gondii* invasion [12], since *T. gondii* invasion was blocked when host cell actin polymerization was inhibited using cytochalasin D [44]. There is evidence that actin organization and actin cytoskeleton integrity may be modulated by host cell vimentin [45].

Herein, in our present study we found that *T. gondii* invasion was affected by host cell vimentin. When vimentin was knocked out in HBMEC cells, the infection rate of *T. gondii* was significantly increased (Figure 4). The result, indicating inhibition of host cell vimentin on *T. gondii* invasion, is opposite to those demonstrated for *Escherichia coli* K1 and Japanese encephalitis virus infection [31, 46], suggesting that vimentin regulates infection of *T. gondii* and other microbial pathogens by different mechanisms.

In our study, we observed variable levels of vimentin protein expression in different mouse tissues, with much lower levels in brain and spleen compared with the other tissues investigated (lung, kidney, liver, and heart; Figure 5). *T. gondii* exhibits a definite neurotropism and much higher number of cysts located in the brain than other organs [47]. Low vimentin expression levels favor *T. gondii* invasion into host cells, which is consistent with the fact of more cysts in the brain than in other organs. Strong and persistent cell-mediated immunity, mainly Th1 responses, is activated by *T. gondii* infection, and can protect the host from rapid tachyzoite proliferation and consequent pathology [48, 49]. CD4+ and CD8+ T cell populations are increased in the spleens of mice immunized with *T. gondii* [50]; therefore, although low expression levels of vimentin were also observed in this organ, it develops no more cysts than other tissues.

Together, the findings of this study indicated that host cell vimentin rearrangement around PVs occurred during *T. gondii* infection, and that phosphorylation and solubility of vimentin were regulated by *T. gondii* infection, but only partially by TgROP18, which may be associated with vimentin reorganization to facilitate *T. gondii* invasion. Moreover, expression levels of vimentin were affected by *T. gondii* infection, but not by TgROP18. Furthermore host cell vimentin was phosphorylated by TgROP18 and played an important role in the inhibition of *T. gondii* invasion. Our findings revealed a new mechanism of *T. gondii* regulation on host cell cytoskeleton reorganization through its effects on vimentin phosphorylation, solubility, and expression, and the contribution of TgROP18 to this regulatory process.

**Materials and Methods**

**Ethics statement**

Animal experiments were approved by the Ethical Committee for Animal Research of Southern Medical University and conducted based on the state guidelines from the Ministry of Science and Technology of China. All KunMing mice (18–22g, female) were purchased from the Laboratory Animal Center of Southern Medical University.

**Parasite culture**

All *T. gondii* strains used in this research were derived from the type I RH line and tachyzoites were maintained by growth in human foreskin fibroblasts (HFF). The HFF cell line was obtained from American Type Culture Collection (ATCC), and grown in Dulbecco’s Modified Eagles Medium (DMEM, Thermo Fisher Scientific) supplemented with 1% fetal bovine serum (FBS) (Gibco, Thermo Fisher Scientific). Transgenic parasites were obtained by electroporation of recombinant plasmids into wild type RH tachyzoites and selection with 3μM pyrimethamine (Sigma). Freshly egressed parasites purified with 3μm nucleopore filters were used in all experiments.

**Cell culture**

HFF and Cos7 cells were cultured in DMEM containing 10% FBS (Thermo Fisher Scientific) and 1% gentamicin (10mg/ml). Human brain microvessel endothelial cells (HBMEC) and HBMEC ΔVim were cultured in RPMI 1640 medium (DMEM, Thermo Fisher Scientific) supplemented with 10% FBS (MUETICELL, WISENT INC.), 1% Non-Essential Amino Acids (NEAA, Thermo Fisher Scientific) and 1% Sodium Pyruvate (Thermo Fisher Scientific).

**Fluorescence Resonance Energy Transfer Experiments**

One group of Cos7 cells were co-transfected with pECFP-ROP18 and pEYFP-Vim plasmids to generate cells overexpressing ROP18 tagged with enhanced cyan fluorescence protein (eCFP) and vimentin tagged with enhanced yellow fluorescence protein (eYFP). Two other groups of Cos7 cells were transfected with pECFP-ROP18 or pEYFP-Vim separately as controls. Simultaneously, a fourth group of Cos7 cells transfected with pEYFP-CFP was used as a positive control, and a fifth group of Cos7 cells co-transfected with pECFP and pEYFP was used as a negative control. FRET between eCFP-ROP18 and eYFP-vimentin was measured using the sensitized emission (SE) program under an Olympus FLUOVIEW FV1000 confocal laser scanning microscope 48h post transfection [51, 52]. Data
information of the images derived from the five groups of samples described above was presented in Table S4. FRET efficiencies were calculated according to the formula indicated in Table S4.

Vimentin solubility assay

Cos7 cells were grown in 6-well culture plates and infected with RH tachyzoites for 0, 1, 2, 6, 12, 24, 36, and 48 h, then lysed with a high salt, detergent buffer (20 mM Tris-HCl, pH 7.4; 600 mM NaCl; 0.5% Triton X-100; 0.1 mM sodium orthovanadate; and protease inhibitors (Roche)), and the soluble fraction separated from the insoluble fraction by centrifugation at 12,000 g for 10 min at 4°C [53]. Equal volumes of soluble and insoluble fractions were loaded for SDS-PAGE and vimentin was detected with mouse anti-vimentin monoclonal antibody by western blotting (WB). Beta-actin was also detected by WB using a monoclonal antibody as a loading control. Details of primary and secondary antibodies are provided in Supplementary Material. The intensity of vimentin bands with non-saturated exposure from three independent experiments was analyzed using Image J software, and the proportion of soluble vimentin to total vimentin was calculated.

Phos-tag assay

Phos-tag acrylamide (Wako Chemicals) based gel analysis was carried out to detect phosphorylated vimentin. Cos7 cells were infected with T. gondii RH or RH Δrop18 strains at a MOI of 3 for different periods of time (0, 1, 2, 6, 12, 24, 36, and 48 h). Cells were then lysed with cell lysis buffer (TransGen Biotech, DE101) and the extracts were loaded on to 10% polyacrylamide gel (separating gel) containing 50 μM Phos-tag acrylamide (Wako Chemicals, AAL-107). As vimentin expression levels changed with the time post T. gondii infection, the amount of cell lysate protein loaded for SDS-PAGE was adjusted to ensure a consistent level of total vimentin protein. After electrophoresis, Phos-tag acrylamide gels were washed three times with gentle shaking in transfer buffer containing 1 mM EDTA for 10 min and then incubated in transfer buffer containing 0.01% SDS without EDTA for 10 min, according to the manufacturer’s protocol. Proteins were then transferred to polyvinylidene difluoride (PVDF) membranes using a standard immunoblotting protocol, and vimentin detected using a specific antibody (Supplementary Material) [54].

Immunofluorescence microscopy

To examine the rearrangement of host cell vimentin after T. gondii infection, cells were grown to 80% confluence on coverslips and then infected with T. gondii at a MOI of 3 for different periods of time. Resulting parasite-infected monolayers were fixed with 4% formaldehyde for 15 min, permeabilized with 0.1% Triton X-100 for 20 min at room temperature, and blocked in 10% FBS for 15 min [55]. Samples were incubated with primary antibodies overnight at 4°C, and then washed in PBS with gentle shaking at RT three times for 5 min, and incubated with secondary antibodies (goat anti-mouse IgG or goat anti-rabbit IgG) conjugated with FITC, Texas Red®, or TRITC, diluted 1:200. Following staining, slides were again washed with PBS (as described above), and then rinsed with ddH2O. Air-dried slides were mounted with DAPI Fluoromount® (Southern Biotech). Samples were visualized and images generated using an Olympus FLUOVIEW FV1000 confocal laser scanning microscope or by fluorescence microscopy.

Immunoblotting and co-immunoprecipitation

For western blotting, cells or parasites were lysed in lysis buffer (Beyotime Biotechnology) for 30 min at 4°C. Cell debris was precipitated by centrifugation at 12,000 g at 4°C for 10 min. Supernatants were boiled in SDS-PAGE sample buffer and loaded for SDS-PAGE. Proteins were separated on polyacrylamide gels, and then transferred to PVDF membranes. Membranes were blocked in PBS containing 5% Bovine Serum Albumin (BSA) and 0.05% Tween-20 and probed with primary antibodies, followed by secondary antibodies conjugated with horseradish peroxidase (HRP) (See Supplementary Material for details). Specific proteins on membranes were visualized by luminescence generated using Clarity™ Western ECL Substrate (Bio-Rad) and photographed with a ChemiDoc™ Touch Imaging System (Bio-Rad).

HFF cells infected with T. gondii RH-ROP18-eGFP-FLAG for 36 h and uninfected controls were lysed and clarified by centrifugation as described above. For co-immunoprecipitation (Co-IP), mouse monoclonal vimentin or FLAG antibodies were added to the clarified cell extracts and incubated at 4°C for 1 h. Then Protein A-Agarose (Santa Cruz) was added to the mix and incubated for 1 h at 4°C with rotation. Beads were collected by centrifugation at 1,000 g for 5 min at 4°C, and then washed for four times with PBS with rotation for 10 min at 4°C. Finally, beads were collected by centrifugation as described above, resuspended in SDS-PAGE sample buffer, and then boiled for 10 min. Boiled samples were centrifuged briefly to eliminate debris, loaded for SDS-PAGE, and then analyzed with rabbit monoclonal anti-vimentin or polyclonal anti-DDDDK (for FLAG tag detection) antibodies by WB (Supplementary Material).
PCR and Quantitative Reverse Transcription PCR (qRT-PCR)

PCRs were performed using premix Taq (Takara) and Pfu DNA polymerase (TransGen Biotech) according to the manufacturer’s directions. The primers used for PCR are presented in Table S2.

Levels of vimentin mRNA were compared by qRT-PCR to determine the efficiency of RNA interference in HFF cells and transcription levels of vimentin in different mouse tissues. Total RNA samples were extracted from tissues or HFF cells using Trizol® Reagent (AmbionTM) and cDNAs were synthesized using the gDNA Removal and cDNA Synthesis SuperMix (TransGen Biotech, China). qRT-PCR was performed using Top Green qPCR SuperMix (TransGen Biotech, China) according to the manufacturer’s instructions. The primers used are listed in Table S2. Beta-actin was used as an internal control for normalization.

T. gondii invasion efficiency assay

Invasion assays were performed based on differential staining of intracellular vs. extracellular parasites, as previously described [8]. T. gondii RH/GFP tachyzoites harvested by mechanical lysis were allowed to invade HBMEC and HBMEC ΔVim monolayers for 30 min at 37°C and then fixed with 4% paraformaldehyde. Extracellular parasites were detected by staining with mouse anti-SAG1 antibody. Both intracellular and extracellular parasites expressing GFP were observed by fluorescence microscopy. Slides were examined by epifluorescence microscopy and the total numbers of parasites (green), extracellular parasites (red), and host cell nuclei (blue) were counted from counts of ≥200 parasites. Values are expressed as the average number of parasite/host cell nuclei and represent means ± SD of three independent experiments.

T. gondii proliferation assay

HBMEC and HBMEC ΔVim cells, or HFF cells pretreated with siRNA, were infected with T. gondii RH/GFP tachyzoites for 30 min, the non-invaded parasites were thoroughly washed off with PBS three times, and cells grown in DMEM complete medium for 22h. Subsequently, cell monolayers were fixed and examined to determine the number of vacuoles containing one, two, four, or eight parasites under a fluorescence microscope (100×). Means ± SD combined from three independent experiments, each performed in triplicate, were analyzed by two-way ANOVA [56].

Induced egress assay

To evaluate the effect of vimentin on the egress of T. gondii, induced egress assays were performed as previously described [57]. Freshly egressed T. gondii RH/GFP were purified as described above and then added to HBMEC and HBMEC ΔVim cells and grown for approximately 30 h at 37°C. Infected host cells were incubated for 7 min at 37°C with serum-free RPMI 1640 medium containing either 3μM of the Ca²⁺ ionophore, A23187 (Sigma, 21186) to induce egress, or DMSO as a control. Cells were fixed with 4% paraformaldehyde and visualized directly by fluorescence microscopy. One hundred vacuoles were counted per group in three independent experiments, and the number of lysed vacuoles was scored.

Kinase assay in vitro

Bacterially derived, purified GST-ROP18 (200ng) was incubated with vimentin (3μg) in reaction buffer (1mM unlabeled ATP, 25mM Tris/HCl (pH 7.5), 15mM MgCl₂) for 30min at 30°C. Then, samples were immediately boiled in sample buffer for 5 min for western blot analysis. Equal purified GST was incubated with vimentin and detected as control according to above description.

Statistical analysis

All experiments were performed at least in triplicate. Data are presented as means ± SD, unless otherwise indicated. SPSS 13.0 software was used for statistical analyses. Differences between groups were analyzed using t-test, or two-way ANOVA. Values of p < 0.05 were considered statistically significant.

Supplementary Material


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Author Contributions

C.H., performed the experiments, analyzed the data and wrote the manuscript. L.K., performed functional assays. L.Z., performed Co-IP. J.X., performed FRET. H.W., cell culture. M.L., generated
ectopically expressed T. gondii. HP., study conception and design, supervision of the research group, funding support, and drafting the manuscript. All authors read and approved the final manuscript.

Competing Interests
The authors have declared that no competing interest exists.

References