Loss of TLR3 aggravates CHIKV replication and pathology due to an altered virus-specific neutralizing antibody response

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Abstract

RNA-sensing toll-like receptors (TLRs) mediate innate immunity and regulate anti-viral response. We show here that TLR3 regulates host immunity and the loss of TLR3 aggravates pathology in Chikungunya virus (CHIKV) infection. Susceptibility to CHIKV infection is markedly increased in human and mouse fibroblasts with defective TLR3 signaling. Up to 100-fold increase in CHIKV load was observed in Tlr3−/− mice, alongside increased virus dissemination and pro-inflammatory myeloid cells infiltration. Infection in bone marrow chimeric mice showed that TLR3-expressing hematopoietic cells are required for effective CHIKV clearance. CHIKV-specific antibodies from Tlr3−/− mice exhibited significantly lower in vitro neutralization capacity, due to altered virus-neutralizing epitope specificity. Finally, SNP genotyping analysis of CHIKF patients on TLR3 identified SNP rs6552950 to be associated with disease severity and CHIKV-specific neutralizing antibody response. These results demonstrate a key role for TLR3-mediated antibody response to CHIKV infection, virus replication and pathology, providing a basis for future development of immunotherapeutics in vaccine development.

Keywords Chikungunya virus; innate immunity; joint inflammation; neutralizing antibodies; TLR3

Subject Categories Immunology; Microbiology; Virology & Host Pathogen Interaction

Introduction

Innate immunity against RNA viruses involves pattern recognition receptors (PRRs) that recognize structurally conserved molecules from diverse pathogens known as pathogen-associated molecular patterns (PAMPs) (Arpaia & Barton, 2011). PRRs include TLRs (particularly TLR3, TLR7 and TLR8) and members of the cytosolic retinoic acid-inducible gene I (RIG-I)-like receptors such as melanoma differentiation-associated protein 5 (MDA5) and RIG-I that detect RNA viruses through their genomic RNA or the double-stranded RNA (dsRNA) viral intermediate generated during replication (Yoneyama et al, 2004, 2005; Gitlin et al, 2006). Activation of these PRRs induces downstream anti-viral type I IFN response, which can also occur independently of viral RNA transcription and replication (Nikonov et al, 2013).

The involvement of TLRs in counteracting RNA virus infection is widely documented (Arpaia & Barton, 2011; Neighbours et al, 2012; Zhang et al, 2013). TLR3 recognizes dsRNA and can influence disease outcomes depending on the type of virus and infection model. TLR3-mediated innate and inflammatory responses were demonstrated to be protective against HIV, CMV and Dengue virus...
infections, while TLR3 stimulation results in detrimental disease outcomes in Influenza A virus and Punta Toro virus infections (Tabeta et al., 2004; Goffic et al., 2006; Gowen et al., 2006; Suh et al., 2007; Nasirudeen et al., 2011). TLR3-dependent response has been shown to be both protective by restricting virus replication in neurons (Daffis et al., 2008) and also detrimental in West Nile virus infection by perturbing TNFR1 signaling to promote virus entry into the brains of mice resulting in lethal encephalitis (Wang et al., 2004). Clinically, patients with impaired TLR3-mediated responses show an increased susceptibility to HSV-1 encephalitis (Zhang et al., 2007; Pérez de Diego et al., 2010; Sancho-Shimizu et al., 2011). Repeated reactivation of HSV-2 that led to the development of Mollaret meningitis has also been reported in an individual with TLR3 deficiency (Willmann et al., 2010).

The significance of TLR-mediated signaling and how TLR molecules influence clinically important re-emerging viruses such as CHIKV remains confounding. CHIKV is an ‘Old World’ alphavirus with a positive sense RNA genome belonging to the Togaviridae family (Deller & Russell, 1968). CHIKV is the causative agent for CHIKF, and over the last decade, it has caused simultaneous outbreaks of unprecedented scale in the Indian Ocean Islands (Josseran et al., 2006), India (Kaur et al., 2008) and subsequently in South East Asia (Laras et al., 2005; AbuBakar et al., 2007; Leo et al., 2009) and Europe (Queyriaux et al., 2008). Serious CHIKF outbreaks have also occurred in Cambodia (Centers for Disease Control & Prevention, 2012; Duong et al., 2012), Laos (Soulaphy et al., 2013) and Sierra Leone (Ansumana et al., 2013). Since 2013, it has finally reached the Americas and triggered ongoing outbreaks in the French West Indies (Enserrink, 2014; Leparc-Goffart et al., 2014). The clinical presentation of the disease is characterized by flu-like symptoms such as fever, rash and muscle aches which subside in 7–10 days (Kam et al., 2009). The hallmark of CHIKV infection is the incapacitating arthralgia that routinely persists for weeks or months after resolution of the acute symptoms and has high costs in terms of both quality of life and healthcare provision/economic loss (Borherini et al., 2008).

The involvement of TLRs in CHIKV replication was previously investigated in murine models (Schilte et al., 2010, 2012), but their role in CHIKV pathology and dissemination was not well established. We show here that TLR3 signaling plays a critical role in the control of CHIKV infection, replication, dissemination and pathology. Complementing an earlier report where CHIKV-infected Trif−/− (Toll/IL-1 resistance domain-containing adaptor inducing IFNβ; an adaptor protein essential for TLR3-mediated signaling) mice showed more pronounced viremia and joint inflammation compared to WT mice (Rudd et al., 2012), this study further demonstrated that infection of cultured primary human TRIF−/− and mouse Tr3−/− fibroblasts resulted in a significant enhancement of virus replication. Notably, infected Tr3−/− mice developed higher viremia and more pronounced joint inflammation, associated with a massive infiltration of myeloid cells such as neutrophils and macrophages when compared to WT mice. Furthermore, monitoring of virus infection using a firefly luciferase (FLuc)-tagged recombinant CHIKV (FLuc-CHIKV) revealed increased CHIKV dissemination in Tr3−/− mice. By infecting bone marrow chimera mice, we showed that TLR3-expressing hematopoietic cells were required for effective CHIKV clearance, but did not directly regulate CHIKV-induced joint inflammation. Mechanistic investigations further demonstrated that TLR3 was required by hematopoietic cells to direct CHIKV-specific antibody response toward important neutralizing linear B-cell epitopes in the E2 glycoprotein. In the absence of TLR3, high levels of CHIKV-specific IgG were still generated, but with substantially diminished neutralizing capacity. The clinical relevance of TLR3 was further investigated in CHIKV-infected patients, where the level of TR3 transcripts was increased in PBMCs of CHIKV-infected patients. Interestingly, SNP genotyping analysis further identified TLR3 SNPs rs3775292 and rs6552950, whose functional effects remain unknown, to be associated with prevalence of CHIKV phenotypes, and in the case of SNP rs6552950, also with disease severity, CHIKV-specific IgG response and antibody neutralizing capacity. Taken together, these results substantiate a role for TLR3 in the control of CHIKV replication, immunity and pathology.

Results

TRIF deficiency increases CHIKV replication

Activation of various TLR signaling pathways including TLR2, TLR3 and TLR4 engage the TRIF adaptor protein to induce expression of downstream anti-viral and pro-inflammatory genes (Yamamoto et al., 2003). To demonstrate a functional role for TRIF signaling in anti-CHIKV response, human primary fibroblasts with homozygous TRIF nonsense mutation (Sancho-Shimizu et al., 2011) were infected with CHIKV in vitro. CHIKV replication was remarkably higher in the TRIF−/− fibroblasts with a 2-log difference when compared to healthy control (Fig 1A–C). This observation is in line with previous findings where TRIF−/− fibroblasts were shown to be more susceptible to infection with HSV-1 and vesicular stomatitis virus (Sancho-Shimizu et al., 2011) due to a defect in type I interferon induction. This suggested that TLRs could be involved in CHIKV infection. We focused our study on TLR3 because TLR3-mediated immunity and polymorphisms to viral infection in human have been demonstrated to regulate disease progression (Pérez de Diego et al., 2010; Reinert et al., 2012; Zhang et al., 2007b).

TLR3 inhibits CHIKV replication

To dissect the significance of TLR3-mediated anti-CHIKV response, primary fibroblasts were isolated from both WT and Tlr3−/− mice and infected with CHIKV ex vivo. The loss of TLR3 significantly increased susceptibility to CHIKV infection that led to marked virus replication when compared to WT fibroblasts (Fig 1D–F). To gain further insights on how TLR3 signaling mediates an anti-CHIKV state, transcriptional profiles of type I IFNs and related IFN-stimulated genes (ISGs) during infection were analyzed (Supplementary Fig S1). Results revealed that the induction of type I IFNs and related IFN-stimulated genes (ISGs) during infection were analyzed (Supplementary Fig S1). Results revealed that the induction of type I IFN response observed here is independent of TLR3.

Loss of TLR3 leads to more pronounced virus dissemination and CHIKV-induced pathology

The importance of TLR3 in CHIKV infection was next examined in vivo in both WT and Tlr3−/− mice via the joint footpad inoculation route (Gardner et al., 2010; Teng et al., 2012). Animals were...
monitored daily for survival, viremia and joint inflammation. Although the loss of TLR3 expression did not affect animal survival, significantly higher viremia was observed throughout the course of disease in Tlr3−/− animals (Fig 2A). Strikingly, Tlr3−/− mice exhibited a remarkable exacerbation of CHIKV-induced inflammation at the joint footpad (Fig 2B). Transcriptional analysis on joint footpad samples harvested at the peak of viremia revealed that the induction of type I IFNs was differentially induced in Tlr3−/− mice and remained detectable until 16 dpi (Fig 2D).

To further study the role of TLR3 on in vivo CHIKV replication and dissemination, we used a recombinant FLuc-CHIKV to infect both WT and Tlr3−/− mice and tracked the kinetics of CHIKV infection by live imaging for a duration of 20 dpi. As expected, the loss of TLR3 once again resulted in more pronounced joint inflammation and viremia (Supplementary Fig S3A–C). Bioluminescence signals indicative of virus replication distinctly reveal the differences in the route of CHIKV dissemination in WT and Tlr3−/− mice during the initial phase of infection (Fig 3C and Supplementary Videos S3 and S4). From the site of infection (i.e. right joint footpad), signals were next detected in the posterior regions (tail, lymph node and left footpad) before moving to the anterior regions in the lower abdomen and head (Supplementary Videos S1 and S2). Notably, CHIKV dissemination was more rapid and extensive in Tlr3−/− mice than in WT mice (Fig 2C and D and Supplementary Videos S1 and S2). High level of virus replication was detected from the whole body by the bioluminescence signals (Fig 2D) and also from the blood by viral RNA quantification (Supplementary Fig S3C). The Tlr3−/− mice exhibited delayed virus clearance throughout the disease. On the contrary, bioluminescence signals were mainly localized at the site of infection in WT mice (Fig 2C and Supplementary Video S1). Notably, bioluminescence signals peaked at 2–3 dpi in the head region of Tlr3−/− mice and remained detectable until 16 dpi (Fig 2D).

To determine whether the pattern of CHIKV dissemination was specific to the loss of TLR3 and not due to the site of virus inoculation, mice were subjected to s.c. inoculation of CHIKV at the dermis region of the ear prior to live imaging analysis (Supplementary Fig S4 and Supplementary Videos S3 and S4). Similar patterns of CHIKV dissemination were observed. Although the level of bioluminescence signals measured by the ear inoculation route was lower, signals detected from the whole body, head region and inoculated ear in Tlr3−/− mice still remained significantly higher than that of the WT mice (Supplementary Fig S4B and Supplementary Videos S3 and S4) and correlated with viremia (Supplementary Fig S3D).

To further evaluate whether a loss of TLR3 expression affects virus tropism in tissues and deeper organs, mice inoculated via the joint footpad route were sacrificed at 3 dpi and dissected in order to quantify bioluminescence signals from the various organs, skeletal body and skin (Fig 2E). Higher bioluminescence signals were
detected from the skeletal body, skin, head, lungs, liver, heart, spleen, kidneys and brain of Tlr3−/− mice (Fig 2E), confirming the role of TLR3 in controlling CHIKV infection and virus dissemination. Interestingly, bioluminescence signals were also detected in the brain in 40% of Tlr3−/− mice (Fig 2E), and positive bioluminescence signals were still detectable in the body, skin and head of Tlr3−/− mice at 6 dpi (Supplementary Fig S5).

Severe joint inflammation in Tlr3−/− mice is associated with a massive infiltration of myeloid cells

In an effort to assess tissue damage in the joints of CHIKV-infected mice, histological assessments revealed an intensified disease score in Tlr3−/− mice that correlated with more pronounced subcutaneous edema and infiltration of CD11b+ myeloid cells at 3 dpi.
(Supplementary Fig S6) and during the peak of disease severity at 6 dpi (Fig 3). The infiltrated myeloid cells consisted primarily of F4/80+ macrophages that were localized in the edema region, while the Ly6G+ neutrophils were localized distinctly in the skeletal muscles. In addition, cellular infiltrates in the joint of virus-infected mice were further analyzed during the peak of inflammation at 6 dpi by flow cytometry. Consistent with histological analysis (Fig 3), the loss of TLR3 resulted in a significant increase in infiltrating CD11b+ myeloid cells and, in particular, CD11b+Ly6G+ neutrophils in Tlr3−/− mice (Fig 4A and Supplementary Fig S7).

This observation was further complemented by transcriptional analysis of the CHIKV-infected joint footpad that revealed a significant induction of neutrophil-associated activation molecules and chemokines such as defensins Defb4, Defb14, myeloperoxidase (Mpo), Il-1b and Il-8r in Tlr3−/− mice at 6 dpi (Fig 4B).

Although both CD4+ and CD8+ T cells were previously demonstrated to infiltrate the joint footpad, only CD4+ T cells were responsible for the pathology (Teo et al, 2013). Therefore, it would
be compelling to assess the number of CD4^+ T cells in Tlr3^{-/-} mice. Interestingly, the number of infiltrating CD4^+ T cells in the joint footpad was significantly reduced by half (Fig 4A). Joint inflammation was significantly reduced both in CHIKV-infected WT and Tlr3^{-/-} mice depleted of CD4^+ T cells when compared to controls (Fig 4C), indicating that joint pathology is mediated by CD4^+ T cells in both groups. However, depleting CD4^+ T cells did not modify the viremia in both WT and Tlr3^{-/-} mice (Figure 4D), demonstrating that virus replication is independent of CD4^+ T cells. Furthermore, CD4^+ T-cell depletion resulted in a significant reduction in Ly6G^+ neutrophils infiltration at 6 dpi (Fig 4E), supporting earlier observations (Figure 3) that neutrophils are a vital mediator of footpad joint inflammation in Tlr3^{-/-} mice.

**TLR3-expressing hematopoietic cells are required for effective CHIKV clearance**

It is well established that hematopoietic and non-hematopoietic cells are engaged in the control of CHIKV infection by the innate immune system (Her et al., 2010; Schilte et al., 2010). To next assess whether
effective CHIKV clearance could be mediated directly by TLR3-expressing hematopoietic cells, bone marrow chimeric mice were generated by lethal irradiation of WT mice, followed by adoptive transfer of either WT (referred to as WT→WT) or Tlr3−/− (referred to as Tlr3−/−→WT) bone marrow cells. Upon successful bone marrow engraftment after 6 weeks, chimeric mice were infected with recombinant FLuc-CHIKV by footpad inoculation and monitored for joint inflammation, viremia and CHIKV dissemination. Strikingly, Tlr3−/−→WT chimeras suffered significantly higher viremia from 5 dpi onward and endured delayed virus clearance till 14 dpi (Fig 5A), but joint inflammation and bioluminescence signals detected in body and CHIKV-infected footpad were not significantly different when compared to WT→WT chimeras (Fig 5B and C). The similar degree of joint inflammation observed between these two groups further substantiate that the differences observed in joint inflammation, viremia and CHIKV dissemination.

FIGURE 5. Tlr3−/−→WT and Tlr3−/−→Tlr3−/− bone marrow chimeras were impaired in CHIKV clearance.

A-D WT and Tlr3−/− mice (n = 4–7 per group) were lethally irradiated and reconstituted with bone marrow cells from WT or Tlr3−/− mice. Chimeric mice were infected with CHIKV [10^5 PFU] by footpad inoculation after 6 wks. Viremia was determined by viral load quantification. Dotted line indicates the limit of viral load detection. Data are representative of two independent experiments and presented as mean ± SD (two-tailed Mann–Whitney U-test, **P = 0.0095 5–7 dpi viremia WT recipient chimeras, *P = 0.0139 8 dpi viremia WT recipient chimeras, **P = 0.0095 10 dpi viremia WT recipient chimeras, *P = 0.0350 4 dpi viremia Tlr3−/− recipient chimeras, **P = 0.0082 5 dpi viremia Tlr3−/− recipient chimeras, *P = 0.0221 6 dpi viremia Tlr3−/− recipient chimeras, **P = 0.0012 7 dpi viremia Tlr3−/− recipient chimeras, **P = 0.0082 8 dpi viremia Tlr3−/− recipient chimeras).

B, E Extent of joint inflammation was measured daily and expressed as disease score relative to day 0 (pre-infection). Data are representative of two independent experiments and presented as mean ± SD (two-tailed Mann–Whitney U-test, *P = 0.0117 4 dpi disease score Tlr3−/− recipient chimeras, **P = 0.0146 5 dpi disease score Tlr3−/− recipient chimeras).

C, F CHIKV dissemination is not modulated in WT and Tlr3−/− chimeras. Bioluminescence signals of whole body was quantified and expressed as average radiance (p/s/cm^2/sr). The lowest limit of detection is 0 p/s/cm^2/sr. Data are representative of one of two independent experiments and presented as mean ± SD (two-tailed Mann–Whitney U-test, *P = 0.0381 1 dpi whole-body WT recipient chimeras, **P = 0.0082 1 dpi whole-body Tlr3−/− recipient chimeras).

Strikingly, Tlr3−/−→WT chimeras suffered significantly higher viremia from 5 dpi onward and endured delayed virus clearance till 14 dpi (Fig 5A), but joint inflammation and bioluminescence signals detected in body and CHIKV-infected footpad were not significantly different when compared to WT→WT chimeras (Fig 5B and C). The similar degree of joint inflammation observed between these two groups further substantiate that the differences observed in joint inflammation, viremia and CHIKV dissemination.

Impaired virus clearance in Tlr3−/− mice is due to diminished recognition of CHIKV E2 glycoprotein

Chimera and T-cell depletion experiments suggested a role for TLR3 signaling on B cells in virus control. The role of B cells in mediating virus clearance was demonstrated with CHIKV-specific antibodies that map to epitopes within the CHIKV E1 and E2 glycoproteins (Kam et al, 2012b; Lum et al, 2013). These antibodies were demonstrated to exhibit high neutralizing activities in both macaque and murine models (Kam et al, 2012b, 2014; Lum et al, 2013). This led us to further examine whether the impaired virus clearance observed in Tlr3−/− mice (Fig 2A and Supplementary Fig S3C and D) was due to a defective CHIKV-specific antibody response. The levels of CHIKV-specific total IgM and IgG antibodies in sera of infected mice collected at 0, 3, 6, 9, 12 and 15 dpi were first determined using virion-based ELISA (Fig 6A and B). CHIKV-specific IgM antibodies were detected on 3 dpi and peaked at 6 dpi with no difference between the WT and Tlr3−/− mice (Fig 6A). CHIKV-specific IgGs were detected later, from 6 dpi, and their levels continued to
Figure 6. TLR3 modulates early CHIKV-specific IgG response in mice.
WT and Tlr3−/− mice (n = 6 per group) were infected with CHIKV (10^5 PFU) by joint footpad inoculation.
A, B Total CHIKV-specific IgM (A) and IgG (B) levels were determined from serum samples collected at 0, 3, 6, 9, 12 and 15 dpi at a dilution of 1:100 and 1:2,000, respectively, using purified CHIKV virion-based ELISA. Data are representative of two independent experiments and presented as mean ± SD (two-tailed Mann–Whitney U-test, **P < 0.0087 6 dpi [ig]).
C, D Neutralizing capacity of pooled sera collected at 6 dpi (C) and 15 dpi (D) from WT mice is significantly higher than from Tlr3−/− mice. Pooled sera were diluted 1:50–1:5,000 and mixed with CHIKV (MOI 10) for 1 h before infection of HEK 293T cells for 6 h. Assays were performed in quintuplicate, and data are expressed relative to virus-only-infected samples without sera. Dotted line indicates the detection limit of assay determined from mock-infected samples. Data are representative of two independent experiments and presented as mean ± SD (two-tailed Mann–Whitney U-test, **P < 0.0079 6 dpi serum dilution, **P = 0.0286 15 dpi serum dilution, *P = 0.0286 15 dpi 1,500 serum dilution, *P = 0.0286 15 dpi 5,000 serum dilution)
E, F Mapping antibody reactivity to linear B-cell epitopes within CHIKV E2 proteome. CHIKV E2 epitopes recognized at 6 dpi (E) and 15 dpi (F) were determined in pooled sera collected from infected mice using ELISA specific for overlapping 18-mer linear peptides spanning the CHIKV E2 proteome. The peptide numbers correspond to the position of the 18-mer linear peptides along the CHIKV E2 proteome. Structural data were retrieved from PDB (id: 3N44 and 2XF8) and visualized using the software CHIMERA (Pettersen et al, 2004). Assays were performed in triplicate and expressed as relative fold change after normalizing to OD_{non} from non-infected sera. Data are representative of two independent experiments and presented as mean ± SD (two-tailed unpaired t-test, *P = 0.0191 6 dpi epitope 381, **P = 0.0386 15 dpi epitope E2EP3, ***P = 0.0004 15 dpi epitope 381, ***P = 0.0001 15 dpi epitope E2EP3).
G Localization of identified CHIKV B-cell epitopes (381, blue; 388, green; E2EP3, red) within CHIKV proteome. Epitopes in the E2 glycoprotein were located based on the structural data obtained from PDB records 3N42. Number corresponds to the region of amino acid sequences in our overlapping 18-mer linear peptide library, along the CHIKV viral genome. All representations are shown in frontal and back view.

increase substantially up to 15 dpi (Fig 6B). These results indicated that the delayed virus clearance observed in Tlr3−/− mice was not due to the inability to produce CHIKV-specific antibodies. Instead, the level of CHIKV-specific total IgG antibodies was significantly higher in Tlr3−/− mice than from WT mice at 6 dpi (Fig 6B), but this did not translate to a higher CHIKV-neutralizing capacity as demonstrated by the in vitro neutralizing assay (Kam et al, 2012b) (Fig 6C). Intriguingly, serially diluted pooled sera taken at 6 dpi from Tlr3−/− mice significantly showed in vitro lower neutralizing activity against CHIKV compared to sera from WT mice (Fig 6C). Similar observations were obtained using pooled sera taken during complete virus clearance at 15 dpi from Tlr3−/− mice (Fig 6D) and Tlr3−/− → Tlr3+/− chimeras (Supplementary Fig S8A). Neutralizing activity was significantly reduced against CHIKV compared to sera from WT and WT→Tlr3+/− chimeras, respectively.

Studies using plasma samples from CHIKV-infected patients and sera from CHIKV-infected WT mice have revealed that most of the linear B-cell epitopes recognized by CHIKV-specific antibodies are localized in the E2 glycoprotein (Kam et al, 2012a,b; Lum et al, 2013). Based on these observations, linear B-cell epitopes were screened to assess whether the decreased neutralizing capacity of anti-CHIKV antibodies from Tlr3−/− mice was associated with diminished epitope recognition. Using the optimized peptide-based ELISA described (Kam et al, 2012b; Lum et al, 2013), linear peptides covering the E2 glycoprotein proteome (Supplementary Table S1) were screened with diluted sera (1:500) from mice collected at 6 and 15 dpi (Fig 6E and F and Supplementary Fig S8B). Linear peptides were screened individually and only three peptides were detected to exhibit significant differences toward the sera between WT and Tlr3−/− mice (Fig 6E and F). Recognition of these three peptides (381, 388 and E2EP3) by CHIKV-specific antibodies from Tlr3−/− mice was significantly diminished compared to WT mice (Fig 6E–G). Specifically, recognition of an early detection serological epitope ‘E2EP3’, which is a dominant linear B-cell epitope in CHIKV-infected patients and animal models (Kam et al, 2012a,b; Lum et al, 2013), was significantly diminished (Fig 6E–G). Moreover, sera from Tlr3−/− → Tlr3+/− chimeras were also less neutralizing and similarly associated with a reduced recognition of the same dominant linear E2 epitopes analyzed in Tlr3−/− mice (Supplementary Fig S8B). These observations strongly indicate that the impaired virus clearance in Tlr3−/− mice is related to the diminished recognition of dominant linear epitopes in the E2 glycoprotein that are functionally important for virus neutralization (Kam et al, 2012b).

TLR3 SNP rs6552950 is associated with disease severity and CHIKV-specific IgG neutralizing antibody capacity in CHIKV-infected patients

To examine the clinical relevance of TLR3-mediated immunity in CHIKV pathogenesis, transcriptional analysis was carried out in PBMCs of patients with CHIKV. TLR3 levels were significantly higher in CHIKV-infected patients when compared to healthy controls during the acute and early convalescent phases of CHIKV infection (Fig 7A). These results suggest that TLR3-mediated signaling forms part of the early innate immune response against CHIKV.

SNP genotyping analysis of 10 TLR3 tagSNPs was performed in 94 CHIKF patients to establish the impact of TLR3 polymorphisms on CHIKV-induced disease outcome. Association with CHIKV phenotypes revealed that SNP rs3775292 and SNP rs6552950 had significant nominal association to disease occurrence when compared to population controls from the 1000 genome project (Table 1) (1000 Genomes Project Consortium et al, 2010). In both SNPs, the minor alleles were associated with increased disease susceptibility as indicated by the odds ratios (OR) (Table 1). Stratified analysis of these TLR3 tagSNPs to disease severity revealed that a non-coding variant SNP rs6552950 had a nominal association with disease severity with an OR of 2.39 (P < 0.05) in severe CHIKF patients when compared to mild CHIKF patients (Table 1 and Fig 7B). In addition, anti-E2EP3 IgG response determined from 69 available patients’ plasma in this study cohort further revealed that patients with SNP rs6552950 genotype associated with severe CHIKF disease severity had significantly less neutralizing antibodies as compared to
patients with AA or AG genotypes (Fig 7D). Taken together, these results suggest that TLR3 polymorphism in humans could influence the neutralizing capacity of CHIKV-specific IgGs by modulating the recognition of CHIKV E2EP3 epitope important for virus neutralization. Specifically, SNP rs3775292 and SNP rs6552950 may be susceptibility factors for CHIKF, with SNP rs6552950 having a possible role in disease severity due to low anti-E2EP3 IgG response.

Discussion

The detection of PAMPs by PRRs to elicit an inflammatory response is an essential process of the host innate immune response against pathogens (Janeway & Medzhitov, 2002). TLR3-mediated immunity to natural infection in humans has been demonstrated to both limit and exacerbate viral disease progression. It has been shown that
Anti-viral IFN-dependent manner and restricts virus replication by inducing mediated through astrocytes which sense HSV-2 in a TLR3-defects in TLR3 signaling axis rendered both humans and mice permissive to HSV-1 encephalitis infection (Zhang et al., 2004; Daffis et al., 2008).

Although studies have emerged demonstrating the functions of RIG-I/MDA5-mediated signaling in anti-CHIKV host response (Schilte et al., 2010; White et al., 2011; Rudd et al., 2012; Olagnier Nile virus infection, TLR3 was either protective or deleterious to the host by limiting virus replication in neurons or promoting efficient virus entry into the brain, respectively, depending on the route of inoculation (Wang et al., 2004; Daffis et al., 2008).

Although studies have emerged demonstrating the functions of RIG-I/MDA5-mediated signaling in anti-CHIKV host response (Schilte et al., 2010; White et al., 2011; Rudd et al., 2012; Olagnier

Figure 7. TLR3 is highly induced in CHIKV-infected patients during the early disease phase and TLR3 SNP rs6552950 is associated with disease severity and specific IgG response.

A Transcriptional profiles of TLR3 in PBMCs of CHIKV-infected patients (n = 23) during acute (median 4 days post-illness onset), and early convalescent (median 10 days post-illness onset) disease. The level of TLR3 gene expression was expressed relative to healthy controls (n = 8) after normalization to GAPDH. Data are presented as mean ± SEM (two-tailed Mann–Whitney U-test, ***P = 0.0001 acute versus healthy controls, ***P = 0.0001 early convalescent versus healthy controls).

B TLR3 SNP rs6552950 is associated with disease severity in 94 CHIKV-infected patients. Histogram shows the percentage of patients in each genotype exhibiting either mild or severe disease phenotype. Severe illness is defined as patients having either a maximum temperature greater than 38.5°C, a maximum pulse rate greater than 100 beats/min or a nadir platelet count less than 100 × 10⁹/L. Mild illness is defined as patients who do not fulfill these criteria. Association between SNP genotype (AA, n = 43; AG, n = 24; GG, n = 11; unknown, n = 16) and disease phenotype (Mild—AA, n = 29; AG, n = 11; GG, n = 4; Severe—AA, n = 14; AG, n = 13; GG, n = 7) is performed using logistic regression analysis (Table 1). *P = 0.02

C TLR3 SNP rs6552950 is associated with the degree of anti-E2EP3 IgG response in 69 CHIKV-infected patients. Plasma collected at early convalescent (median 10 days post-illness onset) was subjected to E2EP3 peptide-based ELISA at a dilution of 1:2,000. Histogram shows the percentage of patients in each genotype exhibiting either low or high anti-E2EP3 IgG response. Low and high anti-E2EP3 IgG response were defined as being below or above the mean value of anti-E2EP3 IgG response, respectively. Association between SNP genotypes (AA, n = 38; AG, n = 20; GG, n = 11) and anti-E2EP3 IgG response (low—AA, n = 16; AG, n = 13; GG, n = 9; high—AA, n = 22; AG, n = 7; GG, n = 2) is performed using logistic regression analysis, *P = 0.0376.

D TLR3 SNP rs6552950 is associated with the CHIKV-specific antibody neutralizing capacity in 69 CHIKF patients. Plasma collected at early convalescent (median 10 days post-illness onset) was diluted 1:100–1:1,000 and mixed with CHIKV (MOI 10) for 2 h before infection of HEK 293T cells for 6 h. Assays were performed in quadruplicate, and data are expressed relative to virus-only-infected samples without sera. Data are presented as mean ± SEM. Comparison among SNP genotypes (AA, n = 38; AG, n = 20; GG, n = 11) is performed using one-way ANOVA analysis followed by Tukey’s multiple comparison test. ***P = 0.0001 1:100 serum dilution GG versus AA, ***P = 0.0001 1:100 serum dilution GG versus AG, *P = 0.0178 1:500 serum dilution GG versus AA, **P = 0.0056 1:500 serum dilution GG versus AG, **P = 0.0037 1:1,000 serum dilution GG versus AA, **P = 0.0041 1:1,000 serum dilution GG versus AG.

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et al., 2014), the precise roles of TLRs as PRRs for CHIKV remain unknown. In this study, TLR3 was demonstrated to be a critical PRR in the control of CHIKV replication, immunity and pathology in humans and mice. The increased susceptibility of both TRIF- and TLR3-/- primary fibroblasts to CHIKV infection demonstrates a role of TLR3 in controlling CHIKV replication. Furthermore, the loss of TLR3 expression in CHIKV-infected mice significantly increased viremia and exacerbated CHIKV-induced inflammation. However, these findings are in contrast to an earlier report where TLR3 was reported to play a modest role in controlling CHIKV infection in young mice (Schilte et al., 2010). This disparity could be due to the age of the animals and the different inoculation routes used. When inoculated intradermally, CHIKV triggered a strong local type I IFN response that was sufficient to locally control virus replication in WT mice and hence no effect on TLR3 signaling (Schilte et al., 2010). However, in this study, the effect of TLR3 signaling is apparent because the type I IFN response is independent of TLR3 and therefore insufficient to locally control CHIKV replication during the early phase of the infection.

By tracking virus dissemination using a recombinant Fluc-CHIKV, the loss of TLR3 clearly resulted in exacerbated CHIKV-induced pathology in mice. Moreover, a marked tropism was observed for skeletal muscle, joints and skin in WT and Tlr3-/- mice. The increased pathology in Tlr3-/- mice was mediated by an increased infiltration of Ly6G+ neutrophils into the infected joint footpad with F4/80+ macrophages, which were previously shown to be the primary cellular infiltrates (Gardner et al., 2010; Rudd et al., 2012; Teng et al., 2012). While the exact role of neutrophils in the development of CHIKV-induced inflammation remains unclear, neutrophils were recently implicated in the control of CHIKV infection in mouse (Dhanwani et al., 2014; Poo et al., 2014) and zebrafish (Palha et al., 2013) models. The increased infiltration of myeloid cells is accompanied by the reduction in CD4+ T-cell infiltration. However, depletion of CD4+ T cells prevented the pathology and abrogated neutrophils without interfering with the development of the viremia. This demonstrated that CD4+ T cells are essential for the pathology in WT and Tlr3-/- mice. One plausible explanation for the exacerbation of joint footpad inflammation in Tlr3-/- mice is that the loss of TLR3 may reduce the number of anti-inflammatory CD4+ T regulatory cells present in the infiltrate. However, this is highly unlikely as demonstrated by bone marrow reconstitution experiments. If TLR3 negatively controlled the number of CD4+ T regulatory cells, the WT mice reconstituted with Tlr3-/- bone marrow would have less CD4+ T regulatory cells. This was not case since WT mice reconstituted with Tlr3-/- bone marrow had the same level of joint inflammation as the WT mice reconstituted with WT bone marrow.

The dissemination of CHIKV into the brains of Tlr3-/- mice is of particular interest because, unlike other organs, the brains of WT mice were not infected. CHIKV is not commonly neurotropic, but the occurrence of CHIKV-associated neurological complications has been increasingly reported in patients since the 2006 Indian Ocean outbreak (Rampal et al., 2007; Wielanek et al., 2007; Das et al., 2010; Kashyap et al., 2010). Although previous studies in mice have shown that CHIKV dissemination into the central nervous system is dependent on type I IFN signaling (Coudere et al., 2008; Abraham et al., 2013), it remains to be elucidated whether this is a bystander effect due to overwhelming CHIKV replication or whether the blood–brain barrier has been breached due to the absence of TLR3, permitting virus entry into the brain.

Infection in bone marrow chimeric mice provided evidence on how TLR3-expressing hematopoietic cells are required to elicit an anti-CHIKV innate immune response and regulate CHIKV infection and pathology. Since the role of CD4+ T cells was shown to be

### Table 1. Association of TLR3 SNPs to CHIKV phenotype and severity.

<table>
<thead>
<tr>
<th>SNP</th>
<th>Chromosome</th>
<th>Position</th>
<th>Minor Allele</th>
<th>STAT</th>
<th>P-value</th>
<th>OR (L95–U95)</th>
<th>STAT</th>
<th>P-value</th>
<th>OR (L95–U95)</th>
</tr>
</thead>
<tbody>
<tr>
<td>rs3775292</td>
<td>4</td>
<td>187003025</td>
<td>C</td>
<td>3.06</td>
<td>0.002*</td>
<td>2.16 (1.31–3.42)</td>
<td>–0.63</td>
<td>0.53</td>
<td>0.80 (0.4–1.6)</td>
</tr>
<tr>
<td>rs6552950</td>
<td>4</td>
<td>186994856</td>
<td>G</td>
<td>2.11</td>
<td>0.03*</td>
<td>1.54 (1.03–2.29)</td>
<td>2.39</td>
<td>0.02*</td>
<td>2.31 (1.16–4.57)</td>
</tr>
<tr>
<td>rs7657186</td>
<td>4</td>
<td>186940439</td>
<td>A</td>
<td>1.62</td>
<td>0.11</td>
<td>1.43 (0.93–2.21)</td>
<td>–1.32</td>
<td>0.19</td>
<td>0.61 (0.29–1.28)</td>
</tr>
<tr>
<td>rs11721827</td>
<td>4</td>
<td>186991137</td>
<td>C</td>
<td>–1.22</td>
<td>0.12</td>
<td>0.70 (0.39–1.24)</td>
<td>–0.20</td>
<td>0.84</td>
<td>0.90 (0.33–2.46)</td>
</tr>
<tr>
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<td>4</td>
<td>18700256</td>
<td>T</td>
<td>1.08</td>
<td>0.28</td>
<td>1.28 (0.82–2.00)</td>
<td>–0.91</td>
<td>0.36</td>
<td>0.71 (0.33–1.5)</td>
</tr>
<tr>
<td>rs7668666</td>
<td>4</td>
<td>187001292</td>
<td>A</td>
<td>0.92</td>
<td>0.36</td>
<td>1.20 (0.81–1.78)</td>
<td>0.12</td>
<td>0.91</td>
<td>1.04 (0.54–2.01)</td>
</tr>
<tr>
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<td>4</td>
<td>187004074</td>
<td>T</td>
<td>0.82</td>
<td>0.41</td>
<td>1.18 (0.79–1.76)</td>
<td>0.47</td>
<td>0.64</td>
<td>1.15 (0.64–2.09)</td>
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<tr>
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<td>4</td>
<td>186994832</td>
<td>A</td>
<td>–0.42</td>
<td>0.68</td>
<td>0.92 (0.60–1.39)</td>
<td>–0.83</td>
<td>0.41</td>
<td>0.72 (0.33–1.56)</td>
</tr>
<tr>
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<td>A</td>
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<td>–0.89</td>
<td>0.37</td>
<td>0.74 (0.38–1.45)</td>
</tr>
</tbody>
</table>

*Population controls consist of 60 CEU, 60 CHB/JPT and 59 YRI individuals sequenced by the 1000 Genomes Pilot Project.

*Severe disease* is defined as patients who had either a maximum temperature greater than 38.5°C, a maximum pulse rate greater than 100 beats/min or a nadir platelet count less than 100 x 10^3/μL. 'Mild disease' is referred to patients who do not fulfill these criteria.

SNP, Single nucleotide polymorphism from TLR3 gene; SNPs rs5743316 and rs5743310 were excluded from further analysis due to failed Sequenom assay design and being monomorphic in control populations, respectively. SNPs rs13108688 was not in Hardy–Weinberg equilibrium in the population.

Chromosome, Position, Chromosome and corresponding position (in base pair) where the SNP is located.

Minor Allele, The rare allelic form of the SNP variant.

STAT, Statistic obtained for the regression analysis for the SNP.

P-value obtained for the regression analysis; *Significant P-value < 0.05 is indicated in bold.

OR(L95–U95), Odds ratio for the SNP as estimated from the minor allele with 95% confidence interval limits.
important for the pathology in the WT and Tlr3−/− mice via the control of neutrophils recruitment, this suggested that B-cell responses could be defective in Tlr3−/− mice. Growing evidence has also shown that activation of TLRs promotes B-cell proliferation and IgG isotype class switch (Ruprecht & Lanzavecchia, 2006; Xu et al., 2008; Sariol et al., 2011). Here, the production of CHIKV-specific IgM and IgG antibodies was not disrupted by the loss of TLR3, and the IgG response was even higher at 6 dpi. Rather, the reduced CHIKV-neutralizing capacity was due to diminished recognition for neutralizing linear B-cell epitopes located in the CHIKV E2 glycoprotein (Kam et al., 2012a,b; Lum et al., 2013). The loss of TLR3 expression could modulate CHIKV antigen (Ag) processing and presentation by APCs, particularly the immunodominant ‘E2EP3’ epitope, to result in an immunodominance shift (Siddiqui & Basta, 2011). Other virus infection models have demonstrated that TLR3 stimulation in dendritic cells decreases production of nitric oxide which in turn increases proteasomal activity and consequently increases viral antigen processing and presentation (Schwarz et al., 2000; Siddiqui et al., 2011).

Figure 8. Proposed model for TLR3-mediated anti-CHIKV response during infection. Upon CHIKV infection of the target cell, TLR3 functions (blue arrows) in synergy with multiple host cytosolic RNA sensors such as RIG-I and MDA5, to vanguard the coordinated endosomal and cytoplasmic recognition of CHIKV RNA, triggering anti-viral type I IFN response and expression of pro-inflammatory genes. This concerted activation acts to restrict CHIKV replication (red arrows). In parallel, the stimulation of cells will up-regulate TLR3 expression and enhance the anti-CHIKV response. Consequently, activation of TLR3 signaling mediates protection from CHIKV-induced pathology by preventing rapid CHIKV dissemination and restricting tissue tropism (red arrows) tied in with assistance from chemokine regulation (blue arrows). TLR3 signaling can also modulate CHIKV immune recognition during the adaptive immune response to promote effective CHIKV clearance through the production of CHIKV-specific immunodominant antibodies by B cells. Therefore, a loss of TLR3 would lead to a reduction in CHIKV antibody neutralizing capacity due to an impaired production of such antibodies, resulting in the persistence of CHIKV viremia in vivo and enhanced/increased disease severity.

Transcriptional analysis of PBMCs isolated from CHIKV-infected patients revealed that TLR3 expression was up-regulated during the acute and early convalescent phase of the disease. Transcriptional analysis of other PRRs such as RIG-I, MDA5 and IPS-1 (adaptor molecule involved in RIG-I and MDA5 mediated signaling) has shown that these molecules were similarly significantly up-regulated and associated with viral load during acute CHIKV infection (Teng et al., 2012), suggesting that the enhanced TLR3 expression is part of a general innate immunity against CHIKV. Previous genetic epidemiological studies have implicated inborn errors in TLR3 immunity and polymorphism in the pathogenesis of HSV-1 encephalitis (Zhang et al., 2007; Herman et al., 2012; Lafaille et al., 2012) and Influenza virus infection (Esposito et al., 2012). While the number of CHIKF patients in our study is too small to demonstrate any direct functional link between TLR3 gene polymorphisms and CHIKV disease outcomes, our findings suggest a possible relationship between the presence of TLR3 SNPs, rs3775292 and rs6552950 with an increased risk of CHIKV disease occurrence. In particular, the SNP rs6552950

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polymorphism was demonstrated to be associated with severe CHIKV disease outcome in patients from the Singapore 2008 CHIKV cohort. Antibody response studies in these patients have demonstrated that naturally acquired neutralizing IgG response is dominated by anti-E2E3 IgG antibodies (Kam et al., 2012b). Furthermore, the low anti-E2E3 IgG response in the SNP rs6552950 GG genotype patients was demonstrated to have reduced neutralizing antibodies, which in turn influenced the severity of disease outcomes in these patients. SNP rs3775292 was previously reported to associate with persisting IgG concentration and serum bactericidal antibody following serogroup C meningococcal polysaccharide-protein conjugate vaccination and an increased risk of colon cancer development (Moore et al., 2011; Slattery et al., 2011). It will be insightful to assess whether these SNPs would lead to a loss of TLR3 function in larger patient cohorts, including the emergent CHIKV outbreaks in the Caribbean islands (Leparc-Goffart et al., 2014).

Collectively, we provided clear evidences for TLR3 in the control of CHIKV infection. The synergy of TLR3 with multiple host RNA sensors such as RIG-I and MDA5 is indispensable for specific interactions with viral dsRNA to restrict CHIKV replication by inducing a rapid anti-viral type I IFN response together with pro-inflammatory gene expression (Fig 8). Essentially, TLR3-mediated immunity controls CHIKV-induced pathology by preventing rapid virus dissemination. The observations reported here also provide further insights on how TLR3-mediated innate responses against CHIKV infection can influence the adaptive immune response, as well as the mechanisms by which TLR3 modulates in vivo CHIKV immune recognition. These findings provide a better understanding on one of the key components that induce protective anti-CHIKV responses and will have critical implications in the future development of novel therapeutic strategies against CHIKV and other clinically important alphaviruses that have serious global health impacts.

Materials and Methods

Study approval

Blood samples from CHIKF patients used in this study were collected by Institute of Infectious Disease and Epidemiology at Tan Tock Seng Hospital from 1 August through September 23, 2008 with approval from the National Healthcare Group’s domain-specific ethics review board (DSRB Reference no. B/08/026). Written informed consent was obtained from all participants in accordance with the Declaration of Helsinki principles. All animal protocols were approved by the Institutional Animal Care and Use Committee (IACUC no. 120714) at the Biological Resource Center at Biopolis, Singapore. All studies involving animals are reported in accordance with the ARRIVE guidelines for reporting experiments involving animals (Kilkenny et al., 2010).

Study population

A total of 94 PCR-confirmed CHIKV-positive individuals were included in this study. There were 48 Chinese, 26 Malay, 10 Indians and 10 of other ethnicity. The median age was 38 with a range of 21 to 67 years of age with 77.7% male and 22.3% female. Based on the clinical parameters defined in Chow et al (2011) and Ng et al (2009), these individuals were further classified into 53 mild and 41 severe cases. Illness was defined as ‘severe’, if a patient had either a maximum temperature greater than 38.5°C, a maximum pulse rate greater than 100 beats/min or a nadir platelet count less than $10^3/\text{l}$. Patients who do not fulfill these criteria are classified as ‘mild’ (Ng et al, 2009; Chow et al, 2011).

Cell culture

African green monkey kidney epithelial cells (Vero-E6), HEK293T, and a mouse hepatocyte cell line (Hepa 1–6) were cultured in DMEM supplemented with 10% FBS (Gibco). Mouse tail fibroblasts were cultured in DMEM supplemented with 10% FBS and 1% penicillin-streptomycin. Aedes albopictus mosquito cell line (C6/36) was cultured in Leibovitz’s L-15 medium (Life Technologies) supplemented with 10% FBS. All cells were maintained at 37°C with 5% CO₂, except for the C6/36 cell line which was maintained at 28°C without CO₂ supplementation.

Genotyping and association analysis

A total of 11 tagSNPs were identified to cover the linkage disequilibrium block for TLR3 gene. Genotyping was performed using matrix-assisted laser desorption/ionization time of flight mass spectrometry (MALDI-TOF MS) for the determination of allele-specific primer extension products using Sequenom’s MassARRAY system and iPLEX technology (Sequenom Inc). One SNP (rs5743316) failed Sequenom assay design. The design of oligonucleotides was carried out according to the guidelines of Sequenom and performed using MassARRAY Assay Design software. Multiplex PCR amplification of amplicons containing the SNPs of interest was performed using QIAGEN HotStart Taq Polymerase using 5 ng of genomic DNA. Primer extension reactions were carried out according to manufacturer’s instructions for iPLEX chemistry. Assay data were analyzed using the Sequenom TYPER software. Clustering of genotype calls was evaluated to determine that the clustering was sufficient for inclusion in the statistical analysis. All SNPs were tested for Hardy–Weinberg equilibrium for quality control and subjected to further statistical analysis. Population controls used for association analysis were the 179 samples sequenced by the 1000 Genomes Pilot Project (60 CEU, 60 CHB+JPT, 59 YRI) (1000 Genomes Project Consortium et al., 2010). Sequenom genotypes for SNPs in reversestrand notation (SNPs rs3775291 and rs3775292) were converted to a forward strand notation using the option –flip provided by the software PLINK v1.07 (Purcell et al., 2007). One of the 10 TLR3 SNPs (rs5743310) was monomorphic in two of the three control populations (CEU and CHB/JPT) and was therefore excluded from further analysis. Association between the other 9 TLR3 SNPs and the CHIKV infection phenotype was computed using a logistic regression model. Association between TLR3 SNPs and severity (mild/severe) was computed using a logistic regression model that included gender and age as covariates. One SNP rs13108688 was not in Hardy–Weinberg equilibrium ($P < 0.005$) in the population; however, this SNP was not significant for association. Both association analyses were performed using the software PLINK v1.07 (Purcell et al, 2007).
lower than 0.05 were considered significant. The distribution of each TLR3 SNP genotype and disease severity for each SNP genotype is listed in Supplementary Table S2.

Virus stocks

CHIKV-IMT isolate used for in vitro infections of human cells was isolated from Reunion Island during the 2006 CHIKF outbreak (Bessaud et al., 2006). Virus stocks were prepared in Vero-E6 cultures, washed and pre-cleared by centrifugation before storing at −80°C. CHIKV-SGP011 isolate used for in vitro and in vivo infections in mouse studies was isolated from an outbreak in Singapore in 2008 at the National University Hospital (Her et al., 2010) and propagated in Hepa 1–6 and C6/36 cultures, respectively. CHIKV variants expressing Fluc was constructed using a full-length infectious cDNA clone of CHIKV LR2006-OPY1 isolate as described (Tsetsarkin et al., 2006; Pohjala et al., 2011; Teng et al., 2012). Infectious viruses were propagated in C6/36 cultures, washed and pre-cleared by ultracentrifugation before storing at −80°C. Virus titer of all virus stocks used was determined using standard plaque assays with Vero-E6 cells (Her et al., 2010; Kam et al., 2012b).

Virus infections

CHIKV infections on primary fibroblasts and continuous cell lines from mouse and human were performed at multiplicity of infection (MOI) 10. Each infection mix consisted of virus suspension prepared from mouse and human stocks used was determined using standard plaque assays with Vero-E6 cells (Her et al., 2010; Kam et al., 2012b).

Flow cytometry and antibodies

Detection of CHIKV Ag was carried out in a two-step indirect intracellular labeling process (Her et al., 2010; Teng et al., 2012). Data were acquired in BD FACSCanto™ II (BD Bioscience) using BD FACSDiva™ software. Dead cells and duplets were excluded in all analysis with FSC/SSC gating. Results were analyzed with FlowJo version 7.5 software (Tree Star, Inc). Antibodies against CHIKV Ag were purchased from Santa Cruz Biotechnology. Antibodies against mouse CD45 (cat# 557659) and CD8 (cat# 553035) were purchased from BD Bioscience. Antibodies against mouse CD11b (cat# 12-0112-82) was purchased from eBioscience. Antibodies against mouse CD3 (cat# 100200), CD4 (cat# 100531) and Ly6G (cat# 127612) were purchased from Biolegend. Mouse CD4-depleting antibody (cat# BE0003-1) and rat IgG2b isotype control (cat# BE0090) were purchased from Bio X cell.

Viral RNA extraction and viral load analysis

Viral RNA was extracted using QIAamp Viral RNA Mini Kit (QIAGEN) according to manufacturer’s instructions. Quantification of CHIKV non-structural protein (nsP) 1 negative-sense RNA was determined according to a quantitative real-time PCR (qRT-PCR) TaqMan assay adapted from Plaskon et al. (Plaskon et al., 2009) using QuantiTect Probe RT-PCR Kit (QIAGEN) in 12.5 µl reaction vol. All reactions were performed using 7900HT Fast Real-Time PCR System machine (Applied Biosciences) with thermal cycling conditions as described previously (Teng et al., 2012). The limit of detection was 10 RNA copies/µl.

Total RNA extraction and gene expression analysis

Total RNA was extracted using RNeasy Mini Kit (QIAGEN) according to manufacturer’s instructions. Quantification of total RNA was performed using NanoDrop 1000 Spectrophotometer (Thermo Scientific), and RNA samples were further diluted to 10 ng/µl. qRT-PCR was performed using QuantiFast™ SYBR Green RT-PCR Kit (QIAGEN) according to manufacturer’s recommendations in 12.5 µl reaction vol. All reactions were performed using 7900HT Fast Real-Time PCR System machine (Applied Biosciences) with thermal cycling conditions as described (Teng et al., 2012). The fold change for each gene between CHIKV-infected and mock-infected was calculated as $2^{-\Delta\Delta Ct}$ (Teng et al., 2012). The primer sequences of the mouse genes analyzed are listed in Supplementary Table S3.

Animal studies

Three-week-old WT or Tlr3−/− C57/BL6 female mice were inoculated s.c. in the ventral side of the right-hind footpad toward the ankle with 10⁶ PFU CHIKV in 25 µl in a non-randomized and non-blinded fashion. Viral RNA extraction was performed from 10 µl of blood collected from the tail, and viremia was determined by qRT-PCR as described (Teng et al., 2012; Lum et al., 2013; Teo et al., 2013). For joint (footpad) inoculated mice, joint inflammation by the measurement of the height (thickness) and the breadth of the footpad using a vernier caliper was calculated as [height × breadth]. The degree of inflammation was expressed as relative increase compared to pre-infection (day 0; d 0) with the following formula: [(x − d 0)/d 0] where x is footpad size measurements for respective dpi as described (Kam et al., 2012b). For joint footpad extraction, mice were sacrificed by terminal anesthesia with ketamine [150 mg/kg]/xyloLAzin [10 mg/kg] followed by intra-cardial perfusion with PBS. Joint footpads were removed and preserved in Trizol (In-vitrogen) at −80°C. Tissues were homogenized using a rotor–stator homogenizer (Xiril Dispomix) at 500 g for 15 s. Homogenized tissues were transferred to clean tubes and mixed with 230 µl of chloroform. Following 2-min incubation, tissue mixtures were centrifuged at 13,523 g for 15 min at 4°C. The aqueous phase was collected, and total RNA isolated as described (Teng et al., 2012). For bone marrow chimera, 6-week-old recipient mice were irradiated twice with 600Rad (3 h apart) and i.v. injected with 10⁶ donor bone marrow cells. Absolute CD45⁺ leukocyte blood count was determined at week 4–6 post-reconstitution using flow cytometry and compared to non-chimeric mice of same age. To test for successful adoptive bone marrow cell transfer, Tlr3 expression in peripheral blood leukocytes from reconstituted mice was determined 6 weeks after reconstitution using quantitative real-time PCR analysis before proceeding with infection. Depletion of CD4⁺ T cells was performed as described (Teo et al., 2013). Each mouse was i.p. injected with 500 µg of either CD4-depleting antibody or rat IgG2b isotype control on −2, −1 and 4 dpi. Complete CD4⁺ T-cell depletion was assessed before CHIKV inoculation (day 0).

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In vivo imaging

Bioluminescence signals were assessed daily from 1 to 8 dpi and subsequently on every alternate day until 20 dpi using an in vivo bioluminescence imaging system (IVIS Spectrum, Xenogen) as described (Teo et al., 2013). Luciferin solution containing the luciferase substrate, D-luciferin potassium salt (Caliper Life sciences), was prepared by dissolving in PBS at a concentration of 5 mg/ml. Mice were shaved and anesthetized in an oxygen-rich induction chamber with 2% isoflurane. Bioluminescence signals were measured 2 min after s.c. injection of 100 μl of luciferin solution. Whole-body imaging was performed with the animals in a ventral position, while the feet and head were imaged with the animals in a dorsal position. Bioluminescence imaging was acquired with a field of view (FOV) of 21.7 cm for whole body (FOV-D) and 13.1 cm for foot and head (FOV-C). The mice were exposed for an initial 60 s, followed by a 4-min delay before another exposure at 60 s. When luminescence readings were above the upper detection limit of machine, the exposure time was reduced and kept consistent across groups. Bioluminescence signals of the region of interest were quantified using the Living Image 3.0 software (Caliper Life sciences) and expressed as average radiance (p/s/cm²/sr). The lowest detection limit is 0 p/s/cm²/sr.

Histology

Mice were terminally anesthetized with ketamine [150 mg/kg]/xylazine [10 mg/kg] and perfused with PBS by in-tra-cardial injection. Tissues were fixed in 4% paraformaldehyde (PFA), decalcified and embedded in paraffin wax before 5-μm-thick sections were cut and underwent H&E or immunohistochemical (IHC) staining against mouse Ly6G (Biolegend, cat# 127602), CD11B (AbCam, cat# ab75476) and F4/80 (DAKO, cat# K3468) using established protocols (Teng et al., 2012).

Virion-based ELISA

CHIKV-specific antibody titers were assessed by virion-based ELISA (Kam et al., 2012c). Polyacrylate 96-well microtiter plates (MaxiSorp, Nunc) were coated with purified CHIKV (10⁶ infectious units per well). Wells were blocked with PBS containing 0.05% Tween 20 and 5% non-fat milk (0.05% PBST + 5% milk) for 1.5 h at 37°C. Serum samples were diluted in 0.05% PBST + 2.5% milk before adding to wells and incubating for 1 h at 37°C. Separately, HRP-conjugated goat anti-mouse IgM (Santa Cruz, cat# sc-2064) or IgG (Santa Cruz, cat# sc-2005) were used to detect mouse antibodies bound to virus-coated wells. Reactions were developed using 3,3′,5,5′-tetramethylbenzidine substrate (Sigma-Aldrich) and terminated by Stop reagent (Sigma-Aldrich). Absorbance at 450 nm was measured using a TECAN Infinite M200 microplate reader and analyzed using Magellan™ software.

Sero-neutralization

Neutralizing capacity of antibodies from CHIKV-infected mice was analyzed by immunofluorescence-based cell infection assays using HEK 293T cells (Kam et al., 2012b). Different amounts of infectious virus required to achieve an infection of MOI 10 were mixed with diluted (1:50–1:5,000), heat-inactivated human plasma or pooled mouse sera and incubated for 2 h at 37°C with gentle agitation (350 rpm). Virus–antibody mixtures were then added to HEK 293T cells seeded on fibronectin-coated 96-well plates and incubated for 1.5 h at 37°C. Virus inoculum was removed and replaced with DMEM supplemented with 10% FBS and incubated for 6 h at 37°C before being fixed with 4% PFA and permeabilized in PBS containing 0.2% Tween 20. Cells were incubated with mouse anti-alphavirus mAb (Santa Cruz, cat# sc-58088) at 1:500 dilution in PBS followed by incubation with Alexa Fluor 488-conjugated goat anti-mouse IgG F(ab)’2 ab (Invitrogen, cat# A11017) at 1:300 dilution in PBS. Nuclei were stained with DAPI (1 μg/μl stock) at 1:10,000 dilution in PBS followed by immunofluorescence quantification using the Cellomics ArrayScan high content analysis reader (Thermo Scientific).

Peptide-based ELISA

Streptavidin-coated polyacrylate 96-well microtiter plates ( Pierce, Thermo Scientific) were blocked with 0.1% PBST supplemented with 1% w/v sodium caseinate (Sigma-Aldrich) for 1 h at RT before being coated with biotinylated 18-mer overlapping peptides synthesized based on consensus E2 glycoprotein sequence (Kam et al., 2012b) for 1 h at RT. Mouse serum diluted 1:500 (mouse) or human plasma diluted 1:2,000 in 0.1% PBST and 0.1% w/v sodium caseinate were added and incubated for 1 h at RT followed by the
addition of relevant HRP-conjugated goat secondary IgG (Santa Cruz, cat# sc-2005) to detect bound antibodies. Reactions were developed using 3,3′,5,5′-tetramethylbenzidine substrate (Sigma-Aldrich) and terminated by Stop reagent (Sigma-Aldrich). Absorbance at 450 nm was measured using TECAN Infinite M200 microplate reader and analyzed using Magellan™ software.

Statistical analysis

Statistics were performed using Prism 5.01 (GraphPad Software). Pairwise comparison between infected and mock-infected primary fibroblasts from human and mice was performed using two-tailed unpaired t-test. Pairwise comparison between WT and Tlr3−/− mice in animal studies was performed using two-tailed Mann-Whitney U-test. TLR3 SNPs association analysis (disease severity and anti-E2EP3 IgG response) was performed using logistic regression analysis, while CHIKV-neutralizing capacity association was performed using one-way ANOVA analysis followed by Tukey’s multiple comparison test. P-values less than 0.05 are considered statistically significant.

Supplementary information for this article is available online: http://embomolmed.embopress.org

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Author contributions

ZH, T-ST, JJLT, T-HT, F-ML, WWLL, CG, RM, AKA, Y-WK, VL, AL, MKW, AC, SKB, Y-SL and AM performed the experiments, provided the clinical and experimental reagents and analyzed the data. ZH, T-ST, JJLT, T-HT, ML, LR and LFPN conceptualized the study, analyzed the data and wrote the manuscript. All authors read and approved the manuscript.

Conflict of interest

The authors declare that they have no conflict of interest.

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