We detected 2 natural, asymptomatic *Plasmodium inui* monoinfections in humans in Malaysia by using nested PCR on concentrated high-volume blood samples. We found a *P. inui*–positive *Anopheles cracens* mosquito in the same site as the human infections. Investigators should use ultrasensitive detection methods to identify simian malaria parasite transmission in humans.

**Zoonotic transmission of simian malaria parasites to humans have been occurring in Southeast Asia and South America. Among the 3 simian malaria parasites in Southeast Asia experimentally shown to infect humans (1), *Plasmodium knowlesi*, *P. cynomolgi*, and *P. inui*, only *P. knowlesi* and *P. cynomolgi* have been reported in cases of natural infection (2). We report 2 natural, asymptomatic *P. inui* human infections detected by using nested PCR (nPCR) on concentrated high-volume blood.

**The Study**

We conducted an epidemiologic and entomologic study at a campsite in Kem Sri Gading, Pahang, Malaysia (3°45′46.24″N, 102°34′20.32″E), because of frequent reports of human *P. knowlesi* infections acquired from this area. Kem Sri Gading is a receptive area, a location in which the ecosystem permits malaria transmission because vector and reservoir host populations both inhabit it.

On March 2, 2020, we obtained <3 mL of venous blood from 71 persons at the camp who provided consent. Participants had undergone training in the forest at Kem Sri Gading during January 27–28, 2020. The Medical Research and Ethics Committee, Ministry of Health Malaysia, approved this study (approval no. NMRR-15-672-23975 for the human study and approval no. NMRR-19-962-47606 for the mosquito study).

The 2 case-patients we report, PMAR0041, a 20-year-old woman, and PMAR0052, a 19-year-old woman, had no previous history of malaria. Before our study, PMAR0041 was in a nonreceptive city in Selangor 1–2 weeks before training at the camp; PMAR0052 regularly entered forested areas ≥2 times per month. During January 29–March 2, 2020, neither case-patient visited any potentially receptive areas. Both persons reported they were healthy before, during, and after blood collection.

Using the amount of DNA equivalent to 500 µL of whole blood (3), we detected *Plasmodium* in the 2 cases in separate nPCR assays (Appendix, https://wwwnc.cdc.gov/EID/article/27/10/21-0412-App1.pdf). We used primers targeting both the asexual and sexual 18S rRNA genes of *Plasmodium* (4). Sequence analysis of the cloned genus PCR products confirmed *P. inui* (Table). We performed species-specific nPCR assays to detect 5 known human malaria parasites, including *P. knowlesi*, and to detect *P. cynomolgi* and *P. inui*, by using previously published primers (4–6). However, the species-specific PCR amplification demonstrated spurious results; we were unable to produce consistent results over repeated tests. Thus, *P. inui* was detected only in case-patient PMAR0041 (Figure 1) because the protocol produced insufficient DNA, which hampered further analyses. However, we found likely trophozoites in thick blood smears of each case during 2 hours of observation (Figure 2).

On October 9, 2020, we obtained a second blood sample from case-patient PMAR0041; case-patient PMAR0052 did not consent to a second blood collection. Between the first and follow-up blood collections, PMAR0052 did not travel to any receptive areas. We did not detect any *Plasmodium* DNA in the second blood sample from PMAR0041 after repeated tests.
We collected *Anopheles craccens*, *An. intortatus*, and *An. barbirostris* sensu lato mosquitoes at the camp by using human landing catches and Mosquito Magnet Independence Trap (Woodstream Corp., https://www.woodstream.com). *An. craccens* was the predominant mosquito species collected. Only 1 nonblood fed *An. craccens* mosquito, caught on August 24, 2020, was *Plasmodium*-positive in its head and thorax by nPCR (4). We found no oocysts upon dissection of the mosquito gut. We were unable to successfully dissect the salivary glands because the mosquito was dead. We used published primers (7) to amplify the *P. inui* 18S rRNA gene and confirmed *P. inui* by sequencing the PCR product (Table). We tested the entomological team by using the same PCR methods described for the case-patients but detected no *Plasmodium*.

Our analyses showed that the *P. inui* sequence obtained from case-patient PMAR0041 was identical to the corresponding region on the asexual type 18S rRNA sequence obtained from the *An. craccens* mosquito (Appendix Figure 2), but the *P. inui* sequence obtained from case-patient PMAR0052 was of the sexual type 18S rRNA (Table). The human *P. inui*-positive cases we detected originated from separate DNA extractions and PCR assays on different days by using dedicated benchtops for different procedures. The case-patients had the only *P. inui*-positive samples, but we identified a few *P. knowlesi*-positive samples among the 71 persons screened at the camp (Appendix Figure 1). The *An. craccens* mosquito was the only *Plasmodium*-positive mosquito we detected. We hypothesize that *P. knowlesi*–positive mosquito was monoinfected with *P. inui*.

<table>
<thead>
<tr>
<th>Sequence source and length, bp (GenBank accession no.)</th>
<th>Description of sequence (GenBank accession no.)</th>
<th>% Identity</th>
<th>% Query cover</th>
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<tbody>
<tr>
<td>Patient PMAR0041, 234 (MW555281)</td>
<td><em>P. inui</em> asexual type 18S rRNA, Celebes (AB287276)</td>
<td>99.57†</td>
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<td><em>P. inui</em> asexual type, 18S rRNA, Thailand (EU400385)</td>
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<tr>
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<td><em>P. inui</em> asexual type 18S rRNA, Taiwan II (FN430725)</td>
<td>99.49</td>
<td>100</td>
</tr>
</tbody>
</table>

†This sequence had only a 1 single-nucleotide mismatch at the forward primer priming site.
‡This sequence had 2 single-nucleotide mismatches, 1 at the forward primer priming site.

**Figure 1.** Species-specific nested PCR amplification products for a study of *Plasmodium inui* infections among humans, Malaysia. Samples were subjected to electrophoresis on a 1.5% agarose gel. A) Results for detection of *P. knowlesi*, *P. falciparum*, *P. vivax*, *P. malariae*, *P. ovale*, and *P. cynomolgi*. Lane 1, human case-patient PMAR0041; lane 2, human case-patient PMAR0052; lane 3, no-template control. B) Results for the detection of *P. inui* in human case-patient PMAR0041. Lane 1, case-patient PMAR0041; lane 2, negative control; lane 3, no-template control. The solid vertical line indicates these are separate parts of the same image. C) Results for the detection of *P. inui* in human case-patient PMAR0052. Lane 1, case-patient PMAR0052; lane 2, no-template control.
with *P. inui* because we found no other *Plasmodium* species in any of them.

**Conclusions**

In experimentally infected humans, patent *P. inui* infections appeared >31 days after infectious mosquito bites (8). Similarly, both cases we report show a patent infection ≈30 days after alleged exposure. *P. inui* undergoes a 72-hour erythrocytic cycle, causing quartan fever (8). Infection by the quartan *P. inui* could be self-limiting in humans because the parasite was not detected in case-patient PMAR0041 ≈8 months after exposure. Indeed, *P. inui* infections in monkeys are usually low-grade and chronic and can be self-limiting (9,10). In addition, the *P. inui* OS strain parasite count in experimentally infected humans was <2,520/µL blood. Symptoms were mild, and parasitemia could be submicroscopic or undetectable for certain periods. Antimalarial intervention was deemed unnecessary in these experimental infections (8).

Natural human *P. inui* infection seems possible, but because of the very low number of parasites and sharp fluctuations between negative and moderate parasitemia by microscopy (8,10), previous studies that used less sensitive methods, including standard PCR, were not able to detect it (11). We show that nPCR on concentrated, high-volume blood was more sensitive at detecting low-grade infection than standard PCR (12), which highlights the need for ultrasensitive detection tools.

We found 2 forms of *Plasmodium* 18S rRNA genes: the asexual type, which is expressed during the parasite's asexual life cycle in the vertebrate host; and the sexual type, which is expressed during its sexual life cycle in the mosquito vector. The *Plasmodium*-genus PCR primers we used amplify asexual and sexual 18S rRNA, but the *P. inui*-specific primers only amplify the asexual type, which explains the negative results from the species-specific nPCR despite the positive amplifications in the *Plasmodium*-genus PCR. Nonetheless, successful PCR amplification is compounded by low levels of parasites and the subsequent chance effect that can lead to occasional spurious results, as we experienced.

*P. inui* sporozoites have been found naturally occurring in *An. cracens* mosquitoes (2). Other mosquito species from the Leucosphyrus group can transmit *P. inui* naturally (2). In addition, laboratory experiments showed *P. inui* adapted to co-indigenous *Anopheles* mosquito species (13).

*P. inui* has a wide geographic range in Asia, including southern India, Southeast Asia, and Taiwan (13). A surveillance study reported that the prevalence of *P. inui* among wild macaques in Pahang was 66.7% (26/39 macaques sampled); 76.9% of these infections were co-infections with other *Plasmodium* species (14). Given the high prevalence of *P. inui* among macaques and natural *Anopheles* mosquito vectors (2), humans could be exposed to *P. inui* via vectorborne transmission from infected macaques, particularly at a location where humans, macaque hosts, and mosquito vectors co-exist. Furthermore, studies report that *P. inui* often occurs in co-infections with *P. knowlesi* and *P. cynomolgi* in monkeys and mosquitoes (2), and that humans frequently can be exposed to a mix of non-human primate malaria sporozoites (15). Because human *P. inui* infections can be asymptomatic, *P. inui* could evolve to efficiently infect humans (2), especially considering patent human infection can be established by just a few parasites (8). Strains from different geographic locations might even exhibit different infection patterns. Investigators should use ultrasensitive methods for epidemiologic and entomological studies of simian malaria transmissions in Malaysia and other countries in malaria elimination efforts.
Acknowledgments
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References

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