Chikungunya disease is a severely debilitating, mosquito-borne, viral illness that has reached epidemic proportions in Africa, Asia, and the islands of the Indian Ocean. A mutation enhancing the ability of the chikungunya virus (CHIKV) to infect and be transmitted by *Aedes albopictus* has increased the geographical range at risk for infection due to the continuing global spread of this mosquito. Research into disease pathogenesis, vaccine development, and therapeutic design has been hindered by the lack of appropriate animal models of this disease. The meticulous study reported in this issue of the *JCI* by Labadie et al. is one of the first reports describing CHIKV infection of adult immunocompetent nonhuman primates. Using traditional and modern molecular and immunological approaches, the authors demonstrate that macaques infected with CHIKV are a good model of human CHIKV infection and also show that persistent arthralgia in humans may be caused by persistent CHIKV infection of macrophages.
A nonhuman primate model of chikungunya disease

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Chikungunya disease is a severely debilitating, mosquito-borne, viral illness that has reached epidemic proportions in Africa, Asia, and the islands of the Indian Ocean. A mutation enhancing the ability of the chikungunya virus (CHIKV) to infect and be transmitted by Aedes albopictus has increased the geographical range at risk for infection due to the continuing global spread of this mosquito. Research into disease pathogenesis, vaccine development, and therapeutic design has been hindered by the lack of appropriate animal models of this disease. The meticulous study reported in this issue of the JCI by Labadie et al. is one of the first reports describing CHIKV infection of adult immunocompetent nonhuman primates. Using traditional and modern molecular and immunological approaches, the authors demonstrate that macaques infected with CHIKV are a good model of human CHIKV infection and also show that persistent arthralgia in humans may be caused by persistent CHIKV infection of macrophages.

Arthrogenic alphaviruses are widely distributed. For example, Ross River virus (RRV) affects thousands every year in Australia, causing polyarthritis disease; Mayaro virus persists in the Amazon rainforest, causing small, sporadic outbreaks; and O’nyong-nyong virus is known to cause explosive outbreaks of thousands of people in central Africa. However, the arthrogenic alphavirus that has become most well known in recent years is chikungunya virus (CHIKV), the subject of the study by Labadie et al. in this issue of the JCI (1).

The comprehensive review “Chikungunya virus disease,” published in volume II of Tom Monath’s landmark series The arboviruses: epidemiology and ecology (2), has remained the classic reference for this virus for more than 20 years. However, after the 2005–2006 CHIKV epidemic in the Indian Ocean Islands, several excellent reviews have been published (3–5). CHIKV was first isolated in Tanzania in 1952, during a severe dengue fever–like epidemic (6). However, chikungunya fever was likely confused with dengue for hundreds of years, with the first possible record of chikungunya fever published in Cairo in 1779 (7). It has been speculated that CHIKV was introduced into the Americas in the early 1800s, but failed to establish a sustainable transmission cycle (8).

Since 1952, laboratory-confirmed outbreaks of chikungunya fever have occurred almost annually in south and central Africa and southeast Asia (2, 4). Between 1990 and 1996, very few outbreaks of chikungunya fever were reported, except in Thailand. Chikungunya fever outbreaks became more frequent and widespread, starting in 2003 in Indonesia. In 2005–2006, the islands of the Indian Ocean, including Seychelles, Mauritius, Madagascar, and La Réunion, were devastated by widespread epidemics of CHIKV infection (4). A chikungunya fever epidemic occurred in India during 2006–2007, where it is estimated that more than 1.4 million people were infected. In July 2007, CHIKV emerged from the tropics and caused a localized outbreak in Ravenna, Italy (9). In 2008–2009, CHIKV was reported in countries of southeast Asia, including India, Thailand, and Singapore. The epidemic continues at the time of this writing, with an estimated 12,000 cases in Indonesia’s Lampung province reported between mid-December 2009 and mid-January 2010.

Clinical presentation of CHIKV infection

CHIKV infection in humans usually causes a nonlethal, self-limiting, febrile illness. Until recently, much of our knowledge of human disease has been descriptive. Patients developed viremias that could exceed 10^9 mouse LD50/ml and last for 4 days, and displayed nonfatal symptoms of variable severity that could persist for many months (10, 11). Chikungunya fever is usually characterized by high fever, arthralgia, and rash. During the acute phase, patients can experience painful and disabling arthritis that can last for 7 days (12). During some outbreaks, two-thirds of patients have had to be hospitalized (12). Unfortunately, after the acute phase, polyarthritis can be recurrent and may persist for up to several years after infection (4). Other symptoms of CHIKV infection can include retro-orbital pain, neurological and hemorrhagic manifestations, and myocarditis. In the most recent epidemics, many nontypical symptoms were reported for the first time, including lethal hepatitis, encephalitis, maternal-fetal transmission, and an increased death rate (4).

The epidemic infecting approximately 300,000 people on the island of La Réunion (a French territory popular with tourists), the importation of cases to many other countries, and establishment of a locally transmitted epidemic in Italy was the impetus for multiple governments to provide a considerable infusion of funding to stimulate major research efforts to improve our understanding of this hitherto largely ignored disease. The apparently increased severity of infection (with reports of approximately 250 fatalities during the Indian Ocean epidemic; ref. 1) may be attributed to several factors, for example, naivety of the afflicted population, preexisting medical conditions, and better reporting of the illness in a modern health care setting. The well-documented A226V mutation in CHIKV, which has arisen independently in several locations, does not seem to be correlated with disease severity, although it has been proven to influence infectivity for the Asian tiger mosquito (Aedes albopictus; refs. 13–16). This adaptation is a cause for concern, as this highly invasive and relatively cold-tolerant mosquito is gradually spreading to new areas in Europe and elsewhere (17). Some climate change proponents have suggested that recent chikungunya fever epidemics might be climate

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Subcutaneous injections of CHIKV in young to provide a good working background for outbred mice, although they induce disease reproducing true human-like symptoms. Similar in symptomology to CHIKV infection, re-creating true human-like symptoms. These symptoms may be correlated with recently published observations of the symptoms in human cases of chikungunya fever (4, 5, 12, 23). In the acute phase, nonhuman primates show high viremia (approximately $10^8$–$10^9$ viral RNA copies/ml), fever, and rash. They also have increased levels of aminotransferase, including aspartate transaminase (AST) and alanine transaminase (ALT), as well as increased expression of IFN, MCP-1, and IL-6. The nonhuman primates also have abnormal blood chemistries, including lymphopenia, monocyteopenia, and granulocytosis. Humans in the acute phase of chikungunya fever present with fever, fatigue, headache, myalgia, and rash as well as increased AST and ALT. Viremia in humans ranges from $10^7$ to $10^8$ viral RNA copies/ml. Leukopenia and thrombocytopenia are common. During the subacute phase in nonhuman primates, there is macrophage infiltration, and CHIKV is present in lymphoid tissue, liver, meninges, joints, and muscle. The subacute phase is not well characterized in humans, but includes the persistence of myalgia, arthralgia, and tenosynovitis. In the chronic phase in nonhuman primates, there was CHIKV persistence in macrophages, lymphoid tissue, and liver tissue up to 2 months after infection. The chronic phase in humans includes persistent and recurrent arthralgia that can last for months to years.

**Mouse models of CHIKV and RRV**

As with dengue fever, the lack of a good animal model replicating human chikungunya fever symptoms and pathology has been a major obstacle for understanding the course of infection, spectrum of disease severity, and persistence of symptoms. A mouse model for RRV, predating the recent work with CHIKV, incorporates both traditional and state-of-the-art molecular and immunological assays that can be readily interpreted by a broad audience in the context of previous work. The synergism of using multiple techniques and the duration of the study has begun to explain the mechanisms underlying the long-term persistence of chikungunya fever symptoms. The reported correlation between dose and disease severity (more severe at a higher dose) is interesting and may explain, at least in part, the range of symptoms observed in humans. The doses used by the authors are within ranges likely delivered by mosquitoes (24). It would have been interesting if some primates had been challenged by CHIKV-infected mosquitoes, given the increasing evidence that chikungunya fever has an immunological component. Mosquito saliva is known to have immunomodulatory activity, and, for several viruses (for example, West Nile virus), virus delivery in the context of saliva can influence infection establishment, dissemination, and severity (25, 26). As Labadie et al. state, the

**Figure 1** Comparison of symptoms resulting from naturally acquired human CHIKV infection and observed symptoms in a laboratory-infected nonhuman primate model of chikungunya disease. In their study in this issue of the JCI, Labadie et al. describe the stages of symptoms of CHIKV infection seen in their primate model as having 3 phases: acute, subacute, and chronic (1). The symptoms and pathology observed in the nonhuman primate model resembled many of those previously reported for CHIKV infection in humans, and so the data are a significant contribution to our understanding of the disease (Figure 1 and refs. 5, 23). It is gratifying that the study was not driven by the use of highly sensitive molecular techniques that do not distinguish between infectious and defective virus, but incorporated more traditional infectivity and immunological assays that can be readily interpreted by a broad audience in the context of previous work.
The macaque model should help in the development of new therapeutic or prophylactic strategies against chikungunya fever and facilitate testing of new interventions (1). This work represents a considerable advance over early primate studies (27, 28), and development of a nonhuman primate model of CHIKV infection has been a focus of several groups. The in utero infections observed during the La Réunion epidemic have been evaluated in a macaque model (29). Importantly, a macaque model has been developed at the NIH laboratories to determine the efficacy of a much-needed CHIKV vaccine and has demonstrated excellent protection against challenge with virulent CHIKV (30). This model, like that reported here by Labadie et al., mimicked key features of human chikungunya disease.

Labadie et al.’s current work in macaques (1) shows a good correlation with previously published work with mice. With little neurological involvement and virus prevalence in immune tissues and muscles, the present model shows promise. The authors admit that a limitation of their model is the lack of joint and muscle pathology in the macaques, but note that the occurrence of these symptoms in humans is restricted to the most severe cases, although recent reports have stated that arthralgia/arthritis actually affects 73%–80% of patients (5). Another limitation of the Labadie et al. paper is that no serology studies were done. It would be interesting to see the level and types of antibodies that the macaques produced in response to CHIKV infection.

**Immunopathology during CHIKV infection**

Labadie et al. present a compelling argument for the importance of macrophages in causing CHIKV-induced disease (1). Macrophages have been shown to be one of the keys to the destructive muscle necrosis, and possibly viral persistence, seen in RRV infection in mice. These early observations of RRV infection in mice are substantiated in the work by Labadie et al. Rulli et al. discussed the recent RRV data showing that macrophages are a key mediator of muscle necrosis and pathology and that, in vitro, RRV can persistently infect macrophages and upregulate release of IFN-γ, TNF-α, and monocyte chemoattractant protein–1 (MCP-1; ref. 31). Macrophage infiltration into muscles and joints in RRV-infected mice was responsible for the production of cytokines and chemokines, such as TNF-α, macrophage inflammatory protein–1α (MIP-1α), MCP-1, IL-1β, and IFN-γ (19). In the CHIKV study by Labadie et al., infected macaques showed increased expression of IFN-α/β, CCL2, CCL3, CCL4, IL-6, and IFN-γ and moderate increases of TNF-α (1). These findings are what would be expected of a viral infection inducing a strong macrophage response. The typical antiviral response in humans is characterized by the induction of cytokines such as IFN-α/β, IFN-γ, IL-12, and IL-18 and the activation of macrophages, NK cells, dendritic cells, neutrophils, and complement. TNF, IL-1, and IL-6 are produced by immune cells as signals of inflammation. Cytokines IL-4 and IL-10, which are not upregulated during viral infection, were not upregulated in the current study (1), again emphasizing the importance of macrophages.

**Wild primates and CHIKV transmission cycles**

The demonstration of *M. fascicularis* susceptibility to CHIKV infection (1) raises an interesting epidemiological question that merits further investigation. Within what can be considered as the endemic/original geographical region of Africa in which CHIKV circulates, primates have been implicated in a transmission cycle that does not involve humans. McIntosh et al. (32) reported CHIKV seropositive, wild-ranging vervets (*Cercopithecus aethiops pygerythrus*) and baboons (*Papio ursinus*) collected in the 1962 endemic area of southern Rhodesia and experimentally recorded that viremias in these animals lasted for 3–4 days in both species, “at levels significantly and consistently higher than those of the African green monkey, *Cercopithecus aethiops*” (33). In Asia, the potential involvement of primates in the transmission cycle is unclear. Inoue et al. reported an IgM seropositive rate of 14.8% for *M. fascicularis* in the Philippines (34), and Marchette et al. reported seropositive monkeys in Malaya — although 15 out of 16 were also positive for other related viruses, which can compromise the tests’ accuracy (35). However, Peiris et al. found no CHIKV positivity among 115 wild toque macaques (*Macaca sinica*) from Sri Lanka (36). The suggestion that there is a sylvatic cycle in Asia must be treated with caution, since evidence of infection in a wild animal must not be interpreted as proof of infection in a transmission cycle. Specific criteria must be met when considering whether a vertebrate is involved in virus transmission. Basically, the vertebrate must be spatially and temporally coincident with a known vector, be naturally fed upon by that vector, and produce viremias of sufficient magnitude and duration to infect the vector. Finally, transmission between the vertebrate and vector must be experimentally demonstrated. To date, these criteria have not been satisfied for primates in Asia, so we cannot assume that primates play any significant role in the Asian CHIKV transmission cycle. This is a gap in our knowledge that needs to be filled, and one to which the study by Labadie et al. contributes (1). *M. fascicularis* is widespread throughout Asia (37) and has been introduced into Hong Kong, Indonesia, Mauritius, and Palau. It is interesting that the species is present in large numbers on Mauritius (30,000 estimated; ref. 38), but there are no compelling data to implicate its involvement in the large chikungunya fever epidemic in the region (C. Chastel, personal communication).

Overall, the development and characterization of primate models for chikungunya fever represent a significant advance in the field that has already increased our understanding of the disease and facilitated evaluation of new potential vaccines. It is hoped that identification of the underlying mechanisms of the disease will be translated into a reduction of the consequences of infection and new preventive strategies that ultimately reduce disease incidence.

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While erudite cell biologists have for many decades described singular immotile appendages known as primary cilia to be present on most cells in our bodies, cilial function(s) long remained an enigma. Driven largely by an ever increasing number of discoveries of genetic defects in primary cilia during the past decade, cilia were catapulted from a long lasting existence in obscurity into the bright spotlight in cell biology and medicine. The study by O’Toole et al. in this issue of the JCI adds a novel “enzymatic” facet to the rapidly growing information about these little cellular tails, by demonstrating that defects in the XP/NPEP3 gene, which encodes mitochondrial and cytosolic splice variants of X-prolyl aminopeptidase 3, can cause nephrophthisis-like ciliopathy. Future studies are in order now to elucidate the cystogenic pathways affected by disrupted enzymatic function of XP/NPEP3 in cilia-related cystic diseases.

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Based on a flood of recent evidence, primary cilia are now heralded as sensory organs for detection and transmission of a broad range of cues from the extracellular environment of cells (1), including mechanical and chemical information as diverse as fluid flow in kidneys, mechanical bone deformation, and light and odorant detection (2). By processing such physical and chemical information from the environment into molecular signals in development and postnatal growth and homeostasis, cilia can affect cell differentiation and polarity and cell cycle control (3).

Genetic damage to primary cilia results in a spectrum of perplexing disorders with seemingly disparate manifestations, now classified as ciliopathies. A growing list of bona fide ciliopathies at present includes Bardet-Biedl syndrome (BBS), nephropathies (NPHP), and Senior-Loken syndrome (SNLS), just to name a few. Common