Natural Human Infections with Plasmodium cynomolgi, P. inui, and 4 other Simian Malaria Parasites, Malaysia

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We detected the simian malaria parasites *Plasmodium knowlesi*, *P. cynomolgi*, *P. inui*, *P. coatneyi*, *P. inui*–like, and *P. simiovale* among forest fringe–living indigenous communities from various locations in Malaysia. Our findings underscore the importance of using molecular tools to identify newly emergent malaria parasites in humans.

oonotic malaria caused by Plasmodium knowlesi, Licommonly found in long-tailed macaques (Macaca fascicularis) and pig-tailed macaques (M. nemestri*na*), is now a major emerging disease, particularly in Malaysia (1,2). Two other simian malaria parasites, P. cynomolgi (2–4) and P. inui (2), have also been shown to have the potential of zoonotic transmission to humans through the bites of infected mosquitoes under natural and experimental conditions. The risk of acquiring zoonotic malaria is highest for persons living at the forest fringe and working or venturing into the forest because of their proximity with the monkey reservoir hosts and the mosquito vectors (5,6). With the aid of molecular methods, we aimed to investigate whether human infections with simian malaria parasites were present among indigenous communities in Malaysia whose villages are situated in the forest or at the forest fringe.

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The Study

We examined 645 archived blood samples that we had collected during 2011–2014 among indigenous populations of various subtribes from 14 villages in 7 states in Malaysia (Appendix Table 1, https://wwwnc. cdc.gov/EID/article/27/8/20-4502-App1.pdf). We first screened the extracted DNA samples at Universiti Malaya (UM) for the presence of Plasmodium with the aid of genus-specific primers (rPLU1 and rPLU5; rPLU3 and rPLU4) (Appendix). Of the 645 indigenous community samples, 102 (15.8%) were positive for Plasmodium. Using species-specific nested PCR assays (Appendix), we identified these infections as monoinfections with *P. knowlesi* (n = 40), *P.* vivax (n = 21), P. cynomolgi (n = 9), P. falciparum (n = 6), P. coatneyi (n = 3), P. inui (n = 3), P. malariae (n = 2), and *P. ovale curtisi* (n = 1) (Table 1). In 17 samples, the species could not be identified despite repeated attempts. Our species-specific primer pairs were designed on the basis of either the asexually (A) or sexually (S) transcribed forms of Plasmodium small subunit (SSU) rRNA genes (7); the genus-specific primer pairs anneal to both asexual and sexual forms of the SSU rRNA genes, and therefore the genus-specific assay is more sensitive.

We further characterized the 55 samples that tested positive for simian malaria parasites by amplifying a longer fragment of the SSU rRNA gene (914 bp-950 bp) for direct sequencing. Phylogenetic analysis using the neighbor-joining method (Figure 1) revealed the presence of *P. knowlesi* (samples PK1-40), *P. coatneyi* (UM1-3), *P. cynomolgi* (UM9, UM11, UM12, UM14, UM15, UM17, UM18), and *P. inui* (UM5-7). Meanwhile, 2 sequences derived from

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¹These authors contributed equally to this article.

<u>intergente de commune</u>	No.	ampiee, sy eta	Human and simian malaria species							
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	samples	No. positive	<i>P</i> .		<i>P</i> .	P. ovale	Р.	<i>P</i> .	<i>P</i> .	
State	tested	samples	falciparum	P. vivax	malariae	curtisi	knowlesi	coatneyi	cynomolgi	P. inui
Pahang	109	5	0	2	0	1	2	0	0	0
Perak	61	55	3	10	2	0	26	3	5	0
Selangor	49	0	0	0	0	0	0	0	0	0
Negeri Sembilan	163	13	1	2	0	0	2	0	2	0
Melaka	32	13	2	3	0	0	1	0	1	1
Kelantan	32	9	0	2	0	0	6	0	1	0
Sarawak	199	7	0	2	0	0	3	0	0	2
Total/overall	645	102† (of	6 (of 102;	21	2	1	40	3	9	3
prevalence		645; 15.8%)	5.9%)	(20.6%)	(2.0%)	(1.0%)	(39.2%)	(2.9%)	(8.8%)	(2.9%)
*0.011 // // //										

Table 1. Human and simian *Plasmodium* malaria species identified by nested PCR at UM targeting SSU rRNA genes among indigenous community blood samples, by state, Malaysia*

*SSU, small subunit; UM, Universiti Malaya.

†102 of 645 (15.8%) indigenous community samples were found positive with *Plasmodium* genus-specific primers; 17 *Plasmodium* genus-positive samples could not be identified up to species level despite repeated attempts.

samples UM10 and UM16 were found to be closely related to *P. simiovale*.

We then reextracted DNA from 15 blood samples that were positive for *P. coatneyi*, *P. cynomolgi*, and *P. inui* and sent these samples (blinded) together with 5 Plasmodium-negative samples to Universiti Malaysia Sarawak (UNIMAS) to confirm their identities by PCR and sequencing of part of the cytochrome c oxidase subunit 1 (COX1) gene. At UNIMAS, using nested PCR assays based on SSU rRNA genes, we found 1 single and 9 double species infections. We could not identify the species of Plasmodium for sample UM6, 4 of the Plasmodium-positive samples from UM were Plasmodium negative, and all 5 Plasmodium-negative samples from UM (UM4, 8, 13, 19, 20) tested negative (Table 2). Furthermore, because both laboratories at UM and UNIMAS had previously extracted DNA from macaque blood to examine for simian malaria parasites, we tested the samples for macaque DNA to rule out the possibility that the simian malaria

parasites detected were the result of contamination with macaque blood. We obtained negative results using nested PCR for detection of macaque DNA for the 20 DNA samples when they were first received at UNIMAS and also when we repeated testing after completing the sequencing of COX1 genes, indicating that these samples were not contaminated with macaque blood upon receipt or during subsequent experiments at UNIMAS.

We then subjected the PCR-positive samples (UM6-7, UM9-12, UM14-18) to amplification and sequencing of partial COX1 genes. Neighbor-joining (Figure 2) phylogenetic inference of these sequences, together with available referral sequences from Gen-Bank, indicated that 32 haplotypes from samples UM9-12 and UM14-18 were genetically indistinguishable from *P. cynomolgi*. Our phylogenetic analyses also demonstrated that sample UM7 had a single infection with *P. inui*-like parasites, whereas UM6 had a double infection with *P. simiovale* and *P. inui*-like

Table 2. Comparison between results of nested PCR and sequencing at UM and UNIMAS for identification of *Plasmodium* malaria species from indigenous community blood samples, Malaysia*

	Identifica	ation at UM	Identification at UNIMAS		
	PCR assays based on	Phylogenetic analysis of	PCR assays based on	Phylogenetic analysis of	
Sample ID	SSU rRNA genes	SSU rRNA genes	SSU rRNA genes	COX1 genes	
UM1	P. coatneyi	P. coatneyi	Negative	ND	
UM2	P. coatneyi	P. coatneyi	Negative	ND	
UM3	P. coatneyi	P. coatneyi	Negative	ND	
UM5	P. inui	P. inui	Negative	ND	
UM6	P. inui	P. inui	Positive	P. inui–like, P. simiovale	
UM7	P. inui	P. inui	P. inui	P. inui-like	
UM9	P. cynomolgi	P. cynomolgi	P. cynomolgi, P. inui	P. cynomolgi	
UM10	P. cynomolgi	Plasmodium spp.	P. cynomolgi, P. inui	P. cynomolgi	
UM11	P. cynomolgi	P. cynomolgi	P. cynomolgi, P. inui	P. cynomolgi	
UM12	P. cynomolgi	P. cynomolgi	P. cynomolgi, P. inui	P. cynomolgi	
UM14	P. cynomolgi	P. cynomolgi	P. cynomolgi, P. inui	P. cynomolgi	
UM15	P. cynomolgi	P. cynomolgi	P. cynomolgi, P. inui	P. cynomolgi	
UM16	P. cynomolgi	Plasmodium spp.	P. cynomolgi, P. inui	P. cynomolgi, P. inui–like, P. simiovale	
UM17	P. cynomolgi	P. cynomolgi	P. cynomolgi, P. inui	P. cynomolgi	
UM18	P. cynomolgi	P. cynomolgi	P. cynomolgi, P. inui	P. cynomolgi	

*Negative, negative for *Plasmodium* DNA and not examined by species-specific nested PCR assays; ND, not done; positive, positive for *Plasmodium* DNA but negative with species-specific nested PCR assays. SSU, small subunit; UM, Universiti Malaya; UNIMAS, Universiti Malaysia Sarawak.

parasites and UM16 had a triple infection with *P. cy-nomolgi*, *P. simiovale*, and *P. inui*-like parasites.

We generated phylogenetic trees of similar topology by the maximum-likelihood method for the SSU rRNA genes (Appendix Figure 1) and by the Bayesian maximum clade credibility method for the COX1 genes (Appendix Figure 2). There were discrepancies between the nested PCR assay results and the sequencing results between our 2 laboratories; mixed species of *Plasmodium* were identified only at UNI-MAS. A possible explanation is that the DNA samples analyzed at UNIMAS were newly extracted and were different from the ones used in the experiments at UM. There might also be a compromise of the sensitivity in detecting the species with lower parasitemia in mixed infections as a result of competition for nest 1 primers by the species with higher parasite loads. Furthermore, for sequencing of the SSU rRNA genes at UM, primers that were specific for the species identified by nested PCR assays were used, whereas for the COX1 genes, both P. cynomolgi-specific primers and primers that could amplify other species of Plas*modium* were used. Therefore, additional species of Plasmodium were identified at UNIMAS in these samples, such as P. simiovale and P. inui-like, for which no species-specific PCR primers exist.

Conclusions

The 40 *P. knowlesi* infections we detected originated from 6 states in Malaysia, thereby confirming the widespread distribution of human *P. knowlesi* malaria cases in Malaysia (1). We detected *P. cynomolgi* infections among indigenous communities in 4 states in Malaysia. Taken together with previous reports of naturally acquired *P. cynomolgi* infections in humans in the states of Terengganu, Sabah, and Sarawak (3,8,9), our findings indicate that human infections caused by *P. cynomolgi* are also widely distributed in Malaysia.

Our study highlights the occurrence of naturally acquired human infections with *P. inui*, *P. inui*-like, *P. coatneyi*, and *P. simiovale*. Natural human *P. inui* infections have not been described (10), although the parasite is experimentally transmissible to humans (2). For *P. coatneyi*, attempts to infect humans with blood from an infected rhesus monkey and through infected mosquitoes were unsuccessful (2). *P. simiovale* is a lesser-studied simian malaria parasite that was previously described only in toque macaques (*Macaca sinica*) of Sri Lanka (2) until it was recently identified, together with *P. inui*-like parasites, in long-tailed macaques from Sarawak in Malaysian Borneo (11). All these simian malaria parasites would have been diagnosed by microsco-









py as human malaria parasites because they share morphological similarities with human malaria parasites. The early blood stages of P. knowlesi resemble those of *P. falciparum*, and the other forms are similar to P. malariae (2,6). P. cynomolgi is morphologically similar to P. vivax (2), and both P. inui and *P. inui*-like parasites are morphologically identical to P. malariae (2,11), whereas P. coatneyi bears morphologic similarities to P. falciparum and P. simiovale bears morphologic similarities to P. ovale (2,12). Besides misdiagnosis of simian malaria parasites as human malaria parasites, there are other limitations of microscopy for diagnosis of malaria; thus, using molecular tools is paramount in generating accurate epidemiology data (6). It is envisaged that screening with molecular tools of other communities living at the forest fringes will demonstrate the widespread distribution of zoonotic malaria and uncover more newly emergent malaria parasites.

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Natural Human Infections with *Plasmodium cynomolgi*, *P. inui*, and 4 other Simian Malaria Parasites, Malaysia

Appendix

Materials and Methods

Ethical Considerations

The study was approved by the Medical Research Ethics Committee of University of Malaya Medical Centre (MEC Ref. No. 920.83). Approval was also obtained from the Department of Orang Asli (Indigenous) Development and the respective heads of the villages before blood sample collection from the indigenous communities. We obtained informed consents from those who agreed to participate, or from parents on behalf of their children.

Source of Archived Blood Samples

We examined 645 archived blood samples that we had collected during 2011–2014 among indigenous populations of various subtribes from 14 villages in 7 states of Malaysia: Pahang, Perak, Selangor, Negeri Sembilan, Melaka, Kelantan, and Sarawak (Appendix Table 1). These indigenous community samples were obtained during previous studies focusing on intestinal parasites. Therefore, information such as body temperature, malaria history, and malaria parasite density were not available.

The indigenous communities we studied here are a diverse group. There are ≥ 95 subgroups distributed in selected states throughout Malaysia, each with its own distinct language and culture. The indigenous population of peninsular Malaysia is separated into 3 main tribal groups, Negrito, Senoi, and Proto Malay (Aboriginal Malay), and consists of 18 subtribes. The largest indigenous groups in Malaysian Borneo are Ibans in Sarawak and the Kadazan Dusuns in Sabah. The indigenous communities that we studied all live in the forest fringe and are engaged with forest and agricultural activities in which there is a greater chance of being exposed to the macaque reservoirs and mosquito vectors (*1,2*).

Molecular Detection of Plasmodium Species at Universiti Malaya (UM)

We extracted genomic DNA from either blood (\approx 3 mL) or blood spots on filter paper using the QIAamp DNA Blood Mini Kit (QIAGEN, https://www.qiagen.com), according to the manufacturer's instructions, and stored the samples at -20° C until further analysis. We first screened the DNA samples at UM for the presence of *Plasmodium* with the aid of genus-specific primers (rPLU1, rPLU5, rPLU3 and rPLU4), as described previously (*3*). We then examined *Plasmodium*-positive samples by nested PCR assays using species-specific primers for *Plasmodium falciparum*, *P. vivax*, *P. malariae*, *P. ovale curtisi*, *P. ovale wallikeri* (*3*), *P. knowlesi* (*4*), *P. coatneyi*, *P. cynomolgi*, *P. inui*, and *P. fieldi* (*5*).

Amplification and Sequencing of SSU rRNA Genes of Plasmodium Species at UM

We amplified and sequenced longer fragments of SSU rRNA genes of simian *Plasmodium* species (914–950 bp) by nested PCR assays with other pairs of species-specific primers (*6*). We performed PCR amplifications in a 50 µL reaction volume consisting of 5 µL DNA template from previously amplified PCR product, 1X PCR buffer (Promega, https://www.promega.com), 0.2 mM dNTPs, 3 mM MgCl₂, 1.5 U Taq DNA polymerase, and 0.5 µM forward and reverse primers. The PCR was carried out in a MyCycler Thermal Cycler (Bio-Rad, https://www.bio-rad.com) under the following conditions: 94°C for 5 min; 35 cycles of 94°C for 1 min, 50–60°C for 90 sec, 72°C for 1 min; 72°C for 10 min. We examined all PCR products (1015–1050 bp) using 1.5% agarose gels before we sent amplicons to a commercial facility for bidirectional sequencing (BigDye Terminator v.3.1 chemistry; Applied Biosystems, https://www.thermofisher.com).

Molecular Detection of Simian Plasmodium Species at Universiti Malaysia Sarawak (UNIMAS)

We subsequently extracted DNA from 15 blood samples we had identified as having *P*. *cynomolgi*, *P*. *coatneyi* and *P*. *inui*, and 5 samples that were malaria-negative at UM. We then sent these samples blind to Universiti Malaysia Sarawak (UNIMAS), where they were first examined by nested PCR assays for *Plasmodium*, and the *Plasmodium*-positive ones were examined with species-specific primers as described previously (3,5,7).

PCR Amplification and Sequencing COX1 Genes at UNIMAS

Sequencing of the partial COX1 genes of *Plasmodium* involved a single-step PCR or a hemi-nested PCR. We amplified 3 samples (UM10, UM11, UM14) with single-step PCR and 3

(UM6, UM7, UM18) with hemi-nested PCR; we used both methods for 4 (UM9, UM12, UM15, UM16).

In the hemi-nested PCR, we amplified the complete COX1 gene using *Plasmodium*specific primers: CYFinF1 (5'-CCTGACATGGATGGATAATACTCG-3') and CYFinR2 (5'-CCATCCATTTAAAGCGTCTGG-3'). We performed Nest 1 PCR amplification in a 50 μ L reaction mixture containing 1× Colorless GoTaq PCR buffer, 2.5 mmol of MgCl₂, 0.2 mmol dNTP mix (Promega, https://www.promega.com), 0.025 U GoTaq DNA polymerase, 0.25 μ mol of each primer (CYFinF1 and CYFinR2), and 5 μ L of purified genomic DNA under the following conditions: 94°C for 4 min; 30 cycles of 94°C for 1 min, 53°C for 1 min, and 72°C for 90 sec; 5 min at 72°C. We then used the Nest 1 amplicons as a template for the hemi-nested PCR assay with *P. cynomolgi*-specific primers: cox1_F1 (5'-CCAAGCCTCACTTATTGTTAAT-3') (8) and CYFinR2 and *Plasmodium*-specific primers: CYFinF3 (5'-

CCAAAGTATAACCGCTGTCGC-3') and CYFinR2. We performed the hemi-nested PCR amplification for each sample in a 30 μ L reaction mixture containing 1× HF colorless PCR buffer, 0.2 mmol dNTP mix, 0.02 U Phusion Polymerase (Promega), 0.5 μ mol of each primer (cox1_F1 and CYFinR2 or CYFinF3 and CYFinR2), and 3 μ L of Nest 1 product under the following conditions: 98°C for 30 sec; 35 cycles at 98°C for 7 sec, 60°C (for cox1_F1 and CYFinR2) and 62°C (for CYFinF3 and CYFinR2) for 20 sec, and 72°C for 22 sec; and 72°C for 10 min.

We performed single-step PCR amplification of *P. cynomolgi* COX1 fragment using *P. cynomolgi*-specific primers: $cox1_F1$ (5'- CCAAGCCTCACTTATTGTTAAT-3') and $cox1_R1$ (5'- ACCAAATAAAGTCATTGTTGATCC-3') (8). We performed amplifications in a 30 µL reaction mixture containing similar concentrations of PCR master-mix components with $cox1_F1$ and CYFinR2 or with CYFinF3 and CYFinR2 primers and 3 µL of purified genomic DNA as the template, using the following parameters: 98°C for 30 sec; 35 cycles at 98°C for 7 sec, 58°C for 20 sec, and 72°C for 28 sec; and 72°C for 10 min.

We performed *Plasmodium sp.* DNA cloning and transformation of the recombinant plasmids using the Zero Blunt TOPO PCR Cloning Kit, with One Shot TOP10 Chemically Competent *E. coli* cells (Invitrogen, https://www.thermofisher.com). We extracted plasmid DNA

using the PureLink Quick Plasmid DNA Miniprep Kit (Invitrogen) and sent plasmids to a commercial facility for bidirectional DNA sequencing.

Phylogenetic Analysis

We trimmed and aligned the SSU rRNA sequences of *Plasmodium* species using the Geneious version 9.1.6 software (9). We constructed phylogenetic trees using the neighborjoining method as described in MEGA v10.0.5 software (10) with bootstrap percentage based on 1,000 replications. We deposited the sequences in GenBank under accession nos. MK351344– MK351383, MK351405–MK351407, MK351409–MK351417, and MK351420–MK351422 (Appendix Table 2).

We used ClustalX v2 to align the partial COX1 sequences. We inferred phylogenetic relationships using the neighbor-joining method (*11*) implemented in MEGA v10.0.5. We reconstructed the neighbor-joining tree with 1,000 bootstrap percentage based on 1,000 replications. We used Tree Annotator to annotate the tree generated by BEAST (https://www.mybiosoftware.com) and visualized the maximum clade credibility tree using FigTree v1.3.1 (https://figtree-1-3-1.software.informer.com). We deposited the *Plasmodium* COX1 sequences generated in GenBank under accession nos. MT992662-MT992702 (Appendix Table 3).

Molecular Detection of Macaque DNA in the Human Samples at UNIMAS

We screened all 20 samples sent from UM to UNIMAS for the presence of macaque DNA. We amplified the cytochrome c oxidase subunit 1 (COX1) gene by PCR using *Macaca* genus-specific primers MacF (5'-CAACGTYATYGTAACGGC-3') and MacR (5'-AGGTAGTATTGAGGTTGC-3'). We performed Nest 1 PCR amplification for each sample using the Applied Biosystems ProFlex PCR System thermocycler (Thermo Fisher Scientific, https://www.thermofisher.com) in a 20 μ L reaction mixture containing 1× colorless GoTaq PCR buffer (Promega), 2 mmol of MgCl₂, 0.2 mmol dNTP mix, 0.25 μ mol of each primer (MacF and MacR), 0.025 U GoTaq DNA polymerase, and 2 μ L of purified genomic DNA under the following conditions: 94°C for 4 min; 35 cycles of 94°C for 30 sec, 59°C for 1 min, and 72°C for 30 sec; and 72°C for 5 min. We used *M. fascicularis*-specific primers MfF (5'-AGGGTTCGGGAACTGACTG-3') and MfR (5'-TGATCAGACAAATAAAGGGGTC-3') and *M. nemestrina*-specific primers MnF (5'-CATACCTATTATGATTGGGGGT-3') and MnR (5'-

GGTGGAGGAGAAGATGATTAGG-3') for subsequent PCR amplification in a 20 μ L reaction mixture containing 1× Green GoTaq PCR buffer (Promega), 2 mmol of MgCl₂, 0.2 mmol dNTP mix, 0.25 μ mol of each primer (MfF and MfR or MnF and MnR), and 0.025 U GoTaq DNA polymerase with 2 μ L of Nest 1 product under the following conditions: 94°C for 4 min; 35 cycles of 94°C for 30 sec, 57°C for 1 min, and 72°C for 30 sec; and 72°C for 5 min.

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State	District	Village	Subtribe	n (%)	Reference
Peninsular Mala	iysia				
Pahang	Pekan	Chini	Proto-Malay (Jakun)	9 (1.4)	Unpublished data
-	Temerloh	Paya Sendayan	Senoi (Jahut)	97 (15.0)	(12)
	Lanchang	Kuala Gandah	Senoi (Che Wong)	3 (0.5)	Unpublished data
Perak	Slim River	Sungail Bil	Senoi (Semai)	40 (6.2)	Unpublished data
	Tapah	Batu 7 1/2	Senoi (Semai)	7 (1.1)	Unpublished data
		Batu 8	Senoi (Semai)	14 (2.2)	Unpublished data
Selangor	Semenyih	Donglai Baru	Proto-Malay (Temuan)	49 (7.6)	(12)
Negeri	Jelebu	Dusun Kubur	Proto-Malay (Temuan)	100 (15.5)	(13)
Sembilan		Ulu Kelaka	Proto-Malay (Temuan)	63 (9.8)	(13)
Melaka	Alor Gajah	Bukit Sebang	Proto-Malay (Temuan)	9 (1.4)	(12)
		Bukit Payung	Proto-Malay (Temuan)	23 (3.6)	(12)
Kelantan	Gua Musang	Kuala Lah	Negrito (Mendriq)	15 (2.3)	Unpublished data
	•	Aring 5	Negrito (Bateq)	17 (2.6)	Unpublished data
Malaysia Borneo	0	Ū			-
Sarawak	Sarikei	Pakan	Iban	199 (30.9)	(14)

Appendix Table 1. Distribution of indigenous community blood samples used in the study of *Plasmodium* infections in Malaysia, according to state, district, village, and subtribe (N = 645).

Appendix Table 2. GenBank accession numbers of partial sequence SSU rRNA gene generated from simian *Plasmodium* species found in study at Universiti Malaya

GenBank accession no.	Plasmodium spp.	Location	Sample ID
MK351344	P. knowlesi	Pakan, Sarikei, Sarawak	PK1
MK351345	P. knowlesi	Pakan, Sarikei, Sarawak	PK2
MK351346	P. knowlesi	Pakan, Sarikei, Sarawak	PK3
MK351347	P. knowlesi	Kg Kuala Gandah, Lanchang, Pahang	PK4
MK351348	P. knowlesi	Kg Chini, Pekan, Pahang	PK5
MK351349	P. knowlesi	Kg Ulu Kelaka, Jelebu, Negeri Sembilan	PK6
MK351350	P. knowlesi	Kg Ulu Kelaka, Jelebu, Negeri Sembilan	PK7
MK351351	P. knowlesi	Kg Sungai Bil, Slim River, Perak	PK8
MK351352	P. knowlesi	Kg Sungai Bil, Slim River, Perak	PK9
MK351353	P. knowlesi	Kg Sungai Bil, Slim River, Perak	PK10
MK351354	P. knowlesi	Kg Sungai Bil, Slim River, Perak	PK11
MK351355	P. knowlesi	Kg Sungai Bil, Slim River, Perak	PK12
MK351356	P. knowlesi	Kg Sungai Bil, Slim River, Perak	PK13
MK351357	P. knowlesi	Kg Sungai Bil, Slim River, Perak	PK14
MK351358	P. knowlesi	Kg Sungai Bil, Slim River, Perak	PK15
MK351359	P. knowlesi	Kg Sungai Bil, Slim River, Perak	PK16
MK351360	P. knowlesi	Kg Sungai Bil, Slim River, Perak	PK17
MK351361	P. knowlesi	Kg Sungai Bil, Slim River, Perak	PK18
MK351362	P. knowlesi	Ko Sungai Bil. Slim River, Perak	PK19
MK351363	P. knowlesi	Kg Sungai Bil, Slim River, Perak	PK20
MK351364	P. knowlesi	Kg Sungai Bil, Slim River, Perak	PK21
MK351365	P. knowlesi	Kg Sungai Bil, Slim River, Perak	PK22
MK351366	P. knowlesi	Ko Batu 7 1/2, Tapah, Perak	PK23
MK351367	P. knowlesi	Kg Batu 7 1/2, Tapah, Perak	PK24
MK351368	P. knowlesi	Kg Batu 7 1/2, Tapah, Perak	PK25
MK351369	P. knowlesi	Kg Batu 7 1/2, Tapah, Perak	PK26
MK351370	P. knowlesi	Kg Batu 8, Tapah, Perak	PK27
MK351371	P. knowlesi	Kg Batu 8, Tapah, Perak	PK28
MK351372	P. knowlesi	Kg Batu 8, Tapah, Perak	PK29
MK351373	P. knowlesi	Kg Batu 8, Tapah, Perak	PK30
MK351374	P. knowlesi	Kg Batu 8, Tapah, Perak	PK31
MK351375	P. knowlesi	Kg Batu 8, Tapah, Perak	PK32
MK351376	P. knowlesi	Kg Batu 8, Tapah, Perak	PK33
MK351377	P. knowlesi	Kg Bukit Sebang, Alor Gaiah, Melaka	PK34
MK351378	P. knowlesi	Kg Kuala Lah, Gua Musang, Kelantan	PK35
MK351379	P. knowlesi	Kg Kuala Lah, Gua Musang, Kelantan	PK36
MK351380	P. knowlesi	Kg Kuala Lah, Gua Musang, Kelantan	PK37
MK351381	P. knowlesi	Kg Aring 5, Gua Musang, Kelantan	PK38
MK351382	P. knowlesi	Kg Aring 5, Gua Musang, Kelantan	PK39
MK351383	P. knowlesi	Kg Aring 5, Gua Musang, Kelantan	PK40
MK351405	P. coatneyi	Kg Sungai Bil, Slim River, Perak	UM1
MK351406	P. coatnevi	Kg Batu 7 1/2, Tapah, Perak	UM2
MK351407	P. coatnevi	Kg Batu 8, Tapah, Perak	UM3
MK351409	P. cynomolai	Kg Sungai Bil, Slim River, Perak	UM9
MK351410	P. cynomolgi	Kg Sungai Bil, Slim River, Perak	UM11
MK351411	P. cynomolgi	Kg Sungai Bil, Slim River, Perak	UM15
MK351412	P. cynomolgi	Kg Batu 8, Tapah, Perak	UM17
MK351413	P. cynomolgi	Kg Batu 8, Tapah, Perak	UM12
MK351414	P. cynomolai	Kg Ulu Kelaka, Jelebu, Negeri Sembilan	UM14
MK351415	Plasmodium spp.	Kg Ulu Kelaka, Jelebu, Negeri Sembilan	UM16
MK351416	P. cynomolai	Kg Bukit Sebang, Alor Gaiah, Melaka	UM18
MK351420	P. inui	Kg Bukit Sebang, Alor Gajah, Melaka	UM5
MK351421	P. inui	Pakan, Sarikei, Sarawak	UM6
MK351422	P. inui	Pakan, Sarikei, Sarawak	UM7

Appendix Table 3. GenBank accession numbers of partial sequence COX1 gene generated from simian Plasmodi	um species
found in study at Universiti Malaysia Sarawak	

GenBank accession no.	Plasmodium spp.	Location	Clone identity*
MT992662	P. cf. inui	Pakan, Sarikei, Sarawak	UM6A_54
MT992663	P. simiovale	Pakan, Sarikei, Sarawak	UM6D
MT992664	P. cf. inui	Pakan, Sarikei, Sarawak	UM7B
MT992665	P. cynomolgi	Kg Sungai Bil, Slim River, Perak	UM9A_1N
MT992666	P. cynomolgi	Kg Sungai Bil, Slim River, Perak	UM9A_2N
MT992667	P. cynomolgi	Kg Sungai Bil, Slim River, Perak	UM9A_2
MT992668	P. cynomolgi	Kg Sungai Bil, Slim River, Perak	UM9A 20
MT992669	P. cynomolgi	Kg Sungai Bil, Slim River, Perak	UM9A_21
MT992670	P. cynomolgi	Kg Sungai Bil, Slim River, Perak	UM9B_66
MT992671	Plasmodium sp.	Kg Sungai Bil, Slim River, Perak	UM9B_75
MT992672	P. cynomolgi	Kg Aring 5, Gua Musang, Kelantan	UM10A 5
MT992673	P. cynomolgi	Kg Aring 5, Gua Musang, Kelantan	UM10A_9
MT992674	P. cynomolgi	Kg Sungai Bil, Slim River, Perak	UM11A_7
MT992675	P. cynomolgi	Kg Sungai Bil, Slim River, Perak	UM11B_8
MT992676	P. cynomolgi	Kg Sungai Bil, Slim River, Perak	UM11B 10
MT992677	P. cynomolgi	Kg Sungai Bil, Slim River, Perak	UM11B_13
MT992678	P. cynomolgi	Kg Batu 8, Tapah, Perak	UM12A 2
MT992679	P. cynomolgi	Kg Batu 8, Tapah, Perak	UM12B_13N
MT992680	P. cynomolgi	Kg Batu 8, Tapah, Perak	UM12B_14N
MT992681	P. cynomolgi	Kg Batu 8, Tapah, Perak	UM12C_107
MT992682	P. cynomolgi	Kg Ulu Kelaka, Jelebu, Negeri Sembilan	UM14A 2
MT992683	P. cynomolgi	Kg Ulu Kelaka, Jelebu, Negeri Sembilan	UM14B_28
MT992684	P. cynomolgi	Kg Sungai Bil, Slim River, Perak	UM15A_2
MT992685	P. cynomolgi	Kg Sungai Bil, Slim River, Perak	UM15A_30
MT992686	Plasmodium sp.	Kg Sungai Bil, Slim River, Perak	UM15B_9
MT992687	P. cynomolgi	Kg Sungai Bil, Slim River, Perak	UM15B_11
MT992688	Plasmodium sp.	Kg Sungai Bil, Slim River, Perak	UM15B_12
MT992689	Plasmodium sp.	Kg Sungai Bil, Slim River, Perak	UM15B_13
MT992690	P. cynomolgi	Kg Sungai Bil, Slim River, Perak	UM15B_14
MT992691	P. cynomolgi	Kg Sungai Bil, Slim River, Perak	UM15B_24
MT992692	P. cynomolgi	Kg Sungai Bil, Slim River, Perak	UM15B_28
MT992693	P. cynomolgi	Kg Sungai Bil, Slim River, Perak	UM15B_34
MT992694	P. cynomolgi	Kg Sungai Bil, Slim River, Perak	UM15B_39
MT992695	P. simiovale	Kg Ulu Kelaka, Jelebu, Negeri Sembilan	UM16A 1
MT992696	P. cf. inui	Kg Ulu Kelaka, Jelebu, Negeri Sembilan	UM16A_9
MT992697	P. cynomolgi	Kg Ulu Kelaka, Jelebu, Negeri Sembilan	UM16B_18
MT992698	P. cynomolgi	Kg Ulu Kelaka, Jelebu, Negeri Sembilan	UM16B_21
MT992699	P. cynomolgi	Kg Ulu Kelaka, Jelebu, Negeri Sembilan	UM16C_12
MT992700	P. cynomolgi	Kg Batu 8, Tapah, Perak	UM17B
MT992701	P. cynomolgi	Kg Bukit Sebang, Alor Gajah, Melaka	UM18A
MT992702	P. cynomolgi	Kg Bukit Sebang, Alor Gajah, Melaka	UM18B

*Clone identity is the identity of the clones derived from samples UM 6–7, UM 9–12, and UM 14–18.



Appendix Figure 1. Maximum-likelihood phylogenetic tree of *Plasmodium* species, based on partial sequence of SSU rRNA genes. Numbers at nodes indicate percentage support of 1,000 bootstrap replicates; only bootstrap values above 70% are displayed. Nucleotide sequences generated from our study are marked with asterisks and are in bold. Scale bar indicates branch length.



Appendix Figure 2. Maximum clade credibility phylogeny of *Plasmodium* species based on partial sequence of COX1 genes inferred using the Bayesian method. Numbers on branches are values of posterior probabilities. Sequences of *P. cynomolgi* are highlighted in yellow, *P. simiovale* in green, and *P. inui*-like in red. Nucleotide sequences generated from the present study are marked with asterisks.