

Plasmodium knowlesi: experimental model, zoonotic pathogen and golden opportunity?

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INTRODUCTION

Plasmodium knowlesi, a parasite adapted to Southeast Asian macaques, occupies a unique position among the *Plasmodium* species. Not only is *P. knowlesi* pathogenic when it crosses from the natural macaque hosts into the human population, it is also permissive in a range of experimental non-human primates (NHPs). In deed *P. knowlesi* is a parasite well known to the malaria research community and has been used extensively as a model for parasite biology, drug and vaccine development since 1932, reviewed in this issue (Butcher and Mitchell, 2016; Pasini *et al.* 2016; Galinski *et al.* 2017). Juxtaposed to research, *P. knowlesi* was virtually unknown to contemporary medical practitioners until 2004 when a large focus of naturally acquired *P. knowlesi* malaria was reported in the human population of Sarawak, Malaysian Borneo (Singh *et al.* 2004). Remarkably detailed information on several experimental lines of *P. knowlesi* existed long before *P. knowlesi* was recognized as an emerging pathogen in the human population of Southeast Asia.

To put the new millennium find into context, the last of the four *Plasmodium* species associated with malaria in the human population, *Plasmodium ovale*, was described in 1922 (Coatney *et al.* 1971a). The other human-host adapted species are *Plasmodium falciparum*, *Plasmodium vivax* and *Plasmodium malariae*. Of these, *P. falciparum* is the most virulent and prevalent species responsible for most of the estimated 212 million (range 148–304 million) new cases of malaria and 429 000 (range 235 000–639 000) malaria-related deaths, in 2015 (WHO, 2016).

Consequently, malaria research was, and still is, underpinned by the quest to understand, control and prevent malaria caused by *P. falciparum*. In addition to virulence as a distinguishing feature, *P. falciparum* has other unique features when compared to the other human-host adapted *Plasmodium* species, the most striking is binding of *P. falciparum*-infected red blood cells to host endothelium, recently reviewed (Mosnier and Lavstsen, 2016; Wassmer and Grau, 2017). While some of the unique characters are more

compelling virulence candidates than others, all are difficult to test.

EXPERIMENTAL MODEL AND ZOONOTIC PATHOGEN

Providing experimentally sound evidence that particular characteristics of *P. falciparum* are indeed related to virulence and taking these forward to develop treatments and augmentative therapies require representative model systems, including modelling *in vivo*. Indeed, an *in vivo* model system remains the missing link in translational research on malaria pathophysiology. Even though the falciparum clade of *Plasmodium* species has recently had new members added, including the multiple variants of *Plasmodium praefalciparum*, *Plasmodium billcollinsi*, *Plasmodium blacklocki*, *Plasmodium adleri* and *Plasmodium gaboni*, *P. falciparum* isolated from human infections is more or less resolutely human host restricted (Loy *et al.* 2017). On this back-drop and, despite huge research efforts to understand the pathophysiology of severe malaria, outputs remain largely hypothetical.

The very recent finding that, like *P. falciparum*, *P. knowlesi* causes severe and fatal infections in the human population opens the possibility for comparative research on malaria parasite virulence. Importantly, and in contrast to *P. falciparum*, *P. knowlesi* is experimentally permissive in a range of NHPs, conferring a means to test virulence (Cox-Singh and Culleton, 2015). Is it possible that *P. knowlesi* may help resolve what is effectively a research bottleneck?

THE PROBLEM WITH MODELS

Phylogenetically *P. knowlesi* falls within the old-world monkey clade, a clade that includes *P. vivax*, and is evolutionarily relatively distant to *P. falciparum*. Perceived wisdom built on *Plasmodium* phylogenies determined that *P. knowlesi*, while a valuable model for parasite biology, was considered too evolutionarily distant from *P. falciparum* to model for malaria pathophysiology and considered more suitable to model for *P. vivax* (Craig *et al.* 2012). A view supported by Martinelli and Culleton, this issue (Martinelli and Culleton, 2016).

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In addition, the most common experimental partner for *P. knowlesi* is the Rhesus macaque and, while used extensively, this model is not optimal to represent severe malaria in the human host. This is because most contemporary laboratory colonies of Rhesus macaques do not mount functional resistance to experimental infections with *P. knowlesi* (Praba-Egge *et al.* 2002). Consequently Rhesus monkeys infected with *P. knowlesi* develop an overwhelming parasitaemia and a disease phenotype that is difficult to reconcile with severe *P. falciparum* or even, latterly, severe *P. knowlesi* infections in the human host (Cox-Singh and Culleton, 2015).

While the hyper-susceptible phenotype in Rhesus macaques re-enforced the perception that *P. knowlesi* is not a suitable model for malaria pathophysiology this phenotype is not universal among Rhesus macaques, as Butcher and Mitchell point out in this issue (Butcher and Mitchell, 2016). Originally there were several *P. knowlesi* – Rhesus macaque models and early research described variable disease phenotypes depending on the experimental *P. knowlesi* line used and the origin of the Rhesus monkey (Butcher and Mitchell, 2016). With time, the highly susceptible Rhesus of Indian origin became the experimental host of choice, and information on diverse disease phenotypes was effectively lost. Furthermore, not all of the earlier laboratory adapted lines of *P. knowlesi* survived. The Indian macaque partnered with a few experimental lines of *P. knowlesi* was established and resulted in an homogenized model system that did not represent severe falciparum malaria. Taken together with the positioning of *P. knowlesi* relatively close to *P. vivax* in the *Plasmodium* phylogenies, the notion that *P. knowlesi* could not model for severe falciparum malaria became ingrained.

THE PROBLEM WITH DOGMAS

Perceived wisdom can be difficult to change. Even reports of a severe disease phenotype in Olive baboons, experimentally infected with *P. knowlesi*, with compelling pathological features similar to severe *P. falciparum* malaria with coma and latterly placental malaria in the same model, did not sway the accepted opinion that *P. knowlesi* is a more representative model for *P. vivax* (Ozwara *et al.* 2003; Onditi *et al.* 2015).

By extension of the same logic, leaning on phylogeny, *P. knowlesi* biology is most often compared with *P. vivax*, even though *P. knowlesi* and *P. vivax* bear little morphological or biological resemblance to each other. For example, hypnozoite liver stages, reticulocyte restriction and enlargement of the infected red blood cell, characteristic of *P. vivax* are absent in *P. knowlesi* as are Schuffner's stippling (Coatney *et al.* 1971b). We now know that *P. knowlesi* can cause severe acute malaria in

the human population similar to *P. falciparum* and furthermore both species express antigenically variant surface antigens on the infected host red blood cell (Brown and Brown, 1965; Howard *et al.* 1983). This type of antigenic variation is the product of complex multiple gene families present in both *P. falciparum*, the *var* gene family, and *P. knowlesi*, the *SICAvar* gene family, and absent in *P. vivax* (Baruch *et al.* 1995; Smith *et al.* 1995; Su *et al.* 1995; al-Khedery *et al.* 1999). In *P. falciparum*, the *var* gene family is strongly implicated in virulence (Mosnier and Lavstsen, 2016; Wassmer and Grau, 2017). Although in *P. knowlesi* the role of the likely orthologous *SICAvar* gene family in precipitating human disease is not yet known, the fact that both species cause severe disease and both have this gene family deserves further investigation. Even more compelling, the WHO criteria for severe malaria fit malaria caused by *P. falciparum* and *P. knowlesi*. The important exception is that, so far, patients with severe malaria caused by *P. knowlesi* do not deteriorate into the classical severe malaria with coma disease phenotype, even though parasite-infected erythrocytes were observed in the brain microvasculature of a fatal case of *P. knowlesi* malaria (Fig. 1) (Cox-Singh *et al.* 2010; Barber *et al.* 2017; Daneshvar *et al.* 2017). Nonetheless, positioning in genetic phylogeny appears to carry more weight than clinical and biological evidence. Perhaps we need to re-consider how we identify and rank the characteristics of representative models for malaria pathophysiology.

CHANGING LANDSCAPES AND MOVING ON

Perhaps it is time to consider *P. knowlesi* as a candidate to model for malaria pathophysiology for the following reasons: *P. knowlesi* is an experimental model parasite *in vivo* that is permissive in a number of NHP hosts (Pasini *et al.* 2016); *P. knowlesi* has been recently adapted to *in vitro* culture in human erythrocytes (Moon *et al.* 2013); *P. knowlesi* is highly amenable to genetic manipulation (Kocken *et al.* 2002; Moon *et al.* 2013); clinical and laboratory data from patients with naturally acquired *P. knowlesi* malaria are accumulating, and matched information on genetic and genomic diversity to identify and interrogate parasite virulence *in vitro* and *in vivo* is available (Daneshvar *et al.* 2009; Barber *et al.* 2011, 2013, 2016; Cox-Singh *et al.* 2011; Ahmed *et al.* 2014; Pinheiro *et al.* 2015; Lapp *et al.* 2017). Taken together surely it is time to characterize *P. knowlesi*, including genetically modified parasites, in several NHP hosts to compare with the differing pathologies that are collectively referred to as severe malaria.

All of that being said, is there an appetite among funders or the malaria research community to support the, by no means trivial, development of a

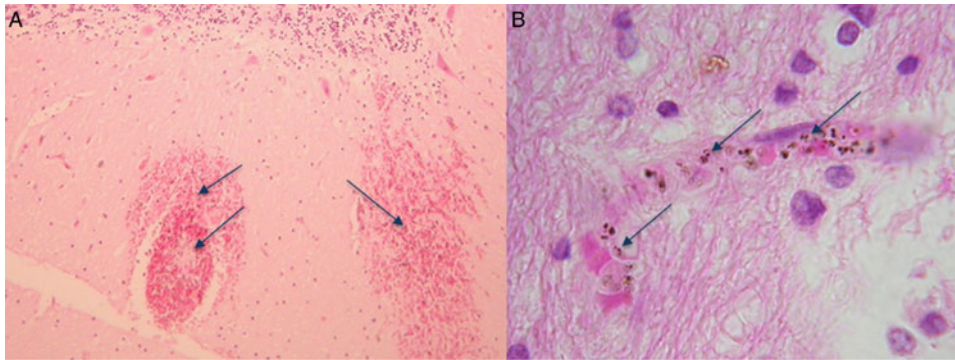


Fig. 1. Haematoxylin and eosin-stained sections from the brain of a fatal case of *Plasmodium knowlesi* malaria. (A) Haemorrhages in white matter containing pigmented parasites (arrows) and (B) at higher magnification showing a capillary congested with pigmented parasite-infected erythrocytes (arrows). The patient was not in coma characteristic of severe falciparum malaria (for detailed descriptions, see Cox-Singh *et al.* 2010).

translational *in vivo* model for malaria pathophysiology? Arguments against this initiative include giving precedence to malaria control rather than defining disease process, an argument further reinforced by the sustainable development goals, Transforming Our World, set by the United Nations in 2016 (<https://sustainabledevelopment.un.org/post2015/transformingourworld>), goals that aim to end the epidemics of infectious disease and neglected tropical diseases, including malaria, by 2030. A target similar to but less specific than that set in 2000 in the millennium development goals with malaria elimination targets set for 2015 (WHO, 2016). Experience suggests that these targets are difficult to achieve and in the interim, while the number of deaths due to malaria have decreased, just under half of the world population remain at risk of malaria (WHO, 2016). Of course for most that risk is negligible, but for some the risk is very real. Particularly at risk are the world's poorest, especially those under the age of 5 years. Perhaps this argument is enough to support research on malaria pathophysiology, including support to develop an *in vivo* translational model.

EMERGING INFECTIONS

A word of caution seems prudent – infectious diseases, including newly emergent infections, tend to be of zoonotic origin. The entry of *P. knowlesi* malaria into the human population in Southeast Asia is a good contemporary example. The entry is on-going and across Southeast Asia. There is no evidence to suggest that malaria caused by *P. knowlesi* is abating and, in terms of research, we remain several steps behind nature. Vythilingam *et al.* this issue highlight just how little is known of the vectors and transmission of *P. knowlesi* (Vythilingam *et al.* 2016). Yakob *et al.*, also this issue, demonstrate the importance of data on vectors to inform mathematical models to describe *P. knowlesi* epidemiology (Yakob *et al.* 2017) Contextually, how feasible is

the goal to end epidemics if they are zoonotic in origin? And yet ambitious goals are set? A case to include zoonotic malaria in the WHO malaria reports to facilitate policy and malaria control strategies in Southeast Asia was made by Barber *et al.* (2017). The need to make a case strongly suggests that zoonoses have perhaps, alarmingly, slipped under the policy makers' radar.

On that backdrop, a dichotomy emerges – funders are willing to support clinical research on severe malaria and research on rodent models for severe disease; yet the same funders are reluctant to engage in discussions on the development of a NHP translational model for severe malaria. Clinical studies collect important data, not least on parasite and host genetic markers of severe disease, markers of immune pathology and immune protection, markers of virulence, yet this work, while valuable, is bottlenecked by the lack of a means to (a) confirm the findings *in vivo*, (b) identify the triggers and pathways to severe disease and (c) translate the results into human health benefit through the rational development of augmentative therapies designed to protect the severely ill through the acute phase of their illness. Translating research on pathophysiology in any disease requires a representative *in vivo* model system. At the risk of repetition, a representative *in vivo* model for severe malaria is not available to perform proof of concept, cause and effective translational research. In the absence of this essential tool, the outputs of clinical and all other research on severe malaria remain and will remain academic. In terms of benefit to communities at risk, money that funds research that cannot be translated could be more beneficial if invested locally to improve access to healthcare, education, safe drinking water and vector control to name but a few.

A POINT OF VIEW

Continued investment in research on severe malaria, which does not include investment in *in vivo* model

development for translational research, is fundamentally flawed. Surely the sole reason to study malaria is to reduce the health impact on communities at risk. If funding and support to develop a vital tool, an *in vivo* model system, is not forthcoming, we need to question support for on-going research on severe malaria. The outputs of current research that are well placed to inform translational research may not be taken forward. If lack of interest and support to develop a translational model persists, then we run the risk of preserving the academic success of the few while failing the many that include the world's poorest who live with the real threat of developing severe life-threatening malaria. Even more frustrating we have the omics revolution at our disposal and a real opportunity to study and interrogate the host–pathogen interface in exquisite detail. Yet without the means to convert the effort into impact-driven research, progress remains grid-locked.

AN OPPORTUNITY

Articles in this special issue highlight the contribution of research on *P. knowlesi* to our current understanding of antigenic variation, parasite invasion of host red blood cells, immunity, drug and vaccine development in malaria (Butcher and Mitchell, 2016; Pasini *et al.* 2016; Galinski *et al.* 2017). Undoubtedly the value of *P. knowlesi* as a laboratory model has waxed and waned over time depending on the needs and particular interests of the research community at any one time. The contemporary finding that *P. knowlesi* has entered the human population from the natural old-world macaque hosts and causes a severe form of malaria with similarity to *P. falciparum* would be expected to heighten interest and application of *P. knowlesi* – the experimental model – to interrogate the many unanswered questions on malaria pathophysiology (Mosnier and Lavstsen, 2016; Wassmer and Grau, 2017). There are obstacles, the number of contemporary research groups with local access to primate research facilities is very small. *Plasmodium knowlesi* while, an interesting parasite, is relatively inaccessible, expensive and ethically more challenging to work with than *P. falciparum* *in vitro* or the *in vivo* rodent models for malaria. Despite the obstacles, the opportunities to exploit omics, genetic manipulation, *in vitro* and *in vivo* experiments that are informed by human disease are huge. *Plasmodium knowlesi* must surely be considered a rational choice for a focused, international initiative to develop, characterize and make available a clinically relevant model system, including a well-matched NHP host, for cause and effect studies on severe malaria (Cox-Singh and Culleton, 2015; Pasini *et al.* 2016; Galinski *et al.* 2017). We have a wealth of information and hypotheses generated from pain-staking clinical and

experimental studies on severe malaria pathophysiology and virulence. Now we need the means to test and translate outputs into health benefit – can *P. knowlesi*, a pioneer parasite in malaria research, help us conquer this difficult frontier?

ACKNOWLEDGEMENTS

In admiration of all those who have worked on *P. knowlesi* as a research tool and to those working in the field informing the research community and policy makers on zoonotic malaria. The author is grateful to the Sasakiwara Foundation for travel support to initiate collaborative discussions on animal model development in Japan.

DISCLAIMER

Any views expressed in this editorial are the personal views of the author and are not endorsed by any other individual or institution.

COVER

Zoonotic malaria transmission – humans are accidental hosts (*Macaca fascicularis* image kindly donated by Cyrus Daneshvar); chromosomal synteny adapted from Figure 2 in Lapp *et al.* (2017) and Giemsa-stained blood film showing mature schizont- and early trophozoite-infected erythrocytes.

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