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Introduction

Four main malaria plasmodia involve human beings. Plasmodium vivax, Plasmodium malariae and Plasmodium ovale had an older evolution paralleling the human species evolution (co-evolution) [1-3], while Plasmodium falciparum has been encountered by humans later during their evolution, since it was acquired by monkeys probably between the end of Mesolithic and the beginning of Neolithic eras [4]. Over twenty species of malaria plasmodia are capable of infecting monkeys (Table 1), but the anecdotal transmission of these organisms to humans has been considered until recently a rare event, with minor public health concerns. The transmission of these plasmodia protozoa predominantly occurred in South-East Asia, China, and Central-Southern America [5]. The expansion of human activities in geographic areas which remained poorly anthropized in the past (i.e. forest zones where productive activities have been recently started), and wild regions, which are increasingly visited by travellers for eco tourism activities, represent objective elements which lead to an expected increase of risk to acquire malaria plasmodia of simian origin. Moreover, the present demographic expansion of local populations in different regions of the Southern Asia and Southern America, which also involved wild areas close to rain forests, especially when villagers carrying out agricultural or breeding activities are interested, bear further epidemiological risk factors. In these environmental contexts, an overlapping of human and simian habitats becomes evident, in presence of a common infestation by Anopheles mosquitoes, which are the needed vehicle for malaria transmission to humans. Among the twenty plasmodia species which may infect monkeys, five of them have been documented as potential infectious agents for humans: they are Plasmodium simium, Plasmodium brasilianum (found in South America), Plasmodium cynomolgi, Plasmodium inui, and Plasmodium knowlesi (retrieved in South-Eastern Asia) [5]. The majority of malaria infection of monkeys tend to cause a mild-to-moderate disease in humans, which is frequently self-limiting, and rarely requires antimalaric chemotherapy, since they are often missed or neglected. The identification of P. knowlesi disease foci confirmed by biomolecular techniques in Sarawak (Malaysian Borneo), Sabah (a state belonging to the Malaysia Confederation), and Pahang (peninsular Malaysia) between the end of the past century and early XXI century [6,7], caught the attention of researchers and clinicians operating in South-Eastern Asia, and more recently of the international health care authorities [5], initially focusing the attention on episodes which presented a typical malaria course, but received a microscopic diagnosis of P. malariae infection. A distinctive feature of these patients was the elevated parasitemia and the severity of clinical course, should a rapid and appropriate treatment was not administered [6,7]; in the records of these authors also four cases with a lethal evolution were present. Later, the availability of specific P. knowlesi primers allowed to pose a definitive diagnosis also in subjects whose erythrocytes proved morphologically similar to those infected by P. malariae, but tested negative when assessed with P. malariae primers. The more severe impact of P. knowlesi malaria is related to the evidence that this Plasmodium has a daily replication cycle and may therefore reach an elevated, life-threatening parasitemia in absence of an effective treatment, whereas P. malariae has a replication cycle every three days (so-called quartan fever), and never reaches elevated blood parasite concentrations [6-8]. Recent entomological studies [9] demonstrated that in the Malaysian geographic context the more efficient vector transmitting P. knowlesi infection to humans is Anopheles latens [10,11] and highest biting rate is registered at the limits of forest (6.74%), while a reduce frequency regards the rain forest (1.85%) and the examined longhouses (the typical Malaysia houses) (0.28%), which never were the site of micro-epidemic disease clusters. Based on these data, it has been hypothesized that humans acquire infection in the forest, when hunting, or returning home at dusk. [10,11]. The potential simian reservoirs are Macaca mulatta, M. nemestina, M. irus, and Saimiri sciurea. After the report of Malaysia foci, where native population was involved, and the identification of single imported cases in Europe and North America by tourists who travelled for short periods close to forest areas of South-Eastern Asia [12-16], the attention of researchers for P. knowlesi rapidly increased. In the year 2008, the Journal "Nature" published the analysis of the nuclear genomic sequencing of P. knowlesi [17]. It was the first complete gene sequencing of a simian Plasmodium, therefore offering the opportunity for a comparison with the complete genome of P. vivax and other already known Plasmodia genomes. In contrast with other plasmodial genomes, putative familiar antigenic variants are scattered throughout the genome and associated with intrachromosomal telomere repeats. One of these families, called KIRs, includes genomic sequences which on the whole match around one-half of the extracellular domain CD99 of the host: this situation suggested that this form could represent an unusual molecular mimicry [17,18]. A relevant information came from recent observations and epidemiological data came from Malaysia. The habitat of humans is more and more overlapping that of primates, and it is primarily due to two aspects: the first is related to the expansion of human activities (agriculture, deforesting, clearing of valued wood, and animal breeding), in areas also inhabited by monkeys. The second emerging issue is related to the relevant increase of touristic activities (i.e. trekking) by international visitors, who spend a proportionally limited time in this environment, and subsequently go back to their countries of origin [8]. In this last situation, the limited knowledge of...
Simian Plasmodium species | Regional distribution | Human species resembling to them
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*P. coatney* | Malaysia, Philippines | *P. falciparum*
*P. cynomolgi* | India, Indonesia, Malaysia, Sri Lanka, Taiwan | *P. vivax*
*P. eylesi* | Malaysia | *P. vivax*
*P. feldi* | Malaysia | *P. ovale*
*P. frangi* | India, Sri Lanka | *P. falciparum*
*P. hylobati* | Indonesia | *P. vivax*
*P. inui* | India, Indonesia, Malaysia, Philippines, Sri Lanka, Taiwan | *P. malariae*
*P. jeffrey* | Indonesia, Malaysia | *P. vivax*
*P. knowlesi* | China, Indonesia, Malaysia, Philippines, Singapore, Thailand, Taiwan | *P. malariae, P. falciparum*
*P. pitheci* | Malaysia | *P. vivax*
*P. simiovale* | Sri Lanka | *P. ovale*
*P. silvaticum* | Malaysia | *P. vivax*
*P. youngi* | Malaysia | *P. vivax*

### Table 1: Malaria species of simian origin isolated in Asia and in Southern America (modified from the reference quotation [4]). Their associated geographical distribution, and the morphologic similarities to one of the four “classical” human plasmodia species (i.e. *Plasmodium falciparum, Plasmodium vivax, Plasmodium ovale*, and *Plasmodium malariae*), are pointed out.

**Figure 1:** The enclosed graph depicts the biological and filogenetical evolutions of *P. vivax, P. malariae, and P. ovale* (Group A), compared with the extremely more recent evolution pathway of *P. falciparum* (group B). *P. knowlesi* (belonging to group C), is the last *Plasmodium* species which steadily entered the human pathology, and is related to productive and touristic human activities carried out in the forest regions of Malaysia and Borneo. The image has been taken and modified from some studies published by L. Capasso (quoted at reference [1]), in particular with the adjunct of group C malaria parasite.
from hominids (group A), toward modern men (group C), with the appearance of *P. falciparum* which entered the epidemiological scenario infecting humans (group B) of mesolithic and early neolithic eras, who spent their lives in cabins located at the edge of rain forest and in the proximity of the first cultured fields. Just the introduction of the first organized human activities which significantly contributed to the expansion of *P. falciparum*, which up to date represents the most relevant malaria pathogen from the pathocenosis of pre-historical populations, unavoidably conditioning both economic and anthropological development. With regard to the appearance of a novel malaria parasite epidemiologically relevant for humans, the hope is that this novel patocenosis remain circumscribed in some countries of South-Eastern Asia, without extension to other areas, which are susceptible due to similar climatic and environmental features. In this evience, the impact of autochtonous populations, not naturally immunized, after the contacts which normally occur during early infancy, may be responsible for particularly severe consequences, not only for local populations lacking of a specific immune memory and exposed to a novel human *Plasmodium* species, but also for the growing number of tourists, who may be exposed to the risk to come back to their countries of origin with a *P. knowlesi* malaria infection, whose diagnosis cannot rely on the routine laboratory examinations.

**References**


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