An Alternative Method for Extracting *Plasmodium* DNA from EDTA Whole Blood for Malaria Diagnosis

Krongkaew Seesui1, Kanokwan Imitawi2, Phimphakon Chanetmahun3, Porttip Laummaunwai1,4*, Thidarut Boonmars1,4

1Department of Parasitology, Faculty of Medicine, Khon Kaen University, Khon Kaen 40002, Thailand; 2Department of Biochemistry, Faculty of Medicine, Khon Kaen University, Khon Kaen 40002, Thailand; 3Wiang Haeng Hospital, Wiang Haeng, Chiang Mai 50350, Thailand; 4Neglected, Zoonosis and Vector-Borne Disease Research Group, Khon Kaen University, Khon Kaen 40002, Thailand

**Abstract:** Molecular techniques have been introduced for malaria diagnosis because they offer greater sensitivity and specificity than microscopic examinations. Therefore, DNA isolation methods have been developed for easy preparation and cost-effectiveness. The present study described a simple protocol for *Plasmodium* DNA isolation from EDTA-whole blood. This study demonstrated that after heating infected blood samples with Tris-EDTA buffer and proteinase K solution, without isolation and purification steps, the supernatant can be used as a DNA template for amplification by PCR. The sensitivity of the extracted DNA of *Plasmodium falciparum* and *Plasmodium vivax* was separately analyzed by both PCR and semi-nested PCR (Sn-PCR). The results revealed that for PCR the limit of detection was 40 parasites/µl for *P. falciparum* and 35.2 parasites/µl for *P. vivax*, whereas for Sn-PCR the limit of detection was 1.6 parasites/µl for *P. falciparum* and 1.4 parasites/µl for *P. vivax*. This new method was then verified by DNA extraction of whole blood from 11 asymptomatic Myanmar migrant workers and analyzed by Sn-PCR. The results revealed that DNA can be extracted from all samples, and there were 2 positive samples for *Plasmodium* (*P. falciparum* and *P. vivax*). Therefore, the protocol can be an alternative method for DNA extraction in laboratories with limited resources and a lack of trained technicians for malaria diagnosis. In addition, this protocol can be applied for subclinical cases, and this will be helpful for epidemiology and control.

**Key words:** *Plasmodium falciparum*, *Plasmodium vivax*, EDTA whole blood, malaria diagnosis, DNA extraction, semi-nested PCR

**INTRODUCTION**

Malaria remains a life-threatening disease. In 2015, the World Health Organization reported that the number of cases globally was estimated at 214 million, accounting for nearly 438,000 deaths [1]. For malaria diagnosis, various reports have shown that microscopy and RDTs have low sensitivity with the limit of detection as low as 100 parasites/µl [2-4]. Therefore, PCR-based methods, including semi-nested PCR (Sn-PCR), have been introduced to overcome these problems [5,6].

DNA extraction is an important step in molecular diagnostic approaches. Numerous protocols have been developed to isolate *Plasmodium* DNA from a variety of blood sources in order to obtain high-quality DNA suitable for downstream applications [7,8]. Commercial DNA extraction kits are commonly used to reduce PCR inhibition caused by hemoglobin in red blood cell [9,10]. However, these kits are not always available in low-resource settings. Another method for DNA extraction in routine use involves the application of phenol and chloroform [11]. However, the drawbacks of the phenol-chloroform-based method are that it is labor-intensive and time-consuming, with a high risk of contamination due to the large number of tube changing steps. In 1994, genomic DNA extraction by isopropanol fractionation with concentrated sodium iodide (NaI) and sodium dodecyl sulfate (SDS) was successfully introduced and performed in a single tube, which reduced the loss and fragmentation of DNA caused by pipetting of DNA during multiple tube changes [12]. The microwave irradiation based DNA extraction has been reported to be rapid, cost-effective and easy to perform with 10 µl of whole blood and the results showed the limit of detection of 5 parasites/µl using nested PCR technique [9]. The limitation of this method is the capacity of tubes to be used should not be less than 0.5 ml, as they can damage when irradiated and might be contaminated.
with biological materials [9], the lysate obtained from heating infected red blood cells with proteinase K and TE buffer was purified by isotachophoresis (ITP) prior to performing PCR and quantitative PCR. This method showed a limit of detection of 500 parasites/µl. Nevertheless, this technique is the high cost because of using the specialized device, microfluidic, for DNA purification.

Therefore, the aim of this study was to demonstrate a simple protocol using TE buffer and proteinase K to extract Plasmodium DNA from EDTA-whole blood for subsequent downstream applications directly, without prior purification methods. The sensitivity of the established method was determined by serial dilution with normal blood and analysis using PCR and Sn-PCR. In the present study, this new protocol was verified by DNA extraction from whole blood of asymptomatic migrant workers who attended the screening program in order to work in Chiang Mai Province. In addition, the steps for extraction of the new protocol were compared with a reference method, commercial DNA extraction kit, and a previously published DNA extraction protocol [12] by focusing on cost-effectiveness and labor-intensiveness.

**MATERIALS AND METHODS**

Whole blood samples were obtained from patients at a hospital in Chiang Mai province, Thailand. EDTA-treated samples were examined for *P. vivax* and *P. falciparum* by microscopy; parasitemia levels in the blood samples were 0.38% and 0.01%, respectively. Normal blood taken from healthy persons with no history of malaria infection was serially diluted with malaria-infected blood. To verify the new method, whole blood samples obtained from 11 asymptomatic Myanmar migrant workers who attended screening program in order to work in Chiang Mai Province. After routine diagnosis, the samples were transported to the laboratory in a cooler and then stored at -20°C until used. The study protocol was approved by the Khon Kaen University Ethics Committee for Human Research (HE591044). Informed consent was obtained from the study subjects using a standard approved procedure.

*P. falciparum* and *P. vivax* DNA was extracted from EDTA-treated whole blood samples using 3 methods: a commercial DNA extraction kit (Qiagen QIAamp DNA Blood Mini Kit, Qiagen, Hilden, Germany); a previously published DNA extraction protocol [12] by focusing on cost-effectiveness and labor-intensiveness.

### Table 1. DNA extraction methods from EDTA whole blood

| Commercial DNA extraction (Qiagen QIAamp DNA Blood Mini Kit) | DNA extraction performed with one tube (Wang et al. 1994) | The new protocol (TE buffer and proteinase K-based method) |
|------------------------------------------------------------|----------------------------------------------------------|----------------------------------------------------------|
| · Add 180 µl of lysis buffer and proteinase K to 100 µl of EDTA whole blood | · Add 500 µl lysis buffer (1% (w/v) Triton X-100, 0.32 M saccharose, 5 mM MgCl₂, and 10 mM Tris–HCl [pH 7.5]) to 100 µl of EDTA whole blood and mix vigorously | · Add 1 ml of sterile distilled water to 100 µl of EDTA whole blood, then vortex vigorously |
| · Incubate at 56°C for 3 hr | · Centrifuge at 12,000 g for 5 min | · Incubate at 4°C for 10 min |
| · Add 200 µl of B3 solution | · Add 0.2 ml of enzyme reaction solution and proteinase K solution (22 mg/ml), and mix occasionally by inversion during incubation at 56°C for 1 hr | · Centrifuge at 12,000 g for 5 min and discard supernatant |
| · Incubate at 70°C for 10 min | · Centrifuge at 12,000 g for 5 min | · Add 40 µl of TE buffer and 10 µl proteinase K solution (22 mg/ml) |
| · Centrifuge at 12,000 g for 5 min | · Precipitate DNA with 0.3 ml of NaI solution and mix gently a few times by inversion | · Incubate at 56°C for 1 hr, and mix occasionally during incubation |
| · Transfer supernatant to a new microcentrifuge tube | · Add 0.5 ml of isopropanol and mix well by inverting the tube repeatedly | · Inactivate proteinase K at 80°C for 10 min |
| · Add 500 µl of absolute ethyl alcohol and transfer the solution into a column tube | · Centrifuge at 12,000 g for 10 min | · Centrifuge at 12,000 g for 5 min |
| · Centrifuge at 12,000 g for 5 min | · Add 200 µl of 70% ethanol, centrifuge at 12,000 g for 10 min, remove supernatant, and dry DNA for 30 min | · Remove and store supernatant at -20°C until used |
| · Discard flow-through and then add 600 µl of washing buffer to the column tube | · Reconstitute DNA with sterile distilled water and store at -20°C until used | · Discard flow-through and then add 600 µl of washing buffer to the column tube |
| · Centrifuge at 12,000 g for 5 min | · Place the column tube into a new microcentrifuge tube and then add 30 µl of elution buffer | · Centrifuge at 12,000 g for 5 min, then store eluted DNA at -20°C until used |
method [12] and the new protocol (TE buffer and proteinase K-based extraction). The protocols for DNA extraction by the 3 methods are given in Table 1. For Polymerase chain reaction, each gene of *P. falciparum* and *P. vivax* DNA was performed in separate PCR reactions using specific primers designed from 18S rRNA, as previously described [13]. A forward universal primer (UF) was designed from a genus-specific region, and reverse primers (FR for *P. falciparum* and VR for *P. vivax*) were designed from a species-specific region of *P. falciparum* and *P. vivax*. The oligonucleotide primers were as follows:

UF, 5’-TCAGCTTTTGATGTTAGGGTATT-3’; FR, 5’–GCACTAAAGATACAAAATATAAGC-3’; VR, 5’-TAAACTCCGAAGAGAATAATTCT-3’.

The PCR tube contained a total volume of 20 μl, consisting of 10 mM KCl buffer, 5 mM Tris–HCl, 2.5 mM MgCl₂, 200 mM dNTP, 0.5 μM of each primer, and 0.1 units of RBC Taq DNA polymerase (RBC Bioscience, New Taipei City, Taiwan). Reactions were performed in a GeneAmp PCR System 9700 thermal cycler (Thermo Fisher Scientific, Waltham, Massachusetts, USA). The cycling conditions for *P. falciparum* and *P. vivax* consisted of an initial denaturation phase at 94°C for 2 min, 40 repetitions at 94°C for 30 sec, annealing of each primer pair at 47°C for 1 min, and extension at 72°C for 1 min, followed by a final extension for 10 min at 72°C. The PCR products were visualized by electrophoresis on 1.5% agarose gel stained with ethidium bromide.

To determine the limitations of DNA extracted by the new method, 2 malaria blood samples with known parasitemia, 0.38% for *P. vivax* and 0.01% for *P. falciparum*, were serially diluted with uninfected blood. The DNA extracted from each dilution was used as a template for conventional PCR and Sn-PCR. For conventional PCR, the conditions for PCR amplification were as described above (UF-VR for *P. vivax* identification, UF-FR for *P. falciparum* identification). The specific primers for Sn-PCR used in this study were designed from a small subunit of the *Plasmodium* ribosomal gene (ssrDNA), as previously described [14]. The primers are shown in Table 2. Sn-PCR amplification of each gene was performed separately, as follows. Nest 1: a total of 20 μl of mixture contained 10 mM KCl buffer, 5 mM Tris–HCl, 2.5 mM MgCl₂, 200 mM dNTP, 0.5 μM of each primer, 0.1 units of RBC Taq DNA polymerase and 2 μl of DNA template. Nest 2: a total of 20 μl of mixture contained 10 mM KCl buffer, 5 mM Tris–HCl, 2.5 mM MgCl₂, 200 mM dNTP, 0.5 μM of each primer, 0.1 units of RBC Taq DNA polymerase and 2 μl of 1:10 diluted DNA from the first reaction. All reactions were performed in a GeneAmp PCR System 9700 thermal cycler. The amplicon was visualized by 1.5% agarose gel electrophoresis and stained with ethidium bromide. To verify the new method, the EDTA-whole blood of 11 asymptomatic Myanma migrant workers who attended the screening program at ChaingMai hospital were used to extract DNA by using the new protocol and analyzed by Sn-PCR, using specific primers as shown in Table 2. The first reaction, which included primers UNR, PLF and HUF; used a total of 20 μl of a mixture consisting of 10 mM KCl buffer, 5 mM Tris–HCl, 2.5 mM MgCl₂, 200 mM dNTP, 0.1 units of RBC Taq DNA polymerase, and 2 μl of DNA template. The amounts of the 3 primers used in this reaction were 0.5 μM for UNR, 0.5 μM for PLF, and 0.05 μM for HUF. The second reaction, which included PLF, FAR and VIR, used a total of 20 μl of a mixture consisting of 10 mM KCl buffer, 5 mM Tris–HCl, 2.5 mM MgCl₂, 200 mM

**Table 2.** List of primers for semi-nested PCR (Sn-PCR) used in this study

| Primers     | Sequences (5’-3’)          | Specificity         | Size of PCR product (bp) |
|-------------|----------------------------|---------------------|-------------------------|
| First reaction Reverse primer UNR | 5’-GAG GGT ATC TGA TCG TCT T-3’ | Universal          |                         |
| Forward primer PLF, HUF | 5’-AGT GTG TAT CAA TCG AGT TT-T-3’ | *Plasmodium* mammals | 783-821*               |
|             | 5’-GAGCGGCCCTGGATAACGC-3’ |                      | 231*                   |
| Second reaction PLF, FAR | 5’-AGT GTG TAT CAA TCG AGT TT-T-3’ | *P. falciparum*     | 395                    |
|             | 5’-AGT TCC CCT AGA ATA GTT ACA-3’ |                 |                         |
| Second reaction PLF, VIR | 5’-AGT GTG TAT CAA TCG AGT TT-T-3’ | *P. vivax*         | 499                    |
|             | 5’-AGG ACT TCC AAG CCG AAG-3’ |                     |                         |

*Position 29 on the published sequence X03205 for human 18S rRNA.
*Size depending upon species: e.g., *P. falciparum* = 787 bp, *P. vivax* = 783 bp.
*231 for human.
dNTP, 0.5 μM of each primer, 0.1 units of RBC Taq DNA polymerase, and 2 μl of 1:10 diluted DNA from the first reaction. All reactions were performed in a GeneAmp PCR System 9700 thermal cycler. The amplicon was visualized by 1.5% agarose gel electrophoresis and stained with ethidium bromide. The estimated cost for each of the 3 methods used to extract DNA from the samples was calculated based on the purchase price of the reagents and kits that were used in US dollars (USD).

RESULTS

*P. falciparum* and *P. vivax* DNA extracted from EDTA whole blood using the new method (TE buffer and proteinase K-based extraction) was successfully amplified by PCR as shown of the reagents and kits that were used in US dollars (USD).

**Fig. 1.** Amplified *P. falciparum* and *P. vivax* gene products after PCR. (A) PCR product of *P. falciparum*. (B) PCR product of *P. vivax*. Lane M, DNA ladder; lane C, negative control; lane 1, PCR product of DNA extracted using a NucleoSpin® Blood; lane 2, PCR product of DNA extracted by a published DNA extraction method; lane 3, PCR product of DNA extracted by the modified method (TE buffer and proteinase K-based extraction). The arrow indicates the expected size of amplicons.

**Fig. 2.** Limit of detection of *P. falciparum* using TE–proteinase K-based method and standard PCR. The numbers above each lane indicate the number of parasites present per µl of blood. Infected whole blood was serially diluted with normal whole blood. The initial parasitemia of *P. falciparum* was 0.01%. Lane M is a DNA size marker.

**Fig. 3.** Limit of detection of *P. vivax* using TE–proteinase K-based method and standard PCR. The numbers above each lane indicate the number of parasites present per µl of blood. Infected whole blood was serially diluted with normal whole blood. The initial parasitemia of *P. vivax* was 0.36%. Lane M is a DNA size marker.
in Fig. 1. The product sizes of the target genes of *P. falciparum* was 1,451 bp and *P. vivax* was 833 bp. PCR products were not obtained from the negative control lacking *Plasmodium* species. This negative result indicates that the reagents had not been contaminated while setting up the process. The sensitivity of PCR and Sn-PCR assays after TE buffer and proteinase K-based extraction was determined based on serial dilution of EDTA-treated whole blood with normal blood. *P. falciparum* was diluted from 1,000 to 0.06 parasites/µl, and *P. vivax* was diluted from 4,400 to 0.06 parasites/µl. Standard PCR was then performed using specific primers, as previously described [13]. The results revealed that the limit of detection was 40 parasites/µl for *P. falciparum* and 35.2 parasites/µl for *P. vivax* (Figs. 2, 3). For Sn-PCR assay, *P. falciparum* was diluted from 5,000 to 0.002 parasites/µl, and *P. vivax* was diluted from 22,000 to 0.01 parasites/µl. The results revealed that the limit of detection was 1.6 parasites/µl for *P. falciparum* and 1.4 parasites/µl for *P. vivax* (Figs. 4, 5).

The new protocol showed successful DNA extraction from EDTA whole blood of 11 asymptomatic Myanmar migrant workers and then analyzed by Sn-PCR as shown in Fig. 6. The product size of the human gene from the first reaction amplified from UNR and HUF primers was 231 bp (Fig. 6A). Thereafter, the amplicon from the first reaction was diluted 1:10 and used as template for the second reaction. The results of the second reaction which using specific primers for *P. falciparum* and for *P. vivax* showed that there were 2 positive samples for *P. falciparum* at the target size of 395 bp (Fig. 6B, lane 7) and 499 bp for *P.
vivax (Fig. 6B, lane 2). Cost per sample was estimated based on the price of reagents or kits used in each of the 3 methods. For the commercial kit, the costs of the previously published DNA extraction method and the method established here were approximately 2.86 US$, 1.86 US$ and 0.15 US$, respectively. For time consuming, the commercial kit took around 4 hr and the published DNA extraction method took time around 2 hr and 30 min, while the new method took time to perform around 1 hr and 30 min. The new method here was, thus, more cost-effective and less time consuming than the 2 with which it was compared.

**DISCUSSION**

Over the years, multiple protocols have been developed and applied for extraction of *Plasmodium* DNA for various purposes, such as diagnostic [15,16], therapeutic uses [17], and epidemiology and control [18]. The aim of this study was to establish a simple protocol for DNA extraction from EDTA whole blood for malaria diagnosis. The protocol consisted of hemolysis by sterile distilled water and partitioning of the DNA from other cell components by incubating at 56˚C after adding TE buffer and proteinase K solution. Following enzyme inactivation and centrifugation, the supernatant was directly used for malaria detection by molecular methods, and no chemicals were required for isolation or purification. This study demonstrated that *Plasmodium* DNA was successfully extracted using this protocol and could be subjected to both standard PCR and Sn-PCR amplification. The sensitivity level of DNA extracted by TE buffer and proteinase K-based method was determined from serial dilutions of blood samples. The results of standard PCR showed a limit of detection of 40 parasites/µl for *P. falciparum* and 35 parasites/µl for *P. vivax*. Sn-PCR technique had greater sensitivity, with a limit of detection of 1.6 parasites/µl for *P. falciparum* and 1.4 parasites/µl for *P. vivax*. Thus, the new protocol in this study could be used to extract *Plasmodium* DNA and apply diagnostic measures to overcome the deficiencies of identification by microscopic examination by using standard PCR or Sn-PCR. Moreover, this protocol was also verified by extraction DNA from EDTA whole blood samples of asymptomatic Myanmar migrant workers who attend the screening program in order to work in Chiang Mai Province and the results demonstrated that this protocol can be used to detect low parasite levels by Sn-PCR and there has been reported that this molecular technique can be applied successfully in cases of asymptomatic infection [14,19]. This will be helpful for epidemiology and control, since asymptomatic persons play an important role in the transmission of malaria.

Techniques that have been employed for malarial DNA extraction from whole blood include rapid boiling, Chelex-100 chelating resin, microwave irradiation, isotachophoresis, and boil and spin method [20]. This is the first study to only use TE buffer and proteinase K for *Plasmodium* DNA extraction from EDTA whole blood. The steps for DNA extraction in this protocol were compared with a commercial DNA extraction
kit and a previously published DNA extraction method [12], as shown in Table 1. In addition, these 2 features are clear advantages of this protocol, which is easy to perform, cost-effective, and less labor-intensive than alternative methods. However, the DNA quality may deteriorate during long-term storage because the sample is relatively unpurified. The effective implementation plans for malaria control are mass testing and prompt treatment. Therefore, governments of the countries should provide funding for malaria control by using effective interventions. As mentioned above, microscopic examination is low sensitivity and need experienced technician to read blood film, molecular techniques have been used to address these problems. The new protocol described offers a simple and cost effective for DNA extraction that can be used for mass testing in malaria control.

The new protocol in this study offers simple, low cost and not labor-intensive. It can be an alternative method for DNA extraction from EDTA whole blood in a clinical laboratory of a hospital with limited resources and lacking an expert microscopist. In addition, this protocol can be applied for detection of low parasitemia or subclinical individuals by using Sn-PCR. This will be helpful for active malaria surveillance.

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CONFLICT OF INTEREST

The authors declare that there is no conflict of interest.

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