THE CIDR1α-PfEMP1 SEQUENCE FROM INDONESIAN PLASMODIUM FALCIPARUM AND ITS POTENTIAL ASSOCIATION WITH THE CEREBRAL OUTCOME

Erma Sulistyaningsih1,2, Yunita Armiyanti1, Rosita Dewi2,3

Correspondence: sulistyaningsih fk@unej.ac.id
1Parasitology Department, Faculty of Medicine, University of Jember, Jember, Indonesia
2Center for Development of Advanced Science and Technology, University of Jember, Jember, Indonesia
3Histology Department, Faculty of Medicine, University of Jember, Jember, Indonesia

ABSTRACT

Background: Plasmodium falciparum Erythrocyte Membrane Protein 1 (PfEMP1) is an important protein responsible for the pathogenesis of severe malaria, including cerebral malaria. The protein is highly diverse. The CIDR1α-PfEMP1 binds endothelial protein receptor (EPCR) and may associated with the brain swelling in childhood malaria.

Objective: To analyze the CIDR1α-PfEMP1 from Indonesian isolate and determine its association with cerebral malaria outcome.

Methods: Fifteen blood samples of clinically mild to severe malaria-patient were collected for DNA extraction. Malaria diagnosis was conducted microscopically by Giemsa-stained thin blood smear. The CIDR1α domain was amplified by PCR using specific primer and PCR product was sequenced. The nucleotide sequences were analyzed by NCBI blast, DNASIS MAX 3 and translated into amino acid sequences using Expasy Translation Tool.

Results: One out of fifteen samples was severe malaria case and infected with P. falciparum, the rest were clinically mild to moderate malaria and infected with pure P. falciparum or mixed infection of P. falciparum and P. vivax. Amplification for CIDR1α domain resulted a single band of +550 bp from a severe sample only. Sequencing of PCR product on both strands read 524 nucleotides and BLAST analysis confirmed as CIDR1α sequence. Multiple alignment showed 74-78% nucleotide sequence similarity with reference sequences, but amino acid sequences presented 23.5% homologous.

Conclusion: An identified CIDR1α domain only from severe case implicating the potential association with the severe outcome including cerebral malaria, but the highly diverse of the domain needs further studies on the interaction with the pathological-causing receptor in the host.

Keywords: Cerebral outcome, CIDR1α, PfEMP1, Plasmodium falciparum

Introduction

Malaria is infectious disease caused by Plasmodium spp and transmitted by female Anopheles mosquito. It is responsible for 228 million cases with approximately 405,000 deaths annually.1 Plasmodium falciparum is the most prevalent malaria parasite in the world, ranging from 50% in South-East Asia Region, 71% in the Western Pacific Region to 99.7% in African Region. It is the most deadly Plasmodium, causing broad clinical symptoms from mild to severe cases even leading to death.1

The important pathology of P. falciparum infection is cytoadherence and rosetting.2,3 There are several proteins involved in these two-central pathogenesis, one of the most important is P. falciparum Erythrocytes membrand Protein 1 (PfEMP1).4 PfEMP1 is a complex protein, contains a highly variable extra-cellular part and a relatively conserved intra-cellular part. The extra-cellular part consist of N-terminal segment (NTS) followed by 2-10 copies of two distinct binding domains: Duffy binding-like (DBL) and Cysteine-rich interdomain regions (CIDR).5 PfEMP1 is encoded by var gene family consisting of approximately 60 variable genes per haploid genome of the parasite.6 Var genes are highly variable in sequences but possess common structural features including conserved DBL and CIDR domains. The CIDR domain consists of semi-conserved stretches and is classified into three different types: α, β, γ and δ.7 The CIDR1α domain of several different PfEMP1 proteins was shown to bind CD36 and endothelial protein C receptor (EPCR).7,8 The expression of CIDR1α-PfEMP1 and the EPCR-binding phenotype are associated with the severe childhood malaria.1,9-10 Studies reported that EPCR-binding CIDR1α domains are highly diverse, even in the EPCR-directly contact residue. In this report, we described the sequence characteristic of the CIDR1α domain from Indonesian P. falciparum isolates and analyzed its potential association with malaria outcome.
Methods

Samples and Study Site
Malaria patients were enrolled from the Primary Health Care in Jember district, East Java, Indonesia. Patients were informed and signed the informed consent before study. The study was received an ethical approval from the Ethical Committee of Faculty of Medicine University of Jember with the reference Nr. 1114/H25.1.11/KE/2017. The inclusion criteria were infection with P. falciparum either pure or mixed infection confirmed with microscopic examination of thin blood smears stained with Giemsa.

DNA extraction and Amplification of CIDR1α domain
Genomic DNA (gDNA) was isolated from blood samples of malaria patients by TIANamp Blood DNA kit (Tiangen Biotech) according to the manufacturer's instructions. The CIDR1α domain was amplified using specific primer according to previous study.(11) The primers were: CIDR-F (5'-CGGGATCCAAATGGAAATGTTATTAG-3') and CIDR-R (5'-GGGGTACCTTTGAGTATTTATCAATT-3'). The cycle conditions for the PCR were as follows: initial denaturation at 95°C for 4 min, followed by denaturation at 95°C for 45 sec, annealing at 46°C for 60 sec and extension at 72°C for 60 sec, for 35 cycles and final extension at 72°C for 10 min. The amplified fragments from PCR were electrophoresed in 1% agarose gel and visualized using UV light transilluminator.

Sequencing of PCR products and Sequence Analysis
PCR products were purified and directly sequenced using the ABI PRISM 3730 Version 3.1 sequencer (Applied Biosystems). The sample was sequenced on both strands, i.e., forward and reverse.

The nucleotide sequences derived from the P. falciparum field isolate were blasted to confirm its identity using Basic Local Alignment Search Tool (BLAST) in the National Center for Biotechnology Information (NCBI) (http://www.ncbi.nlm.nih.gov/BLAST.cgi). The sequences were aligned and analyzed for sequence similarity by NCBI domain. The nucleotide sequences were translated into amino acid sequences using Expasy Translation Tool (http://www.expasy.ch/tool/dna.html). Percentage sequence similarity and phylogenetic tree building was carried out based on a Neighbour-Joining methods in DNASIS MAX 3.

Results

Characteristic of samples
As many as fifteen blood samples of malaria patients from Jember, East Java, Indonesia were enrolled in the study after written informed concern. Fourteen out of 15 patients (93.3%) were male and the rest was female (6.7%). One out of fifteen samples was severe malaria case and infected with P. falciparum, the rest were clinically mild to moderate malaria and infected either single infection of P. falciparum or mixed infection of P. falciparum and P. vivax, as shown in Table 1. Microscopic examination as a gold standard for malaria diagnosis confirmed 9 samples (60%) pure of P. falciparum infection, the rest patients (40%) showed mixed infection of both P. falciparum and P. vivax. There were neither P. malariae nor P. ovale infection. Clinical manifestation showed a wide range of symptom, where only a patient with P. falciparum infection showed a severe malaria with anemia as a prominent symptom, the rest were either moderate or mild malaria. Based on WHO classification, malaria patient is categorized as having severe malaria when there are at least one symptom, either severe anemia, prostration, convulsion and respiratory distress, metabolic acidosis or cerebral malaria with impaired consciousness and coma.(12) In this study, the severe malaria patient had a symptom of severe anemia with the Hb < 5g/dL and decreased of consciousness.

<table>
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Amplification and Sequence Analysis of the CIDR-1α Domain
The amplification of CIDR1α sequence using specific primer resulted a single band of approximately 550 bp in a severe malaria sample only, as shown in Fig. 1. The band is similar with the previous report on the CIDR domain that the CIDR sequence were approximately 600 bp and 520 bp from the cDNA amplification of iTG2.F6 strain and 510 bp from the gDNA of K1 strain.11 The study also analyzed the CIDR1α domain by sequencing. Sequencing from both strands resulted 524 nucleotides. BLAST-ing analysis showed that the resulted sequence had 74-78% sequence similarity with previous sequences of P. falciparum isolates in the GenBank. The phylogenetic tree analysis showed in Figure 2. The sequence had 82-84% identity with P. falciparum reference sequences (LR129699.1 and LR131409.1) with the query coverage of 79-87%. It also showed 78% identity and 99% query coverage with the KX154955.1, this is P. falciparum isolate 1994-3 and 1734-2 from Tanzania which is found from children with severe malaria, 74% identity and 98% query coverage with the CIDR1α of 3D7 genome (LN999947.1 and XM_001349402.1), 77% identity and 80% query coverage with P. falciparum from Papua New Guinea ((AF050740.1), and 73% identity and 67% query coverage with FCQ strain from Malayan Camp (AF008980.1).
Translation of the nucleotide sequences into amino acid sequences using the Expasy Translation Tool resulted 174 amino acids. Analysis using Protein BLAST showed 98-100% coverage and 51-64% identity with several reference sequences (PF3D7, PFDOG_01745_Dd2, PFFCH_05578, PFMALIP_05783, PFNF135_01540, PFRAJ116, PFTANZ_03634, PFTANZ_06110, PFUGT51). Multiple alignment using DNASIS MAX 3 resulted 23.5% homologous amino acid sequences as shown in Figure 3.

Discussion

Malaria is still major health problem in Indonesia, and mostly caused by P. falciparum which has a broad spectrum of clinical outcome from symptomatic, mild, moderate, severe until life threatening and causing death. One protein has a major role in the severe pathogenesis including cerebral malaria is PfEMP1. This study analysed the CIDR1α-PfEMP1 from Indonesian malaria patients and determined its association with severe malaria outcome, including cerebral malaria.

As many as 15 malaria patients were enrolled in the study, and the characteristic was shown in Table 1. Fourteen out of 15 patients were male and the rest one was female. This result is in accordance with the previous report that in some societies, men have a greater occupational risk of contracting malaria than women in mines, fields or forests at peak biting times of mosquitoes, or migrate to areas of high endemicity for work. Most of malaria patient in our study were migrant working from Java to Papua, Kalimantan and Nusa Tenggara, where the three areas were categorized as moderate to high endemic malaria areas in Indonesia. They were infected in those migrant areas and returned as a malaria-infected person or referred from hospital to receive an appropriate treatment. Migrant workers as well as non-immune travellers are vulnerable to severe malaria, irrespective of the endemicity of the area where their infection was acquired.

Table 1 also showed that one out of fifteen samples was severe malaria case and infected with P. falciparum, the rest were clinically mild to moderate malaria and infected either single infection of P. falciparum or mixed infection of P. falciparum and P. vivax. As previously mentioned that P. falciparum is the majority cause of malaria throughout the world, which resulted broad spectrum of clinically malaria outcome from asymptomatic, mild/uncomplicated malaria to severe malaria.

The severe malaria outcome in this study was severe anaemia with haemoglobin < 5 g/dl and impaired of consciousness. Based on WHO epidemiological and research purposes, severe malaria is defined as one of more of the following symptoms, i.e., impaired consciousness, acidosis, hypoglycaemia, severe malarial anaemia, renal impairment, jaundice, pulmonary oedema, significant bleeding, shock, or hyperparasitaemia, occurring in the absence of an identified alternative cause and in the presence of P. falciparum in the blood.

As previously reported that PfEMP1 plays a major role in the pathogenesis of severe malaria, we tried to identify the presence of the protein in the sample. Amplification of the CIDR1α-PfEMP1 using CIDR specific primer resulted in a single band of approximately 550 bp only from severe malaria sample, as presented in Figure 1. The band is similar with the CIDR domain previously reported, which were approximately 600 bp, 520 bp for CIDR of the cDNA amplification of iTG2.F6 strain, and 510 bp from the gDNA of K1 strain.

The PCR product of CIDR1α-PfEMP1 was further analysed by sequencing on both strands and resulted in 524 nucleotides. The sequence identification by blasting confirmed it as CIDR1α-PfEMP1. The sequence had 78% identity with the KX154955.1, this is P. falciparum isolate 1994-3 and 1734-2 from Tanzania which is found from children with severe malaria. The study also found that the expression of the CIDR1α which is bind to EPCR consistently found in both children suffering from severe malarial anaemia or cerebral malaria. The result consistent with our finding that the only detected CIDR1α domain was from patient suffering from severe malaria with the symptom of anaemia and impaired

Figure 1. Amplification result of the CIDR1α domain using specific primer. There are a single band of approximately 550 bp band (orange line) from only one severe patient (lane 2: sample 2). M: 1 kb DNA marker; 1-15: Code number of samples; All samples except sample 2 showed negative result. C-: Negative control for PCR.
The phylogenetic tree on Figure 2 demonstrated that the CIDR1α domain from Indonesian isolate had a close relationship with the sequence of 3D7 strain, *P. falciparum* isolate 1994-3 and 1734-2 erythrocyte membrane protein (var) gene from Tanzanian children, FCQ strain from Malayan Camp and *Plasmodium falciparum* from Papua New Guinea isolates. Although it is known that the var gene family-encoding *PfEMP1* is highly diverse gene, it is likely that there is no clustering of the CIDR1α sequences based on geographical origin. Furthermore, the N-terminal DBL-CIDR head structure of *PfEMP1* has diverged molecular insight into its protein diversification, i.e. group A proteins diversified into those that bind EPCR (CIDR1α domain) and non-EPCR binders (CIDR7/10/14 domains) and group B and C encode for binding CD36 (CIDR2-6 domains). Our CIDR1α-PfEMP1 sequence had high identity with the FCQ strain from Malayan Camp (AF008980.1), it is the *P. falciparum* FCG-27 clone which express the PfEMP1 region which is bind to CD36. The CIDR1α is the domain mediating binding to CD36. Previous studies reported that there is a highly conserved shape of the domain which mediates adherence to CD36, particularly cysteine residues. Binding to CD36 is interesting as it is a feature of many parasite isolates. CD36 is a glycoprotein scavenger receptor found on the surface of various cells including platelets, macrophages, monocytes, leukocytes, dendritic cells, epithelial cells and microvascular endothelial cells. CD36 expression on cerebral endothelium of cerebral malaria patients was very little, but there was ubiquitously on lung, liver, kidney, skin and muscle vasculature. In the study we found that the severe patient showed an anaemia as the prominent symptom beside the impaired consciousness. The variability of the *PfEMP1* is further confirmed by its homologous. The nucleotide sequences of CIDR1α-PfEMP1 were translated into amino acid sequences using the Expasy Translation Tool and yielded 174 amino acids. Analysis using Protein BLAST showed 98-100% coverage and 51-64% identity with several reference sequences (PF3D7, PFDG_01745_Dd2, PFFCH_05578, PMALIP_05783, PFNF135_01540, PFRAJ116, PFTANZ_03634, PFTANZ_06110, PFUGT5.1). Figure 3
Figure 3. Multiple alignment of CIDR1α domain of PfEMP1 from Indonesian isolate with several CIDR1α reference sequences (PF3D7: P. falciparum 3D7; PFDD0_01745_Dd2: P. falciparum Dd2 isolate; PFFCH: P. falciparum from Philippines; PFMALIP_05783: P. falciparum genome from Mali; PFNF135_01540: P. falciparum genome strain NF135, PFNF135_01541: P. falciparum genome strain NF135; PFRAJ16: P. falciparum genome strain RAJ116; PPTANZ: P. falciparum genome from Tanzania; PFUGT5: P. falciparum genome strain UGT5.)
presented the CIDR1α-PfEMP1 amino acids multiple alignment using DNASIS MAX 3. It showed only 23.5% homologous amino acid sequences. This results further confirmed the variability of the CIDR1α-PfEMP1. 

Conclusion

We have reported the characteristic of CIDR1α domain of PfEMP1 from Indonesian isolate which have 524 nucleotides in length and/or 174 amino acids sequences, and the closely relation with broad sequences from different origin. The fact that the domain only amplified from severe case implicating its role in clinically outcome. The interaction of CIDR1α and EPCR which commonly found in human brain endothelial cells is suggested as the key mechanism of severe malaria outcome especially related to impairment consciousness, but the highly diverse of the domain needs further studies on the interaction with the receptors causing the pathomechanism in the host

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