Immune response against toxoplasmosis—some recent updates RH: *Toxoplasma gondii* immune response

Madiha Sana¹,*, Muhammad Rashid²,*, Imran Rashid¹, Haroon Akbar¹, Jorge E Gomez-Marin³ and Isabelle Dimier-Poisson⁴

Abstract

**Aims:** Cytokines, soluble mediators of immunity, are key factors of the innate and adaptive immune system. They are secreted from and interact with various types of immune cells to manipulate host body’s immune cell physiology for a counter-attack on the foreign body. A study was designed to explore the mechanism of *Toxoplasma gondii* (*T. gondii*) resistance from host immune response.

**Methods and results:** The published data on aspect of host (murine and human) immune response against *T. gondii* was taken from Google scholar and PubMed. Most relevant literature was included in this study. The basic mechanism of immune response starts from the interactions of antigens with host immune cells to trigger the production of cytokines (pro-inflammatory and anti-inflammatory) which then act by forming a cytokinome (network of cytokine). Their secretory equilibrium is essential for endowing resistance to the host against infectious diseases, particularly toxoplasmosis. A narrow balance lying between Th1, Th2, and Th17 cytokines (as demonstrated until now) is essential for the development of resistance against *T. gondii* as well as for the survival of host. Excessive production of pro-inflammatory cytokines leads to tissue damage resulting in the production of anti-inflammatory cytokines which enhances the proliferation of *Toxoplasma*. Stress and other infectious diseases (human immunodeficiency virus (HIV)) that weaken the host immunity particularly the cellular component, make the host susceptible to toxoplasmosis especially in pregnant women.

**Conclusion:** The current review findings state that in vitro harvesting of IL12 from DCs, Np and MΦ upon exposure with *T. gondii* might be a source for therapeutic use in toxoplasmosis. Current review also suggests that therapeutic interventions leading to up-regulation/supplementation of SOCS-3, IL12, and IFNγ to the infected host could be a solution to sterile immunity against *T. gondii* infection. This would be of interest particularly in patients passing through immunosuppression owing to any reason like the ones receiving anti-cancer therapy, the ones undergoing immunosuppressive therapy for graft/transplantation, the ones suffering from immunodeficiency virus (HIV) or having AIDS. Another important suggestion is to launch the efforts for a vaccine based on GRA6Nt or other similar antigens of *T. gondii* as a probable tool to destroy tissue cysts.

¹Department of Parasitology, University of Veterinary and Animal Sciences, Lahore, Pakistan
²Department of Parasitology, Faculty of Veterinary and Animal Sciences, The Islamia University of Bahawalpur, Pakistan
³Grupo Gepamol, Centro de Investigaciones Biomedicas, Universidad del Quindio, Armenia, CO, South America
⁴Université de Tours, Institut national de recherche pour l’agriculture, l’alimentation et l’environnement (INRAE), Unité mixte de recherche 1282 (UMR1282), Infectiologie et santé publique (ISP), Tours, France

*The first and second authors equally contributed

**Corresponding author:** Imran Rashid, Department of Parasitology, University of Veterinary and Animal Sciences, Sheikh Abdul Qadir Jillani Road, Lahore 54200, Pakistan. Email: imran.rashid@uvas.edu.pk

Creative Commons Non Commercial CC BY-NC: This article is distributed under the terms of the Creative Commons Attribution-NonCommercial 4.0 License (https://creativecommons.org/licenses/by-nc/4.0/) which permits non-commercial use, reproduction and distribution of the work without further permission provided the original work is attributed as specified on the SAGE and Open Access pages (https://us.sagepub.com/en-us/nam/open-access-at-sage).
Introduction

Toxoplasmosis is a zoonotic infectious disease caused by an intracellular protozoan pathogen called Toxoplasma gondii (T. gondii). It causes the abortion or congenital abnormalities (hydrocephalus and retinocochoroiditis) in pregnant women which are more susceptible to Toxoplasma infection. Generally, the immune system of immunocompetent individuals builds a protective immune response upon the interaction of antigen with antigen-presenting cells (APC) which induces the translocation of nuclear factor kappa (NF-κB) to initiate production of pro-inflammatory cytokines (IL1b, IL12, IL18, and IFNγ). The host lymphocytes and myeloid cells not only secrete a network of cytokines for signaling pathways upon exposure to antigen but also up-regulate certain chemokines (CXCl, C, and CX3C) and toll-like receptors (TLR) on their surfaces for acting as signal-recipients against any antigen. In response to specific intracellular signals, various pro-inflammatory (IL1b, IL12, IL18, TNFα, IFNγ) and anti-inflammatory (IL4, IL10, TGFβ) cytokines give rise to a cytokinome that acts for specific immunological stimulus to develop immune response for susceptibility or resistance. Immunological studies on human infections have clearly concluded that cell-mediated immunity and IFNγ are paramount in the control of any infection particularly caused by the intracellular pathogen (T. gondii). The inability of the humoral response alone (antibodies) to prevent Toxoplasma reactivation is evident by the fact that most of the HIV-infected people lack effective immunity hence exhibit symptoms of T. gondii infection and reactivation, even in the presence of high titers of specific IgG.

Pro-inflammatory cytokines are among the key factors to initiate and maintain innate as well as acquired immunity to restrict proliferation of Toxoplasma. A variety of cytokines are produced upon activation of APCs and cells of the adaptive immune system (B and T cells). The differences in cytokinome can be speculated at the different stages of infection, due to intra- or extra-cellular nature of pathogens as well as due to diversity of the host genetic makeup. The indirect role of cytokines against T. gondii in leading to either resistance or susceptibility also depends upon the stage of parasite in the host and the induction and modulation of pro-inflammatory cytokines driven by the particular parasite strain as well as the robustness of the host’s immune profile. Different typical and atypical strains of T. gondii exist globally, and have been specifically studied in America and Europe. This parasite is identified having three distinct genotypic lineages in humans: type I strain (RH-88), type II strain (ME49, and DEG), and type III strains (CEP and VEG). The genetic moieties in these strains result into a highly varied level of virulence which inflicts diverse pathological effects in host by a variety of cytokine pathways as well as owing to the wide range of interactions through a vast diversity of host and parasite molecules interacting each other, with some known and some unknown footprints. The known footprints include but not limited to inherent-oxidative stress, a diversity of IRGs (Immunity Related GTPases) of host with a locus on chromosome 11, Z-DNA binding protein 1 (ZBP1), receptor-interacting serine/threonine-protein kinase 3 (RIPK3) of host. Likewise, the wide range of parasite molecules includes a diversity of rhoptry proteins (e.g. ROP5, ROP18), dense granule proteins (e.g., GRA5, GRA12, GRA16, GRA24); small GTPase immunity-associated proteins (GIMAPs 4, 5 & 6) of T. gondii. This review describes the cytokinome in toxoplasmosis and their interactive role for development of host’s susceptibility and resistance toward T. gondii infection.

Database search

A search was conducted on Science Direct (https://www.sciencedirect.com/science/search) and PubMed (https://www.ncbi.nlm.nih.gov/pubmed) using “host immune response and resistance of toxoplasmosis” to achieve the relevant literature for this study. The searched period ranged from 1980 until 15 September 2021, yielding 649 publications. The literature having mechanism of immune response and pathogen resistance were included in this study. Whereas for the last 5 years, the data were searched from PubMed using the same key words as mentioned above and majority of the articles with novel insights into the immune response mechanism as well as host parasite interactions were included in this review. Although maximum efforts have been done to review the literature on “host immune response and resistance of toxoplasmosis”, certainly there are limitations of this review. Hence, this should not be considered a review encompassing all the new literature on ‘Toxoplasma gondii” and “Toxoplasmosis”. This is because of the fact that 24,912 and 13,928 results are retrieved from Science Direct and PubMed, respectively, while using a key word “Toxoplasmosis” with the same time period as mentioned above. Likewise, 23,158 and 15,750 results are displayed from Science Direct and PubMed, respectively, with a key word “Toxoplasma gondii” for the same time period. Similarly, slightly modified key words, that is, “host immune response of toxoplasmosis” displays almost 7000 (6969) results in Science Direct.

Keywords
Toxoplasma gondii, cytokinomes, host immune response, host resistance, susceptibility
1) **Virulence of T. gondii strains**

In humans, the regulatory cytokines profile depends on the T. gondii strain. Among three notable strains (type I, II and III) of T. gondii, the type I is more virulent as compared to type II and type III strains. In type I (RH strain) infected cells, the translocation of NF-κB does not take place resulting in the production of anti-inflammatory (IL10, IL27, and TGFβ1) cytokines which are higher as compared to uninfected cells.\(^{19-23}\) These anti-inflammatory cytokines enhance the proliferation of T. gondii. It was found that with a significantly high level of TGFβ1 in the blood as well as in the aqueous humor of the acutely intraocular Toxoplasma-infected host, it may adversely interfere with the effective cellular immune response leading to an increased mortality and extensive ocular tissue damage with tachyzoites observed in the pigment epithelium layers.\(^{24}\) Consequently, Th2 and Treg responses are enhanced in comparison with a primary ocular infection.\(^{25}\) ME49 is a type II strain that induces the translocation of NF-κB-light-chain-enhancer from the cytoplasm to the nucleus of activated B cells, splenocytes\(^{26}\) and bone marrow-derived MΦ\(^{27}\) which induce the production of pro-inflammatory cytokines (IL1β, IL12, IL18, TNFα, IFNγ, and IL12p40) by thioglycolated MΦ/cell lines\(^{19,28}\). Whereas, the production of anti-inflammatory cytokines are lower in ME49-infected cells than that of uninfected cells\(^{21}\) speculating CD36-associate dense granule proteins in modulating parasitic virulence while interacting the host body’s resistance mechanisms like GRA12 of Toxoplasma was identified as a major virulence factor to counter the host’s IFNγ.\(^{30}\)

**Host immune response**

**A) Innate immune response**

In early stages of infection, dendritic cells (DC), macrophages (MΦ), natural killer (NK), and neutrophils (Np) interact in a coordinated way to provide the first line of defense in the form of innate immune response leading to develop adaptive immunity.\(^{31-33}\) The innate immune response is elicited against toxoplasmosis in the form of IL12 production upon interaction with antigen (Ag). The release of IL12 from MΦ, DCs, and Np is essential for the release of IFNγ from NK cells (innate immune response) and T lymphocytes (adaptive immune response) via antigen presentation.\(^{34}\) IFNγ has been shown to induce guanylate binding proteins (GBPs) in a murine model of toxoplasmosis, thereby, these GBPs accumulate on the surface of intracellular parasites potentially causing parasitic destruction, thus displaying an active role of intracellular autonomous immunity.\(^{35}\) The increased susceptibility toward T. gondii infection is due to the depletion of NK cells, MΦ, or DCs which have a significant involvement for innate immune response against the infection.\(^{36}\) The mechanism of innate immune response initiate upon interaction of toll-like receptors (TLRs) with ligands expressed on T. gondii surface, thereby starting intracellular signaling pathways through engagement of the myeloid differentiation domain-88 (MyD88). These are the universal adaptor proteins involved in signaling of all TLRs except TLR-3. The study on MyD88 deficient mice model was impaired to induce primary protection in acute infection of T. gondii (RH strain).\(^{37}\) Studies on mouse with targeted inactivation of MyD88 showed that DCs work as antigen-presenting cells (APC) and are responsible for the increased susceptibility to T. gondii infection. However, there was no effect on MΦ and Np. The MyD88 deficient mice masked the production of IL12 from DCs and IFNγ by Nks to initiate innate immune response. It explains a central role of DCs in the coordination of innate immune response against T. gondii infection and predicts the increased susceptibility towards infection if DCs are recognized defective in early encounter of T. gondii infection.\(^{38}\)

**B) Cellular immune response**

In T. gondii infection, the strong resistance to re-infection as well as the hindrance to reactivation of chronic infection is based on the host’s cell-mediated immunity.\(^{39,40}\) The synergistic role of CD4\(^+\) and CD8\(^+\) T cells for the development of acquired immunity was understood from targeted experiments on C57BL/6 mice vaccinated with temperature sensitive mutant strain of T. gondii (ME49).\(^{41,42}\) The development of complete immunity against a virulent strain (type I) is dependent on IFNγ synthesis from NK and T cells. The immunocompetent host activates either T cells or NK cells for encountering parasitic invasion. In a previous study, the MHC I (lack of CD8\(^+\) cell stimulation) impaired mice (beta 2m-deficient mice) surprisingly showed high resistance against T. gondii following vaccination. This enhanced immunological response in the absence of CD8\(^+\) cells showed the involvement of NK cells activated by IL12 upon parasitic infection.\(^{43}\)

Generally, the CD8\(^+\) T cells are the major source of IFNγ production against most of the T. gondii strain.\(^{44}\) The c-Rel expression regulated by NF-κB is widely dominant in hematopoietic cells,\(^{45}\) which play a critical role in the development of resistance against T. gondii (ME49) infection.\(^{46}\) The role of c-Rel in influencing the CD8\(^+\) cell response was investigated in mice model (c-Rel\(^{−/−}\) mice) with a special infection of T. gondii strain (replication-deficient strain) in C57BL/6, CD45.1, and Thy1.1 mice. The CD8\(^+\) cells impair to replicate in the absence/
deficiency of c-Rel. Likewise, the c-Rel deficient mice remained unable to survive during infection with the replicating pathogen.47,48

CD8⁺ T immune cells having a TCR Vβ8.1, 8.2+ phenotype produce protection against tissue cyst development in mouse model. Transfer of CD8⁺ cells induced by N-terminal of dense granule protein-6 (GRA6Nt) of parasite has been shown to clear T. gondii cysts from the brains of infected mice that were deficient in T cells (Sa et al. 2017), further highlighting the role of cytotoxic T cells in the induction of protective immunity against T. gondii. A genetically resistant strain of mice having a H-2d haplotype helped discover these specific type of cytotoxic T cells, whereas H-2Ld was found as a major antigen-presenting molecule to CD8⁺ T cells to achieve this objective of tissue cyst elimination.

The immune response of CD8⁺ cells is more dominant alike effector cells than CD4⁺ cells. Nevertheless, CD4⁺ helper T cells are direly required for an effective functioning molecule to CD8⁺ T cells to achieve this objective of tissue cyst elimination.

Regulatory T (T reg) cells are required for the maintenance of immunological self-tolerance and immune homeostasis by actively suppressing the pathological and physiological immune responses. An IL2 knocked-out mouse model orally infected with lethal dose of T. gondii, showed highly Th1 cell type-polarized mucosal immune response. Such effect contributed to the incapacity of T reg cells to perform effector responses and consequently led to immuno-pathogenesis.54,55 Besides this, T reg cells of infected mice expressed lower levels of Bcl-2 and increased levels of apoptotic markers than that of naive mice. It is suggested that de-regulation in the T reg cells is a consequence of these impaired cells turnover.54 Besides this, there was found a gradual weight loss and significant delayed mortality in T reg-transferred toxoplasmosis-infected mice associated with lower level of IFNγ and TNFα. Additionally, higher cyst number and parasite load in brain of those mice were observed.55 Furthermore, activity of T reg cells results in the death of proliferating T cells which favors the multiplication of pathogen in the murine model.57 In a pregnant T. gondii-infected mouse model, numbers of splenic CD4⁺CD25⁻ T reg cells and placental Foxp3⁺ cells decreased synchronously. During infection, the reduction of splenic CD4⁺CD25⁻ T reg cells was associated with apoptosis (Bcl-2) induced by proliferating T cells.58 Additionally, injection of pregnant mice with excretory–secretory antigens (ESA) of T. gondii also causes fetal death associated with apoptosis of CD4⁺CD25⁺ T reg cells by down-regulating their Bcl-2 expressions and Bcl-2/Bax ratio. It could be partly prevented by adoptive transfer of CD4⁺CD25⁺ T reg cells from normal to infected pregnant mice.59

The high level of Th1 cytokines (IL2 and IFNγ) were reported to be produced by CD4⁺ cells upon interaction with tachyzoite.50 It is found that MyD88 is effectively involved in the development of Th1 response.60 Generally, the Th1 cell-mediated immunity builds the resistance against the T. gondii infection by IFNγ production via Th1 effector cells.61 The role of Th1 cytokines (IFNγ, IL12, and TNFα) for susceptibility to toxoplasmosis has been witnessed with the absence of any of these pro-inflammatory mediators as previously studied.42,62 Moreover, other cytokines like IL2, IL6, IL7, IL15, IL18, and IL23 are also associated with the development of strong immunogenic response.60 Recent evidence demonstrates TLR-11–independent activation of inflammasome for driving CD4⁺ T-cell–derived IFNγ-mediated host resistance to T. gondii.7

GRA24-driven protective immunity mediated through p38 MAPK activation, IL12 production, and independent of MyD88 pathway has been evidenced, recently, through use of bicistronic IL12YFP reporter mice on MyD88+/+ and MyD88–/– genetic backgrounds, MyD88+/- and MyD88–/- bone marrow–derived macrophages as well as exploiting parasites species named as uracil auxotrophic Type-I stain of T. gondii cps1-1 and cps1-1:Δgra24.63

In contrast, various experiments have demonstrated that the cytokines involved in Th2 response also play a detrimental role for enhancing the susceptibility to T. gondii infection.66 The modulation of Th2 response is mainly carried out by IL4 and IL10. Both cytokines increase the host susceptibility to T. gondii in early infection.103 Nevertheless, the regulatory function of Th2 cytokines has been unveiled. The evidence from T. gondii infection (Type II stain) to the IL4-knockout mice resulted in less susceptibility to toxoplasmosis.65 Shoot-up levels of inflammatory cytokines were detected in IL10 knockout mice causing early resistance to Toxoplasma.64,66

C) Cytokines and other inflammatory mediators playing a role against T. gondii

Interferon-γ: Interferon-γ (IFNγ) is reckoned as the main pillar of cytokines induced by T cells (CD4⁺ and CD8⁺), γδT cells, and NK cells as protective immunity against either the acute or chronic phase of T. gondii infection.42,67,68 The neutralization of INFγ (anti-INFγ antibodies) in in vivo makes the mice (Swiss-Webster) susceptible to the primary infection (acute infection) and reactivates parasite (ME49 strain) in chronic infection.69 The IFNγ is produced from activated MΦs to exhibit its specific immunological functions against T. gondii (C56 strain).70 The latest evidence demonstrating the CD4⁺, CD8⁺, γδT cells, and NK cells as the main producers of IFNγ during toxoplasmosis was based on the use of a newly developed mouse line named as “GREVEN” an IFNγ reporter mouse having a fusion protein of Venus and
NanoLuc to analyze IFNγ producing cells. Likewise, the Lck-Cre/Ifngf1f1 mice were found highly susceptible to toxoplasmosis, further strengthening the role of T-cell-IFNγ in protection against toxoplasmosis. At the site of infection, the maintenance of IFR8+ inflammatory DCs is essential provision of host resistance against intracellular T. gondii, whereas this requires the production of IFNγ by ILC1 and NK cells through T-bet involvement. The extensive role of IFNγ was partly dependent on the release of TNFα by activated Mφs. In the host, most documented immunological pathway against T. gondii (virulent RH strain) is nitric oxide–dependent. It involves the collective role of IFNγ and TNFα for the production of nitric oxide that curtails the development of microbial pathogens. IFNγ is involved in an alternative mechanism via inducing Ag-specific CD8+ (CTL) cells-dependent immunity. INFγ directs the antimicrobial response such as STAT-1, TNFα, IL1β, and CD40L by activating the transcription factor (NF-κB) signaling pathway. This signaling pathway has been believed to rely mainly through involvement of inducible nitric oxide synthase (iNOS) and immunity related GTPase (IRGs) which make grounds to resist T. gondii (ME49) infection, the nitric oxide-independent intracellular resistance mechanism has been evidenced recently through use of IFNγ-stimulated bone marrow–derived macrophages (BMDM). Likewise, recent evidence has shown IFNγ as an inducing agent of guanylate binding proteins (GBP) that accumulate on surface of intracellular stage of T. gondii; thus, mediating the role of an intracellular check on the excessive growth of this parasite that may become lethal to mouse host in the absence of GBP as has been demonstrated by the early death of the mice deficient in murine guanylate binding protein-7 (mGBP7), by T. gondii infection. The intracellular pathogen (T. gondii) has been highly adapted for combating host immune response by interfering with NF-κB signaling pathway. It also has the ability to inhibit IFNγ signaling by curtailing the function of STAT-1 and by augmenting the levels of IFNγ signaling suppressor molecules, named as suppressor of cytokine signaling molecule-1 (SOCS1). Most of the Th2 cytokines function antagonistically to IFNγ.

**Tumor necrosis factor-α (TNFα):** TNFα is a pyrogenic factor, produced by Mφs, T lymphocytes and basophils, found to be responsible for the production of acute inflammatory response. It has the ability for microbialic activity in Mφs via the production of IFNγ from NK cells. TNFα functions synergistically with IFNγ for the development of resistance against T. gondii (C56 strain) infection. Hence, it is suggested to have a crucial role in the protective immunity against toxoplasmosis. However, certain researchers explained the role of TNFα to be doubtful. TNFα has also been reported to elicit cerebral and hepatic autoimmunity. It has also been found to assist in intra-cerebral dissemination of T. gondii (ts-4 strain and virulent RH) in mice ((TNF(−/−), L'Talphα(−/−)), and TNF/L'Talphα(−/−)).

**Interleukin-1:** It is an acute phase response mediator cytokine that plays a synergistic role with TNFα enhancing inflammation during infection with T. gondii. TNFα has been found to be associated with IL1β in regulating endothelial cells for the immunological and inflammatory role. In vitro studies elucidated an effect of these cytokines in hindering the intracellular multiplication of T. gondii in murine peritoneal Mφs or human fibroblast. However, in vivo studies on mice showed the protective role of recombinate TNFα and/or recombinate IL1β during infection with tachyzoites of T. gondii (C56 strain and RH strain). In female mouse models (BALB/c, and Swiss-Webster), IFNγ induced the anti-toxoplasmic activity by augmenting the production of TNFα when treated with recombinate cytokines (TNFα and IL1β).

**Interleukin-2:** Interleukin-2 is exclusively produced by CD4+ T cells. It is initially regarded as the primary T cell growth factor having significant role in the proliferation as well as in the development of antigen stimulated CD8+ T cells. Various studies carried out on viral infections revealed an indispensable role of IL2 during primary response of T cells via effector cytotoxic T cell development and restoration of CD8+ memory T cells. IL2 has also been found to induce T cells proliferation for IFNγ production during the infection with T. gondii (virgin). Whereas, the role of IL2 in secondary immune response to T. gondii (ME49) was disclosed by Sa et al., (2013) on CD8+ T cell hybridoma clones from the spleens of chronically infected female mice (BALB/c, BALB/c-background Rag1−/−, and Swiss-Webster). They found an increased production of INFγ with exogenous input of IL2 to CD8+ T cell hybridomas. Various studies on murine models proved the protective role of IL2 against the infection with T. gondii. Increased survival rates and reduced number of cysts in the brains of mice were observed when treated with recombinate IL2. It also triggered the lytic activity of Mφs and NK cells.

**Interleukin-4:** Interleukin-4 is a Th type 2 cytokine that down-regulates the effect of Th type 1 cytokines. The progressive toxoplastic encephalitis found to be linked with the presence of mRNA transcripts in the brains of infected mice (C57BL/10 ScSn). In acute infection of T. gondii, IL4 exerted a protective role by antagonizing the products of Th1 cells that reduced the number of mortalities. However, the prolonged exposure of IL4 made the mice (IL-4R/2) susceptible to chronic toxoplasmosis with increased multiplication of parasite cysts in brain.

**Interleukin-6:** Mainly, it is involved in early development of acute phase response, maintenance of hematopoiesis and immune barriers in ocular as well as cerebral Toxoplasmosis. It makes the NK cells to increment their cytotoxic activity and is also involved in the maturation of
antibody secreting B lymphocytes and the differentiation of T lymphocytes. Different immune cells such as MΦs, endothelial cells, monocytes, myelomatous, and fibroblasts are involved in the production of IL6. This cytokine functions synergistically with IL1β and TNFα. Hence, it is regarded as a remarkable pyrogenic factor mediating dominantly the production of hepatocyte based acute inflammatory proteins. The signaling pathway of IL6 is associated with gp130 signal transducing component that leads toward the activation of STAT1 and STAT3 via the signaling of JAK1, JAK2, and TYk2. This pathway is referred as gp130-mediated JAK/STAT signaling pathway which is most effectively regulated by socs-3 signaling. The host transcription factor STAT3, antagonizes host response and strengthens parasite survival which is activated by the cytokines (IL6 and IL10) during infection with certain intracellular pathogens. Interestingly, T. gondii (ROP16-deficient type I) has been found to activate this factor by phosphorylation caused by rhoptry kinase. This event is accompanied with impaired production of IL12p40 and TNFα response. To investigate the mechanism involved in the invasion of immune strategies, it was found that socs-3 up-regulates IL6 and IL10 which stimulate activation of STAT-3. The role of IL6 is more critical as in normal defense mechanism, socs-3 can curtail function of this cytokine by formation of gp130/IL6 receptor complex. In the work conducted by Whitmarsh et al., (2011), the mice deletion with socs-3 in MΦs and neutrophils unexpectedly resulted in increased susceptibility to toxoplasmosis by reducing the levels of IL12 and IFN-γ. This study suggested more pronounced anti-inflammatory role of IL6 in MΦs particularly in the absence of socs-3.

Figure 1. Outcome of toxoplasmosis in case of resistance and susceptibility.
Moreover, the gp130 transducing component has common structural features to IL6 and IL27.116 This IL27 is critically involved in curtailing the infection induced inflammatory pathology and functions antagonistically to IL6. The mice lacking gp130 signal transducer (gp130 Y757F mice) upon infection with *T. gondii* (ME49), showed high parasite burdens and increased mortality having low IL12 and IFNγ titer.117 IL6 is a cytokine that more pronouncedly functions to resist the *Toxoplasma*-induced encephalitis in murine model.118 Previously, the protective role of IL6 was questioned in various studies that illustrated the role of IL6 in increased intracellular multiplication of *T. gondii*.110 Later studies defined the critical role of IL6 in progression of *T. gondii* infection as the IL6-deficient mice rapidly switched to severe states of toxoplasmosis such as *Toxoplasma* encephalitis119 (Figure 2).

**Interleukin-7:** It plays a crucial role for the development of memory CD8+ T cells120 which are the key producers of INF-γ in acquired immunity.41 The IL2, IL7, and IL15 fall in the family of γ-chain cytokines that are involved in building CD8+ memory T cell.121,122 The development of memory cells in the form of CD8+ T cells has been disclosed to be dependent on the significant role of IL7 as well as IL15. The *in vivo* neutralization of IL15 affected CD8+ memory T cells productions which are more susceptible to *T. gondii* (76K) re-infection.123 In memory cells development process, IL7 is recruited to provide the survival signals to naïve and memory CD8+ T cells.124 In a study conducted by Bhadra et al., (2010), the synergistic role of IL7 and IL15 was explored. Their findings indicated a severe impairment of memory cells in case of the absence of both of these cytokines. However, the absence of any of these cytokines resulted in minimal impact on the maturation of splenic CD8+ T cells.125

**Interleukin-10:** It induces pleiotropic effect on cells and is found to be a suppressive cytokine that is purely an anti-inflammatory in its action.126 Immune cells including Mφs, CD4+, B cells, DCs, and mastocytes are the source of this cytokine.127–129 The rapid production of IL10 by localized Mφs was reported upon infection with high doses of virulent (RH) strain in mice.130 However, it keeps a check on the protective functioning of CD4+ T cells in acute phase infection.130,132 IL10 is also reported to be responsible for suppressive microbicidal activity of Mφs and neutrophils by minimizing the function of nitric oxide (NO) synthase enzyme, IFNγ and IL12. The NO inhibits the production of O2 free radicals and prostaglandins.133–135 The simultaneous production of these antagonizing cytokines may lead to demolish the defensive mechanism. Hence, it was reported that the production of IL10 is supported by an “activation signal” after the release of IL12. The IL12 lets the production of IFNγ by Th1 cells not only triggering the effector cells to play protective role but also passing a signal for the reactivation of IL10 gene expression. The study conducted by Gaddi et al.,136 (2007) explained the negative feedback mechanism for the poised state of these pathogenic and regulatory cytokines via CD4+ T cell lineage. In the mice (BALB/c) susceptible to *T. gondii*, the increased levels of IL10 were found in their lymph nodes and central nervous system causing chronic toxoplasmic encephalitis.134 In a study, the detrimental role of IL10 was disclosed as to promote the development of intracellular tachyzoites by inhibiting Mφ L-arginine-based killing.133 However, making a comparison, severe combined immune-deficient (SCID) IL10 knockout *T. gondii* (RH strain) infected mice lived longer than the infected SCID mice.66

**Interleukin-12:** In innate immune response, certain cell populations (DCs, MΦ, and Np) are reported to produce IL12 in *vitro* against *T. gondii* (a virulent strain).130,137,138 It is the central inducer of IFNγ in developing the protective immunity against *T. gondii*. It is evident from various experiments that the complete absence of immunity in INFγ as well as IL12p40, IL12p35, and STAT4 deficient mice leads to deaths in early acute infection.139–141 The increased mortalities were reported when diphtheria toxin were used to deplete DCs in *T. gondii* (ME49 strain) infected organism which abolish the production of IL12.142 The absence of p40 chain in IL12 heterodimer is responsible for the poor production of IFNγ in mice (IL12-deficient).143 During chronic Toxoplasmosis, the 12/15 lipoxygenase (12/15-LOX) is involved in the oxidation of unsaturated fatty acids in Mφs which deprives the production of IL12. In addition, the 12/15-LOX deficient mice were enabled to produce comparable levels of IL12 when stimulated with lipopolysaccharide (LPS). This finding was explained by the involvement of neutrophils and particularly DCs in inducing IL12 production in acute phase infection.144

**Interleukin-15:** It belongs to Th1 immune response and plays an effective role in enhancing the function and development of CD8+ T cells.145 IL15 serves to play a key role in the development of various lymphocyte population more importantly NK, CD8+ T cells and intraepithelial lymphocytes (IELs).146 The memory (CD8+) T cell response was found inadequate identified in IL15 knockout mice.147,148 However, a study conducted by Lieberman et al., (2004) reported no role of IL15 for development of memory immune response. They found the mice deficient with this cytokine withstood the severe *T. gondii* (ME49) infection.149

**Interleukin-17:** The IL17A, IL17F, and IL22 are secreted from Th17 cells reported by Wu et al., (2018). But a variety of immune cells including CD8+, γδ, and NK cells also secrete IL17. It is an inflammatory cytokine that provides innate immunity from the recruitment of neutrophils which make the host resistant against *T. gondii* infection.152 A subset of CD4+ T cells has been identified which produces IL6, IL17A, IL17F, and TNF in response to IL23.157,158 In a study conducted by
Kelly et al., (2005), IL17 knockout mice remained successful in developing a normal acquired immunity against *T. gondii.* A study carried out on T and B lymphocytes deficient mice revealed that NK cells are the major IL17 producers. These cells are influenced to produce IL17 in the same manner as T cells are triggered. In addition, the researchers revealed a key role of IL6 to target NK cells for secretion of IL17. The IL17A neutralization by antibodies had a partial protective effect against fatal *T. gondii*-associated inflammation. Severe South American ocular toxoplasmosis is associated with decreased level of intraocular IFNγ and IL17A. But in ocular toxoplasmosis (with less severe clinical presentation and infected by non-virulent strains) in France, the IL17A level was augmented in toxoplasmic uveitis. Neutralizing IL17A decreased intraocular inflammation and parasite load in mice (Swiss-Webster). It is suggested that the local IL17A production by resident cells plays a central role in the pathology of ocular toxoplasmosis. Finally, there was observed a lower level of IL17 expressing CD4+ and CD8+ T lymphocytes in cells cultures from sero-negative and seropositive pregnant and non-pregnant women, respectively upon stimulation with tachyzoites. A recent study demonstrated an essential role of T cells expressing class I-restricted T cell-associated molecule (CRTAM), for IL17 production during toxoplasmosis. The study also highlights the importance of IL17 in regulating immunopathology whereby deficiency of IL17 can cause dysbiosis through the production of antimicrobial peptides as well as through translocating gut-bacterial flora to spleen and mesenteric lymph nodes. Overall, these results suggest that IL17-mediated responses may be useful for both protective and pathogenic effects.

**Interleukin-18:** It is a pleiotropic cytokine produced in a non-specific manner. IL18 is a potent cytokine involved in the production of IFNγ by NK cells and T lymphocytes. Hence, it is involved in building innate as well as acquired immune response against *T. gondii* infection. Attributable to its identical role in developing resistance, IL18 is referred as a potential enhancer of IL12 activity. Structurally, it is closely related to IL1β cytokine family. Moreover, similar to IL1β signaling pathway, IL18 precede the activation of NF-κB that requires STAT4 factor for its activation. The impaired role of IFNγ was observed in IL18 deficient mice with intracellular infection. Interestingly, NK cell has certain receptors for IL18 which has synergistic role with IL12 in developing innate immunity against *T. gondii* infection. However, endogenous role of IL18 on SCID mice was demonstrated to be trivial having less influence on IFNγ production when the infected mice were treated with anti-IL18. In contrary, the exogenous role of IL18 was reported to increment the production of IFNγ ultimately boosting the resistance against *T. gondii.* IL18 reported to be involved in the immunopathology of intestine in mice accompanied by IL12.

**Interleukin-33:** The host damage protein IL-33 has recently been shown to play an important role in the immune response by affecting the local environment of brain in the favour of both host and parasite survival through engagement of astrocytes via IL-33 Receptor. IL-33 is a host damage protein that is produced locally in the brain tissue from the oligodendrocytes and astrocytes. The evidence to this effect is very strong as it is based on the use of mice having IL-33 receptor-deficient astrocytes. Transforming Growth Factor-β: Transforming growth factor-β (TGFβ) is another immunosuppressant cytokine that plays a critical role in antagonizing the action of TNFα, TNFβ, IFNγ, and IL12. Moreover, it is considered to be involved in limiting the immune-pathologies incited by Th1 cytokines specifically in central nervous system (CNS) as well as in intestine. TGFβ has been reported to induce immunopathological effects on retinal cell line in an *in vitro* study by increasing the replication of *T. gondii.* The intraepithelial lymphocytes are the main producers of TGFβ that is involved in the down-regulation of pro-inflammatory cytokines (IFNγ, TNFα, IL1β, IL12, IL15, and IL18) in case of pathogenic lamina propria lymphocytes (LPL) response (hyper-Th1 response). When wild type mice were treated with active transfer of IELs, they showed no sign of ileitis. This study revealed the modulating role of IELs for LPL produced Th1 response in the intestine.

In spite of gastrointestinal sites, TGFβ also has a dominant role as an anti-inflammatory agent in brain and eyes. TGFβ signals the spleen cells to secrete anti-inflammatory cytokine such as IL10 that is synergistically involved in checking the pro-inflammatory secretions from NK cells and CD4+ T cells in these immune-privileged sites (eyes, brain, and placenta). The production of IL6 by innate immune response functions antagonistically to TGFβ and suspends its protective role for immune privileged sites (eyes and brain) susceptible to hyper-inflammation.

**CCL22:** *T. gondii* induces expression of CCL22, a chemokine that has been linked to Toxoplasma-induced activation of Wnt/beta-catenin signaling pathway, for resisting cellular check on parasite replication and favoring the parasite survival within Toxoplasma-exposed naïve BMDMs. This cellular-check (induced by IFNγ and independent of nitric oxide) significantly reduced intracellular parasite load when Wnt/beta-catenin signaling pathway was chemically antagonized using IWR-1-endo in naïve BMDMs, thus strengthening the hypothesis that Toxoplasma induces this signaling pathway to survive intracellular anti-parasitic immunity. This got further support when a significant reduction in the intracellular parasite load of RHASPS5, a mutant strain of *T. gondii* lacking ability to secrete dense proteins into the host cell, was seen in naïve BMDMs.
Additionally, T. gondii invading the BMDMs pre-stimulated with IFNγ, switched on its bradyzoite gene profile.27

2) Toxoplasmosis and pregnancy

Toxoplasmosis is more important in pregnant women and immune compromised patients with respect to abortion, hydrocephalus, and retinochoroiditis. In pregnancy, Th2 immune response becomes activated which favors the proliferation of Toxoplasma. Briefly, in acute phase of T. gondii infection, certain cytokines (TGFβ1, TNFα, IL4, IL5 IL7, IL10, and IL17A) and chemokines (CXC, C, and CX3C families) play an important role as protective immune response.191,192 These pro-inflammatory cytokines down-regulate anti-inflammatory cytokines which travel to immune-privileged sites (brain, eyes, and placenta) to favor the existence of corpus luteum in the presence of low progesterone and 17β estradiol in pregnant women.23 Apoptosis of placental cells may end up in fetal resorption, congenital anomalies (hydrocephalus and retinochoroiditis), or abortion191 (Figure 1). Briefly, toxoplasmosis with lymphadenitis has been reported with higher levels of chemokines (CXCL8/IL8, CXCL9, and CXCL10) in pregnant women. Additionally, levels of VCAM1, CCL2, and CCL5 are lower in pregnant than in non-pregnant women.193 The levels of ICAM1, CXCL9, CXCL10, MCSF, and TNFβ were up-regulated in acutely toxoplasmosis-infected Colombian pregnant women. Whereas, the levels of Eotaxin (Et), TGFβ, TNFα, IFNγ, IL2, IL4, IL15, CXCL1, and stem cell factor (SCF) were down-regulated in pregnant American acute cohorts.194 In congenital toxoplasmosis, it was found that serum levels of IFNγ and IL5 were greatly increased during active stage of retinochoroiditis. In contrast, IL10 production was low during inflammatory stage and significantly higher in patients with inactive lesions.195 The cytokine profile of acute toxoplasmosis-infected patients varies with geographical localities.

4) Perspectives for immunomodulation, therapy, vaccine, and other anti-parasitic challenges

The exploration of deep knowledge on the role of cytokines in toxoplasmosis should open new avenues for therapeutic measures based on immunomodulation. For instance, the use of IL17A antagonist inhibited the ocular toxoplasmosis in European patients.196 Similarly, inhibition of parasite kinases in South American toxoplasmosis patients enhances the expression of IFNγ197-199 This difference in inhibition sites might be strain dependent. Recent analysis of the cytokines profile in congenital toxoplasmosis200 indicates that modulation of cytokines through immuno-modulatory peptides could be assayed as immune adjuvants.201 Such approaches need to be explored for the control of toxoplasmosis in humans. Etanercept (a soluble TNF-receptor fusion protein), widely used to treat autoimmune disease, activates the conversion of bradyzoites (chronic toxoplasmosis) to tachyzoites (acute toxoplasmosis) through down-regulation of pro-inflammatory cytokines (TNF, IL-1beta, and IL6).202 It would be interesting to try to achieve a sterile immunity in an experimental model of chronic toxoplasmosis, at first transforming bradyzoites to tachyzoites through use of Etanercept but not too long after this, treating the tachyzoites to eliminate the parasite from the host body.

A recently identified drug target for T. gondii is an endonuclease named as cleavage and polyadenylation specificity factor subunit-3 (CPSF3) that has a role in mRNA processing in eukaryotes. This has been demonstrated by strong in vitro anti-parasitic activity by use of benzoxaborole (AN3661), a drug molecule that targets wild-type CPSF3. The parasites that were found resistant to this drug molecule displayed mutations in the TgCPSF3. Recapitulation of the similar resistant phenotype of the parasite through generation of mutations in the wild-type CPSF3 while exploiting CRISPR/Cas9, further strengthened the importance of this new therapeutic target against T. gondii.203

One of the most exciting areas of research is to explore the means and effects of intervention strategies on how various strains of T. gondii can modulate host’s transcriptome204 and non-coding RNAs including microRNA and long non-coding RNA.205 Similarly, exploring how T. gondii exploits exosomes in modulating host immune response206 as well as how therapeutic interventions designed for heme-deficient conditions affect infection outcome,207 remains interesting areas of research.
Given the assumed fact that around one third population of world is harboring Toxoplasma in chronic form, that is, tissue cysts, why not to plan a vaccine to eliminate the tissue cysts from human population and other hosts seropositive to this infection, with a vaccine (based on GRA6Nt or other similar antigens) that should be capable of eliminating tissue cysts.

**Conclusions**

Different factors are responsible for the pathogenesis of Toxoplasmosis and the survival of host. These factors include versatile genetic makeup of different strains of T. gondii, complicated immunological background of hosts, biochemical interaction among certain cytokines, invasion...
strategies of parasite as well as the immunogenicity of antigens encountered with host’s immune cells. The type of cytokines production depends on the strain of *Toxoplasma*. The IL10 and TGFβ1 production were higher in type I strain and lower in type II and III strain of toxoplasmosis. The production of IL12 was higher upon exposure of pathogen to DCs, MΦ, Np, NK cells, and T cells which is essential for the release of IFNγ. The production of IL12 switches the NK cells for release of IFNγ which develops resistance against *T. gondii* infection in host. Impairment in the production of IL12 may lead to demolish IFNγ resulting to develop host sensitivity for *T. gondii* infection. Moreover, IL6 also has critical role for gp130 signaling pathway for the up- and down-regulation of SOCS-gene which is responsible for the susceptibility and resistance of toxoplasmosis (Figure 2). The basic switching of pro- and anti-inflammatory cytokines in acute and chronic phases of toxoplasmosis is direly required for understanding the development of disease. Such cytokines are involved in the development of resistance and susceptibility of *Toxoplasma* in host. Agonist and antagonist effect of host cytokines network leads to the chronic condition of disease. *In vivo* up- and down-regulation of desired cytokines (IFNγ, IL6, IL12, and SOCS-3) could be helpful to boost up the immune response of host for the control of toxoplasmosis. Moreover, the synergistic and antagonistic relations among cytokines need to be comprehended on molecular and biochemical basis. The most compelling results are related with a Th2-deviated response associated to virulent strains in South American patients. Type II strain has the ability to translocate NF-κB in the nucleus of mouse splenocytes and bone marrow–derived MΦ. It is the reason that *Toxoplasma* type I strain survives from host immune response rather than type II and III but the complete defeat of host’s immune response is not in the favor of parasite’s survival in the ecosystem. The survival of the host after entry of *T. gondii*, is essential for ensuring existence of both the host and the parasite as if parasite defeats the host’s immune response, it not only marks the death of the host but also of the parasite as parasite needs a viable host to ensure its own survival as well as for its transmission to next generations of the same host as well as to other host species. The current review findings state that *in vitro* harvesting of IL12 from DCs, Np and MΦ upon exposure with *T. gondii* might be a source for therapeutic use in toxoplasmosis. Current review suggests that therapeutic interventions leading to up-regulation/supplementation of SOCS-3, IL12, and IFNγ to the infected host could be a solution to sterile immunity against *T. gondii* infection. This would be of interest particularly in patients passing through immunosuppression owing to any reason like the ones receiving anti-cancer therapy, the ones undergoing immunosuppressive therapy for graft/transplantation, the ones suffering from immunodeficiency virus (HIV) or having AIDS.

Acknowledgements

N. Cardona and M. Murillo, Centro de Investigaciones Biomedicas, Universidad del Quindio, Avenida Bolivar 12N, Armenia (Q), Colombia, South America helped to add some important studies in the manuscript. Mr. Adeel Mumtaz Abbasi (MPhil Student) is acknowledged for drawing of figures of the manuscript.

Declaration of Conflicting Interests

The author(s) declared no potential conflicts of interest with respect to the research, authorship, and/or publication of this article.

Funding

The author(s) received no financial support for the research, authorship, and/or publication of this article.

ORCID iDs

Imran Rashid https://orcid.org/0000-0002-0280-9335
Haroon Akbar https://orcid.org/0000-0002-8294-5862

References

1. Chen XQ, Zhou CX, Elsheikha HM, et al. (2017) Profiling of the perturbed metabolic state of mouse spleen during acute and chronic toxoplasmosis. *Parasites Vectors* 10(1): 339.
2. White MW, Radke JR and Radke JB (2014) Toxoplasma development - turn the switch on or off? *Cell Microbiol* 16(4): 466–472.
3. Fisch D, Clough B and Frickel E-M (2019) Human immunity to Toxoplasma gondii. *PLoS Pathogens* 15(12): e1008097.
4. Costantini S, Castello G and Colonna G (2010) Human Cytokine: a new challenge for systems biology. *Bioinformation* 5(4): 166–167.
5. Sarciron M and Gherardi A (2000) Cytokines involved in Toxoplasmosis encephalitis. *Scand J Immunol* 52(6): 534–543.
6. Akbar H, Germon S, Berthon P, et al. (2012) Depletion of CD25+ cells during acute toxoplasmosis does not significantly increase mortality in Swiss OF1 mice. *Memórias do Instituto Oswaldo Cruz* 107(2): 155–162.
7. López-Yglesias AH, Camanzo E, Martin AT, et al. (2019) TLR11-independent inflammasome activation is critical for CD4+ T cell-derived IFN-γ production and host resistance to Toxoplasma gondii. *PLoS Pathogens* 15(6): e1007872.
8. Sullivan WJ Jr. and Jeffers V (2012) Mechanisms of Toxoplasma gondii persistence and latency. *FEMS Microbiol Rev* 36(3): 717–733.
9. Vidal JE, Hernandez AV, de Oliveira AC, et al. (2005) Cerebral toxoplasmosis in HIV-positive patients in Brazil: clinical features and predictors of treatment response in the HAART era. *AIDS Patient Care STDS* 19(10): 626–634.
10. Schmitt N and Ueno H (2015) Regulation of human helper T cell subset differentiation by cytokines. *Curr Opin Immunol* 34: 130–136.
11. Saeij JP, Boyle JP and Boothroyd JC (2005) Differences among the three major strains of Toxoplasma gondii and their specific interactions with the infected host. Trends Parasitol 21(10): 476–481.

12. Murillo-León M, Müller UB, Zimmermann I, et al. (2019) Molecular mechanism for the control of virulent Toxoplasma gondii infections in wild-derived mice. Nat Commun 10(1): 1–15.

13. Witola WH, Kim CY and Zhang X (2013) Inherent oxidative stress in the Lewis rat is associated with resistance to toxoplasmosis. Infect Immun 85(10): e00289–17.

14. Hassan MA, Olijnik A-A, Frickel E-M and Saeij JP (2019) Clonal and atypical Toxoplasma strain differences in virulence vary with mouse sub-species. Int J Parasitol 49(1): 63–70.

15. Cervantes PW, Martorelli Di Genova B, Erazo Flores BJ, et al. (2021) RIPK3 facilitates host resistance to oral Toxoplasma gondii infection. Infect Immun 89(5): e00021–21.

16. Rashid I, Moiré N, Héraut B, et al. (2017) Enhancement of the protective efficacy of a ROP18 vaccine against chronic toxoplasmosis by nasal route. Medical microbiology and immunology 206(1): 53–62.

17. Hu L-Y, Zhang N-Z, Zhang F-K, et al. (2017) Resistance to Chronic Toxoplasma gondii infection induced by a DNA vaccine expressing GRA16. BioMed Research International 2017: 1295038.

18. Kim CY, Zhang X and Witola WH (2018) Small GTPase immunity-associated proteins mediate resistance to Toxoplasma gondii infection in Lewis rat. Infect Immun 86(4): e00582-e00517.

19. Butcher BA, Kim L, Johnson PF, et al. (2001) Toxoplasma gondii tachyzoites inhibit proinflammatory cytokine induction in infected macrophages by preventing nuclear translocation of the transcription factor NF-kappa B. J Immunol 167(4): 2193–2201.

20. Butcher BA, Kim L, Panopoulos AD, et al. (2005) IL-10-independent STAT3 activation by Toxoplasma gondii mediates suppression of IL-12 and TNF-alpha in host macrophages. J Immunol 174(6): 3148–3152.

21. Angeloni MB, Guirelli PM, Franco PS, et al. (2013) Differential apoptosis in BeWo cells after infection with highly (RH) or moderately (ME49) virulent strains of Toxoplasma gondii is related to the cytokine profile secreted, the death receptor Fas expression and phosphorylated ERK1/2 expression. Placenta 34(11): 973–982.

22. Butcher BA, Kim L, Johnson PF, et al. (2001) Toxoplasma gondii tachyzoites inhibit proinflammatory cytokine induction in infected macrophages by preventing nuclear translocation of the transcription factor NF-κB. J Immunol 167(4): 2193–2201.

23. Iqbal J and Al-Awadhi M (2016) Toxoplasmosis role of cytokines in disease modulation & tissue pathology. Ann Clin Pathol 4(7): 1090.

24. Iqbal J, Al-Awadhi MA and Raghupathy RG (2016) TGF-beta1 levels and intraocular tissue alterations in mice infected with a virulent type I RH Toxoplasma gondii strain. Exp Parasitol 162: 57–63.

25. Sauer A, Pfaff AW, Villard O, et al. (2012) Interleukin 17A as an effective target for anti-inflammatory and antiparasitic treatment of toxoplasmic uveitis. J Infect Dis 206(8): 1319–1329.

26. Dobbin CA, Smith NC and Johnson AM (2002) Heat shock protein 70 is a potential virulence factor in murine toxoplasmosis infection via immunomodulation of host NF-kappa B and nitric oxide. J Immunol 169(2): 958–965.

27. Robben PM, Mordue DG, Truscott SM, et al. (2004) Production of IL-12 by macrophages infected with Toxoplasma gondii depends on the parasite genotype. J Immunol 172(6): 3686–3694.

28. Schade B and Fischer HG (2001) Toxoplasma gondii induction of interleukin-12 is associated with acute virulence in mice and depends on the host genotype. Vet Parasitol 100(1–2): 63–74.

29. Zhao Y, Reyes J, Rovira-Diaz E, et al. (2021) Cutting Edge: CD36 Mediates Phagocyte Tropism and Avirulence of Toxoplasma gondii. J Immunol 207(6): 1507–1512.

30. Fox BA, Guuevara RB, Rommereim LM, et al. (2019) Toxoplasma gondii parasitophorous vacuole membrane-associated dense granule proteins orchestrate chronic infection and GRA12 underpins resistance to host gamma interferon. mBio 10(4): e00589–19.

31. Sayles PC and Johnson LL (1996) Exacerbation of toxoplasmosis in neutrophil-depleted mice. Nat Immun 15(5): 249–258.

32. Hunter CA (1996) How are NK cell responses regulated during infection? Exp Parasitol 84(3): 444–448.

33. Denkers EY, Butcher BA, Del Rio L, et al. (2004) Neutrophils, dendritic cells and Toxoplasma. Int J Parasitol 34(3): 411–421.

34. Denkers EY, Bzik DJ, Fox BA, et al. (2012) An inside job: hacking into Janus kinase/signal transducer and activator of transcription signaling cascades by the intracellular protozoan Toxoplasma gondii. Infect Immun 80(2): 476–482.

35. Steffens N, Beuter-Gunia C, Kravets E, et al. (2020) Essential role of mGBP7 for survival of Toxoplasma gondii infection. mBio 11(1): e02993–19.

36. Goldszmid RS, Bafica A, Jankovic D, et al. (2007) TAP-1 indirectly regulates CD4+ T cell priming in Toxoplasma gondii infection by controlling NK cell IFN-gamma production. J Exp Med 204(11): 2591–2602.

37. Scanga CA, Aliberti J, Jankovic D, et al. (2002) Cutting edge: MyD88 is required for resistance to Toxoplasma gondii infection and regulates parasite-induced IL-12 production by dendritic cells. J Immunol 168(12): 5997–6001.

38. Hou B, Benson A, Kuzmich L, et al. (2011) Critical coordination of innate immune defense against Toxoplasma
gondii by dendritic cells responding via their Toll-like receptors. *Proc Natl Acad Sci U S A* 108(1): 278–283.
39. Frenkel JK (1988) Pathophysiology of toxoplasmosis. *Parasitol Today* 4(10): 273–278.
40. Gomez Marin JE, Pinon JM, Bonhomme A, et al. (1997) Does human toxoplasmosis involve an imbalance in T1/T2 cytokines? *Med Hypotheses* 48(2): 161–169.
41. Gazzinelli R, Xu Y, Hiency S, et al. (1992) Simultaneous depletion of CD4+ and CD8+ T lymphocytes is required to reactivate chronic infection with Toxoplasma gondii. *J Immunol* 149(1): 175–180.
42. Denkers EY and Gazzinelli RT (1998) Regulation and function of T-cell-mediated immunity during Toxoplasma gondii infection. *Clin Microbiol Rev* 11(4): 569–588.
43. Denkers EY, Gazzinelli RT, Martin D, et al. (1993) Emergence of NK1.1+ cells as effectors of IFN-gamma dependent immunity to Toxoplasma gondii in MHC class I-deficient mice. *J Exp Med* 178(5): 1465–1472.
44. Suzuki Y, Sa Q, Gehman M, et al. (2011) Interferon-gamma and perforin-mediated immune responses for resistance against Toxoplasma gondii in the brain. *Exp Rev Mol Med* 13: e31.
45. Grumont RJ and Gerondakis S (1990) The murine c-rel proto-oncogene encodes two mRNAs the expression of which is modulated by lymphoid stimuli. *Oncogene Research* 5(4): 245–254.
46. Mason NJ, Artis D and Hunter CA (2004) New lessons from old pathogens: what parasitic infections have taught us about the role of nuclear factor-kappaB in the regulation of immunity. *Immunol Rev* 201: 48–56.
47. Dzierszinski F, Pepper M, Stumhofer JS, et al. (2007) Presentation of Toxoplasma gondii antigens via the endogenous major histocompatibility complex class I pathway in nonprofessional and professional antigen-presenting cells. *Infect Immun* 75(11): 5200–5209.
48. Jordan KA, Dupont CD, Tait ED, et al. (2010) Role of the NF-kappaB transcription factor c-Rel in the generation of CD8+ T-cell responses to Toxoplasma gondii. *Int Immunol* 22(11): 851–861.
49. Sa Q, Ochiai E, Tiwari A, et al. (2017) Determination of a key antigen for immunological intervention to target the latent stage of Toxoplasma gondii. *J Immunol* 198(11): 4425–4434.
50. Gazzinelli RT, Denkers EY and Sher A (1993) Host resistance to Toxoplasma gondii: model for studying the selective induction of cell-mediated immunity by intracellular parasites. *Infect Agents Dis* 2(3): 139–149.
51. Shirahata T, Yamashita T, Ohta C, et al. (1994) CD8+ T lymphocytes are the major cell population involved in the early gamma interferon response and resistance to acute primary Toxoplasma gondii infection in mice. *Microbiol Immunol* 38(10): 789–796.
52. Ohkura N, Kitagawa Y and Sakaguchi S (2013) Development and maintenance of regulatory T cells. *Immunity* 38(3): 414–423.
53. Akbar H, Dimier-Poisson I and Moiré N (2015) Role of CD4+ Foxp3+ regulatory T cells in protection induced by a live attenuated, replicating type I vaccine strain of Toxoplasma gondii. * Infect Immun* 83(9): 3601–3611.
54. Oldenhove G, Bouladoux N, Wohlfert EA, et al. (2009) Decrease of Foxp3+ Treg cell number and acquisition of effector cell phenotype during lethal infection. *Immunity* 31(5): 772–786.
55. Benson A, Murray S, Divakar P, et al. (2012) Microbial infection-induced expansion of effector T cells overcomes the suppressive effects of regulatory T cells via an IL-2 deprivation mechanism. *J Immunol* 188(2): 800–810.
56. Olguin JE, Fernandez J, Salinas N, et al. (2015) Adoptive transfer of CD4+(+)Foxp3(+) regulatory T cells to C57BL/6J mice during acute infection with Toxoplasma gondii down modulates the exacerbated Th1 immune response. *Microb Infect* 17(8): 586–595.
57. Salinas N, Olguin JE, Castellanos C, et al. (2014) T cell suppression in vitro during Toxoplasma gondii infection is the result of IL-2 competition between Tregs and T cells leading to death of proliferating T cells. *Scand J Immunol* 79(1): 1–11.
58. Ge YY, Zhang L, Zhang G, et al. (2008) In pregnant mice, the infection of Toxoplasma gondii causes the decrease of CD4+CD25+ -regulatory T cells. *Parasite Immunol* 30(9): 471–481.
59. Chen JL, Ge YY, Zhang J, et al. (2013) The dysfunction of CD4+(+)CD25(+) regulatory T cells contributes to the abortion of mice caused by Toxoplasma gondii excreted-secreted antigens in early pregnancy. *PLoS One* 8(7): e69012.
60. Chen M, Aosai F, Norose K, et al. (2002) Involvement of MyD88 in host defense and the down-regulation of anti-heat shock protein 70 autoantibody formation by MyD88 in Toxoplasma gondii-infected mice. *J Parasitol* 88(5): 1017–1019.
61. Hunter CA, Villarino A, Artis D, et al. (2004) The role of IL-27 in the development of T-cell responses during parasitic infections. *Immunol Rev* 202: 106–114.
62. Alexander J and Hunter CA (1998) Immunoregulation during toxoplasmosis. *Chem Immunol* 70: 81–102.
63. Mercer HL, Snyder LM, Doherty CM, et al. (2020) Toxoplasma gondii dense granule protein GRA24 drives MyD88-independent p38 MAPK activation, IL-12 production and induction of protective immunity. *PLoS Pathog* 16(5): e1008572.
64. Gazzinelli RT, Wysocka M, Hieny S, et al. (1996) the absence of endogenous IL-10, mice acutely infected with Toxoplasma gondii succumb to a lethal immune response dependent on CD4+ T cells and accompanied by overproduction of IL-12, IFN-gamma and TNF-alpha. *J Immunol* 157(2): 798–805.
66. Neyer LE, Grunig G, Fort M, et al. (1997) Role of interleukin-10 in regulation of T-cell-dependent and T-cell-independent mechanisms of resistance to Toxoplasma gondii. *Infect Immun* 65(5): 1675–1682.

67. Nishiyama S, Pradip A, Ma JS, et al. (2020) T cell-derived interferon-γ is required for host defense to Toxoplasma gondii. *Parasitol Int* 75: 102049.

68. López-Yglesias AH, Burger E, Camanzo E, et al. (2021) T-cell-depended interferon-γ mediates DC1-dependent host resistance against Toxoplasma gondii. *PLoS Pathog* 17(1): e1008299.

69. Suzuki Y, Conley FK and Remington JS (1989) Importance of endogenous IFN-gamma for prevention of toxoplasmonic encephalitis in mice. *J Immunol* 143(6): 2045–2050.

70. McCabe RE, Luft BJ and Remington JS (1984) Effect of murine interferon gamma on murine toxoplasmosis. *J Infect Dis* 150(6): 961–962.

71. Chang HR, Grau GE and Pechere JC (1990) Role of TNF and IL-1 in infections with Toxoplasma gondii. *Immunology* 69(1): 33–37.

72. Green SJ, Nacy CA and Meltzer MS (1991) Cytokine-induced synthesis of nitrogen oxides in macrophages: a protective host response to Leishmania and other intracellular pathogens. *J Leukoc Biol* 50(1): 93–103.

73. Sharma DP, Ramsay AJ, Maguire DJ, et al. (1996) Interleukin-4 mediates down regulation of antiviral cytokine expression and cytotoxic T-lymphocyte responses and exacerbates vaccinia virus infection in vivo. *J Virol* 70(10): 7103–7107.

74. Andrade RM, Portillo JA, Wessendorp M, et al. (2005) CD40 signaling in macrophages induces activity against an intracellular pathogen independently of gamma interferon and reactive nitrogen intermediates. *Infect Immun* 73(5): 3115–3123.

75. Lieberman LA, Banica M, Reiner SL, et al. (2004) STAT1 plays a critical role in the regulation of antimicrobial effector mechanisms, but not in the development of Th1-type responses during toxoplasmosis. *J Immunol* 172(1): 457–463.

76. Martens, Parvanova I, Zerrahn J, et al. (2005) Disruption of Toxoplasma gondii parasitophorous vacuoles by the mouse p47-resistance GTPases. *PLoS Pathogens* 1(3): e24.

77. Gossner A and Hassan MA (2020) Transcriptional analyses identify genes that modulate bovine macrophage response to Toxoplasma infection and immune stimulation. *Frontiers in Cellular and Infection Microbiology* 10: 437.

78. Butcher BA, Greene RI, Henry SC, et al. (2005) p47 GTPases regulate Toxoplasma gondii survival in activated macrophages. *Infect Immun* 73(6): 3278–3286.

79. Shapiro S, Harb OS, Caamano J, et al. (2004) The NF-kappaB signaling pathway: immune evasion and immunoregulation during toxoplasmosis. *Int J Parasitol* 34(3): 393–400.

80. Zimmermann S, Murray PJ, Heeg K, et al. (2006) Induction of suppressor of cytokine signaling-1 by Toxoplasma gondii contributes to immune evasion in macrophages by blocking IFN-gamma signaling. *J Immunol* 176(3): 1840–1847.
Filisetti D and Candolfi E (2004) Immune response to Toxoplasma gondii. *Ann Ist Super Sanita* 40(1): 71–80.

O’Shea JJ, Gadina M and Schreiber RD (2002) Cytokine signaling in 2002: new surprises in the Jak/Stat pathway. *Cell* 109(Suppl): S121–S131.

Nicholson SE, De Souza D, Fabri LJ, et al. (2000)Suppressor of cytokine signaling-3 preferentially binds to the SHP-2-binding site on the shared cytokine receptor subunit gp130. *Proc Natl Acad Sci U S A* 97(12): 6493–6498.

Babon JJ, Sabo JK, Soetopo A, et al. (2008) The SOCS box domain of SOCS3: structure and interaction with the elonginBC-cullin5 ubiquitin ligase. *J Mol Biol* 381(4): 928–940.

Beaman MH, Hunter CA and Remington JS (1994). Enhancement of intracellular replication of Toxoplasma gondii by IL-6. Interactions with IFN-gamma and TNF-alpha. *J Immunol*, 153(10): 4583–4587.

Bermudez LE, Wu M, Petrofsky M, et al. (1992) Interleukin-6 antagonizes tumor necrosis factor-mediated mycobacteriostatic and mycobactericidal activities in macrophages. * Infect Immun* 60(10): 4245–4252.

Murray PJ (2007) The JAK-STAT signaling pathway: input and output integration. *J Immunol* 178(5): 2623–2629.

Yamamoto M, Standley DM, Takashima S, et al. (2009) A single polymorphic amino acid on Toxoplasma gondii kinase ROP16 determines the direct and strain-specific activation of Stat3. *J Exp Med* 206(12): 2747–2760.

Kubo M, Hanada T and Yoshimura A (2003) Suppressors of cytokine signaling and immunity. *Nat Immunol* 4(12): 1169–1176.

Whitmarsh RJ, Gray CM, Gregg B, et al. (2011) A critical role for SOCS3 in innate resistance to Toxoplasma gondii. *Cell Host & Microbe* 10(3): 224–236.

Taga T and Kishimoto T (1997) Gp130 and the interleukin-6 family of cytokines. *Annu Rev Immunol* 15: 797–819.

Silver JS, Stumhofer JS, Passos S, et al. (2011) , 187, pp. 350–360. IL-6 mediates the susceptibility of glycoprotein 130 hypermorphs to Toxoplasma gondii.*J Immunol*.

Deckert-Schluter M, Albrecht S, Hof H, et al. (1995) Dynamics of the intracerebral and splenic cytokine mRNA production in Toxoplasma gondii- resistant and -susceptible congenic strains of mice. *Immunology* 85(3): 408–418.

Suzuki Y, Rani S, Liesenfeld O, et al. (1997) Impaired resistance to the development of toxoplasmic encephalitis in interleukin-6-deficient mice. *Infect Immun* 65(6): 2339–2345.

Kasper LH, Matsuura T and Khan IA (1995). IL-7 stimulates protective immunity in mice against the intracellular pathogen, Toxoplasma gondii. *J Immunol*, 155(10): 4798–4804.

Ma A, Koka R and Burkett P (2006) Diverse functions of IL-2, IL-15, and IL-7 in lymphoid homeostasis. *Annu Rev Immunol* 24: 657–679.

Schluns KS, Williams K, Ma A, et al. (2002). Cutting edge: requirement for IL-15 in the generation of primary and memory antigen-specific CD8 T cells. *J Immunol*, 168(10): 4827–4831.

Combe CL, Moretto MM, Schwartzman JD, et al. (2006) Lack of IL-15 results in the suboptimal priming of CD4+ T cell response against an intracellular parasite. *Proc Natl Acad Sci U S A* 103(17): 6635–6640.

Prlc M, Lefrancois L and Jameson SC (2002) Multiple choices: regulation of memory CD8 T cell generation and homeostasis by interleukin (IL)-7 and IL-15. *J Exp Med* 195(12): F49–F52.
125. Bhadra R, Guan H and Khan IA (2010) Absence of both IL-7 and IL-15 severely impairs the development of CD8 T cell response against Toxoplasma gondii. *PLoS One* 5(5): e10842.

126. Adib-Conquy M, Petit AF, Marie C, et al. (1999) Paradoxical priming effects of IL-10 on cytokine production. *Int Immunol* 11(5): 689–698.

127. Mosmann TR (1994) Properties and functions of interleukin-10. *Adv Immunol* 56: 1–26.

128. Moore KW, de Waal Malefyt R, Coffman RL, et al. (2001) Interleukin-12 in the absence of endogenous IFN-gamma, mice develop uncontrolled acute Toxoplasma gondii infection. *J Immunol* 165(2): 628–631.

129. Levings MK, Bacchetta R, Schulz U, et al. (2002) The role of IL-10 and TGF-beta in the differentiation and effector function of T regulatory cells. *Int Arch Allergy Immunol* 128(4): 263–276.

130. Bliss SK, Butcher BA and Denkers EY (2000) Rapid recruitment of neutrophils containing prestored IL-12 during microbial infection. *Journal of immunology (Baltimore, Md)* 165(8): 4515–4521.

131. de Waal Malefyt R, Yssel H, Roncarolo MG, et al. (1992) Interleukin-10 and the interleukin-10 receptor. *Annu Rev Immunol* 19: 683–765.

132. Rennick D, Berg D and Holland G (1992) The role of IL-10 and TGF-beta in the differentiation and effector function of T regulatory cells. *Int Arch Allergy Immunol* 129(4): 263–276.

133. Gazzinelli RT, Oswald IP, James SL, et al. (1992) IL-10 inhibits parasite killing and nitrogen oxide production by IFN-gamma-activated macrophages. *Journal of immunology (Baltimore, Md)* 148(6): 1792–1796.

134. Hunter CA, Litton MJ, Remington JS, et al. (1994) Immunocytochemical detection of cytokines in the lymph nodes and brains of mice resistant or susceptible to toxoplasmosic encephalitis. *J Infect Dis* 170(4): 939–945.

135. Schluter D, Kaefer N, Hof H, et al. (1997) Expression pattern and cellular origin of cytokines in the normal and Toxoplasma gondii-infected murine brain. *Am J Pathol* 150(3): 1021–1035.

136. Gaddi PJ and Yap GS (2007) Cytokine regulation of immunopathology in toxoplasmosis. *Immunol Cell Biol* 85(2): 155–159.

137. Gazzinelli RT, Wysocka M, Hayashi S, et al. (1994) Parasite-induced IL-12 stimulates early IFN-gamma synthesis and resistance during acute infection with Toxoplasma gondii. *J Immunol* 153(6): 2533–2543.

138. Bliss SK, Zhang Y and Denkers EY (1999) Murine Neutrophil Stimulation by Toxoplasma gondii Antigen Drives High Level Production of IFN-gamma-Independent IL-12. *J Immunol* 163(4): 2081–2088.

139. Scharton-Kersten TM, Wynn TA, Denkers EY, et al. (1996) The absence of endogenous IFN-gamma, mice develop unimpaired IL-12 responses to Toxoplasma gondii while failing to control acute infection. *J Immunol* 157(9): 4045–4054.

140. Yap G, Pesin M and Sher A (2000) Cutting Edge: IL-12 Is Required for the Maintenance of IFN-gamma Production in T Cells Mediating Chronic Resistance to the Intracellular Pathogen, Toxoplasma gondii. *J Immunol* 165(2): 628–631.

141. Lieberman LA, Cardillo F, Owyang AM, et al. (2004). IL-23 provides a limited mechanism of resistance to acute toxoplasmosis in the absence of IL-12. *J Immunol*, 173(3): 1887–1893.

142. Liu CH, Fan YT, Dias A, et al. (2006) Cutting edge: dendritic cells are essential for in vivo IL-12 production and development of resistance against Toxoplasma gondii infection in mice. *J Immunol* 177(1): 31–35.

143. Magram J, Connaughton SE, Warrier RR, et al. (1996) IL-12-deficient mice are defective in IFN gamma production and type 1 cytokine responses. *Immunity* 4(5): 471–481.

144. Middleton MK, Rubinstein T and Pure E (2006) Cellular and molecular mechanisms of the selective regulation of IL-12 production by 12/15-lipoxygenase. *J Immunol* 176(1): 265–274.

145. Khan IA and Kasper LH (1996) IL-15 augments CD8+ T cell-mediated immunity against Toxoplasma gondii infection in mice. *J Immunol* 157(5): 2103–2108.

146. Fehniger TA and Caligiuri MA (2001) Interleukin 15: biology and relevance to human disease. *Blood* 97(1): 14–32.

147. Lodolce JP, Boone DL, Chai S, et al. (1998) IL-15 receptor maintains lymphoid homeostasis by supporting lymphocyte homing and proliferation. *Immunity* 9(5): 669–676.

148. Kennedy MK, Glaccum M, Brown SN, et al. (2000) Reversible defects in natural killer and memory CD8 T cell lineages in interleukin 15-deficient mice. *J Exp Med* 191(5): 771–780.

149. Lieberman LA, Villegas EN and Hunter CA (2004) Interleukin-15-deficient mice develop protective immunity to Toxoplasma gondii. *Infect Immun* 72(11): 6729–6732.

150. Intlekofer AM, Banerjee A, Takemoto N, et al. (2008) Anomalous type 17 response to viral infection by CD8+ T cells lacking T-bet and comesdermin. *Science* 321(5887): 408–411.

151. Tajima M, Wakita D, Noguchi D, et al. (2008) IL-15 augments CD8+ T cell-dependent spontaneous proliferation is required for the induction of colitogenic IL-17-producing CD8+ T cells. *J Exp Med* 205(5): 1019–1027.

152. Lockhart E, Green AM and Flynn JL (2006) IL-17 production is dominated by gammadelta T cells rather than CD4 T cells during Mycobacterium tuberculosis infection. *J Immunol* 176(1): 31–35.

153. Michel ML, Keller AC, Paget C, et al. (2007) Identification of an IL-17-producing NKT.I.1(neg) iNKT cell population involved in airway neutrophilia. *J Exp Med* 204(5): 995–1001.

154. Rachitskaya AV, Hansen AM, Horai R, et al. (2008). Cutting edge: NKT cells constitutively express IL-23 receptor and RORgammat and rapidly produce IL-17 upon receptor ligation in an IL-6-independent fashion. *J Immunol*, 179(9): 5167–5171.

155. Xue ML, Thakur A and Willcox M (2002) Macrophage inflammatory protein-2 and vascular endothelial growth factor regulate corneal neovascularization induced by...
infection with Pseudomonas aeruginosa in mice. *Immunol Cell Biol* 80(4): 323–327.

156. Kelly MN, Kolls JK, Happel K, et al. (2005) Interleukin-17/interleukin-17 receptor-mediated signaling is important for generation of an optimal polymorphonuclear response against Toxoplasma gondii infection. *Infect Immun* 73(1): 617–621.

157. Aggarwal S, Ghilardi N, Xie MH, et al. (2003) Interleukin-23 promotes a distinct CD4 T cell activation state characterized by the production of interleukin-17. *J Biol Chem* 278(3): 1910–1914.

158. Langrish CL, Chen Y, Blumenschein WM, et al. (2006) Interleukin-12 (IL-12) and IL-18 synergistically induce the production of interferon-gamma by natural killer. *J Exp Med* 201(2): 233–240.

159. Amadi-Obi A, Yu CR, Liu X, et al. (2007) TH17 cells contribute to uveitis and scleritis and are expanded by IL-2 and inhibited by IL-27/STAT1. *Nat Med* 13(6): 711–718.

160. Stumhofer JS, Laurence A, Wilson EH, et al. (2006) Interleukin 27 negatively regulates the development of interleukin 17-producing T helper cells during chronic inflammation of the central nervous system. *Nat Immunol* 7(9): 937–945.

161. Passos ST, Silver JS, O’Hara AC, et al. (2010) IL-6 promotes NK cell production of IL-17 during toxoplasmosis. *Journal of immunology (Baltimore, Md)* 184(4): 1776–1783.

162. Guiton R, Vasseur V, Charron S, et al. (2010) Interleukin 17 receptor signaling is deleterious during Toxoplasma gondii infection in susceptible BL6 mice. *J Infect Dis* 202(3): 427–435.

163. de-la-Torre A, Sauer A, Pfaff AW, et al. (2013) Severe South American ocular toxoplasmosis is associated with decreased Ifn-gamma/IL-17a and increased IL-6/IL-13 intraocular levels. *PLoS Neglected Tropical Diseases* 7(11): e2541.

164. Sauer A, Villard O, Creuzot-Garcher C, et al. (2015) Intracellular levels of interleukin 17A (IL-17A) and IL-10 as respective determinant markers of toxoplasmosis and viral uveitis. *Clin Vaccine Immunol : CVI* 22(1): 72–78.

165. Silva JL, Rezende-Oliveira K, da Silva MV, et al. (2014) IL-17-expressing CD4(+) and CD8(+) T lymphocytes in human toxoplasmosis. *Mediat Inflamm* 2014: 573825.

166. Cervantes-Barragan L, Cortez VS, Wang Q, et al. (2019) CRTAM protects against intestinal dysbiosis during pathogenic parasitic infection by enabling Th17 maturation. *Front Immunol* 10: 1423.

167. Hunter CA, Timans J, Pisacane P, et al. (1997) Comparison of the effects of interleukin-1 alpha, interleukin-1 beta and interferon-gamma-inducing factor on the production of interferon-gamma by natural killer. *Eur J Immunol* 27(11): 2787–2792.

168. Kim YM, Talanian RV, Li J, et al. (1998) Nitric oxide prevents IL-1beta and IFN-gamma-inducing factor (IL-18) release from macrophages by inhibiting caspase-1 (IL-1beta-converting enzyme). *Journal of immunology (Baltimore, Md)* 161(8): 4122–4128.

169. Bazan JF, Timans JC and Kastelein RA (1996) A newly defined interleukin-1? *Nature* 379(6566): 591.

170. Matsumoto S, Tsuji-Takayama K, Aizawa Y, et al. (1997) Interleukin-18 activates NF-kappaB in murine T helper type 1 cells. *Biochem Biophys Res Commun* 234(2): 454–457.

171. Thomassen E, Bird TA, Renshaw BR, et al. (1998) Binding of interleukin-18 to the interleukin-1 receptor homologous receptor IL-1Rlp1 leads to activation of signaling pathways similar to those used by interleukin-1. *J Interferon Cytokine Res* 18(12): 1077–1088.

172. Cai G, Radzanowski T, Villegas EN, et al. (2000), 165, pp. 2619–2627. Identification of STAT4-dependent and independent mechanisms of resistance to Toxoplasma gondii. *J Immunol* 162(10): 5894–5901.

173. Wei XQ, Leung BP, Niedbala W, et al. (1999) Altered immune responses and susceptibility to Leishmania major and Staphylococcus aureus infection in IL-18-deficient mice. *Journal of immunology (Baltimore, Md)* 163(5): 2821–2828.

174. Hyodo Y, Matsui K, Hayashi N, et al. (1999) IL-18 up-regulates perforin-mediated NK activity without increasing perforin messenger RNA expression by binding to constitutively expressed IL-18 receptor. *Journal of immunology (Baltimore, Md)* 162(3): 1662–1668.

175. Walker W, Aste-Amezaga M, Kastelein RA, et al. (1999). IL-18 and CD28 use distinct molecular mechanisms to enhance NK cell production of IL-12-induced IFN-gamma. *J Immunol*, 162(10): 5894–5901.

176. Zhang T, Kawakami K, Qureshi MH, et al. (1997) Interleukin-12 (IL-12) and IL-18 synergistically induce the fungicidal activity of murine peritoneal exudate cells against Cryptococcus neoformans through production of gamma interferon by natural killer cells. *Infect Immun* 65(9): 3594–3599.

177. Vossenkamper A, Struck D, Alvarado-Esquivel C, et al. (2004) Both IL-12 and IL-18 contribute to small intestinal parasite control. *Infect Immun* 71(8): 4122–4128.

178. Still KM, Batista SJ, O’Brien CA, et al. (2020) Astrocytes promote a protective immune response to brain Toxoplasma gondii infection via IL-33-ST2 signaling. *PLoS Pathogens* 16(10): e1009027.

179. Hunter CA, Bermudez L, Beemink H, et al. (1995) Transforming growth factor-beta inhibits interleukin-12-induced production of interferon-gamma by natural killer cells: a role for transforming growth factor-beta in the regulation of T
cell-independent resistance to Toxoplasma gondii. Eur J Immunol 25(4): 994–1000.

183. Langermans JA, Nibbering PH, Van Vuren-Van Der Hulst ME, et al. (2001) Transforming growth factor-beta suppresses interferon-gamma-induced toxoplasmatastic activity in murine macrophages by inhibition of tumour necrosis factor-alpha production. Parasite Immunol 23(4): 169–175.

184. Buzoni-Gatel D, Debbabi H, Mennechet FJ, et al. (2001) Murine ileitis after intracellular parasite infection is controlled by TGF-beta-producing intraepithelial lymphocytes. Gastroenterology 120(4): 914–924.

185. Schluter D, Bertsch D, Frei K, et al. (1998) Interferon-gamma antagonizes transforming growth factor-beta expression in human retinal pigment epithelial cells is enhanced by Toxoplasma gondii: a possible role in the immunopathogenesis of retinochoroiditis. Clin Exp Immunol 128(2): 372–378.

186. Nagineni CN, Detrick B and Hooks JJ (2002) Transforming growth factor-beta1, beta2, and beta3 antagonizes interleukin-10 and interferon-gamma in human retinal pigment epithelial cells. Invest Ophthalmol Vis Sci 43(9): 2668–2673.

187. Mennechet FJ, Kasper LH, Rachinel N, et al. (2004) Intestinal intraepithelial lymphocytes prevent pathogen-driven inflammation and regulate the Smad/T-bet pathway of lamina propria CD4+ T cells. Eur J Immunol 34(4): 1059–1067.

188. Okamoto S, Hara Y and Streilein JW (1995) Induction of anterior chamber-associated immune deviation with lymphoreticular allogeneic cells. Transplantation 59(3): 377–381.

189. Lu F, Huang S and Kasper LH (2003) Interleukin-10 and interferon-gamma antagonizes transforming growth factor-beta in human retinal pigment epithelial cells. Invest Ophthalmol Vis Sci 44(10): 2591–2596.

190. Ohta K, Yamagami S, Taylor AW, et al. (2000) IL-6 antagonizes TGF-beta and abolishes immune privilege in eyes with endotoxin-induced uveitis. Invest Ophthalmol Vis Sci 41(9): 2591–2596.

191. Rezende-Oliveira K, Silva N, Mineo J, et al. (2012) Cytokines and Chemokines production by mononuclear cells from parturient women after stimulation with live Toxoplasma gondii. Placenta 33(9): 682–687.

192. Zlotnik A and Yoshie O (2000) Chemokines: a new classification system and their role in immunity. Immunity 12(2): 121–127.

193. Pomares C, Holmes TH, Estran R, et al. (2017) Cytokine profiles in patients with toxoplastic lymphadenitis in the setting of pregnancy. Cytokine 90: 14–20.

194. Pernas L, Ramirez R, Holmes TH, et al. (2014) Immune profiling of pregnant Toxoplasma-infected US and Colombia patients reveals surprising impacts of infection on peripheral blood cytokines. J Infect Dis 210(6): 923–931.

195. de Araujo TE, Coelho-Dos-Reis JG, Bela SR, et al. (2017) Early serum biomarker networks in infants with distinct retinochoroidal lesion status of congenital toxoplasmosis. Cytokine 95: 102–112.

196. Lahmar I, Abou-Bacar A, Abdelrahman T, et al. (2009) Cytokine profiles in toxoplastic and viral uveitis. J Infect Dis 199(8): 1239–1249.

197. Alvarez C, de-la-Torre A, Vargas M, et al. (2015) Striking Divergence in Toxoplasma ROP16 Nucleotide Sequences From Human and Meat Samples. J Infect Dis 211(12): 2006–2013.

198. de-la-Torre A, Pfaff AW, Grigg ME, et al. (2014) Ocular cytokinome is linked to clinical characteristics in ocular toxoplasmosis. Cytokine 68(1): 23–31.

199. Menees KL, Haskins BE and Denkers EY (2019) Impact of etanercept in mice chronically infected with Toxoplasma gondii. Front Immunol 10: 2822.

200. Camero AC, Machado AS, Bela SR, et al. (2016) Cytokine signatures associated with early onset, active lesions and late cicatricial events of retinochoroidal commitment in infants with congenital toxoplasmosis. J Infect Dis 213(12): 1962–1970.

201. Gomez Marin JE (2016) Possibilities for immunomodulation in congenital toxoplasmosis. J Infect Dis 214(4): 656.

202. Yang J, Wang L, Xu D, et al. (2018) Risk assessment of etanercept in mice chronically infected with Toxoplasma gondii. Front Microbiol 9: 2822.

203. Palencia A, Bougdour A, Brenier-Pinchart M-P, et al. (2017) Targeting Toxoplasma gondii CPSF3 as a new approach to control toxoplasmosis. EMBO Mol Med 9(3): 385–394.

204. Garfoot AL, Cervantes PW, Knoll LJ, et al. (2019) Transcriptional Analysis Shows a Robust Host Response to Toxoplasma gondii during Early and Late Chronic Infection in Both Male and Female Mice. Infect Immun 87(5): e00024–19.

205. Menard KL, Haskins BE and Denkers EY (2019) Impact of Toxoplasma gondii Infection on Host Non-coding RNA Responses. Frontiers in Cellular and Infection Microbiology 9(132).

206. Varikuti S, Jha BK, Holcomb EA, et al. (2020) The role of vascular endothelium and exosomes in human protozoan parasitic diseases. Vessel Plus 4.

207. Kloehn J, Harding CR and Soldati-Favre D (2021) Supply and demand-heme synthesis, salvage and utilization by Apicomplexa. FEBS J 288(2): 382–404.

Appendix

Abbreviation

| Acronym | Description |
|---------|-------------|
| Ag | Antigen |
| APC | Antigen-presenting cells |
| BMDM | bone marrow-derived macrophages |
| CNS | Central nervous system |
| CPSF3 | Cleavage and polyadenylation specificity factor subunit-3 |
| DCs | Dendritic cells |
| ESA | Excretory–secretory antigens |
| Abbreviation | Description |
|--------------|-------------|
| GBPs         | Guanylate Binding Proteins |
| GIMAPs       | small GTPase immunity-associated proteins |
| HIV          | Human immune deficiency virus |
| IELs         | Intraepithelial lymphocyte |
| iNOS         | Inducible nitric oxide synthase |
| LPL          | Lamina propria lymphocytes |
| IRGs         | Immunity-related GTPases |
| LPS          | Lipopolysaccharide |
| MyD88        | Myeloid differentiation domain-88 |
| MΦ           | Macrophages |
| NK           | Natural killer |
| NP           | Neutrophils |
| RIPK3        | receptor-interacting serine/threonine-protein kinase 3 |
| SCF          | Stem cell factor |
| SCID         | Severe combined immune-deficient |
| SOCs1        | Suppressor of cytokine signaling molecule-1 |
| T reg        | Regulatory T |
| T. gondii    | Toxoplasma gondii |
| TGFβ         | Transforming growth factor-β |
| Th2          | T helper cell 2 |
| Th17         | T helper cell 17 |
| TLR          | Toll-like receptors |
| ZBP1         | Z-DNA binding protein-1 |