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# Analysis of *Plasmodium vivax* Apical Membrane Antigen-1 (PvAMA-1) Haplotypes among Iranian Isolates

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Plasmodium vivax apical membrane antigen-1(PvAMA-1) is a surface protein with polymorphic sites. This study was aimed to analyze the polymorphic amino acid residues at PvAMA-1 in different infected age groups. 92 blood samples were collected from the south and southeast of Iran. The DNA coding for the domain I (DI), DII, and partial DIII of this antigen was amplified by Nested-PCR, and sequenced. Nucleotide mutations were found in 49 sites and based on the amino acid sequence, 30 variable sites were detected. Age distribution of malaria cases showed that the majority of the patients were between 10 to 30 years old. The scattering plot haplotypes by age showed an increasing incidence rate with age during childhood, whereas, incidence was the lowest in patients under five years old. Comparison of the polymorphic sites of PvAMA-1 in Iranian isolates with those found in other geographic regions of the world indicated nine common variable positions. In addition, a significant dependence was found between some particular substitutions and age categories. Dependence between particular substitutions and age groups suggests that certain residues in AMA-1 are responsible for clinical attacks in different ages, likely as a result of host immune pressure. The crystal structure of the PvAMA-1 showed that the amino acid substitutions that changed the protein charge were exclusively located in loops and turns where, the interactions with antibodies could occur. These data provide the necessary information for an AMA-1 based malaria vaccine design to be effective across all ages.

**Key words**: Apical membrane antigen-1 (AMA-1), DNA sequencing, genetic polymorphism, haplotype, malaria transmission, *Plasmodium vivax*.

alaria is the most important infectious disease, and continues to be a major global health problem in the world (1). According to the latest report from World Health Organization

(WHO) around 214 million cases of malaria were reported globally with an estimated 438,000 deaths (2). Of the five *Plasmodium* species causing malaria in humans, *P. vivax* is responsible for the

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most prevalent form of malaria outside Africa (3). Therefore, development of a vaccine against P. vivax is a necessary priority, particularly due to increased resistance of this parasite to the antimalarial drugs such as chloroquine, primaquine and pyrimethamine (4-7). Identification of target antigens that induce protective immune response is a pre-requisite for vaccine development. Many reports indicate apical membrane antigen1 (AMA-1) as one of the promising malaria vaccine candidates (8-11). AMA-1 is a type I integral membrane protein found in most of the Plasmodium species (11). AMA-1 expression is maximal in late schizogony during asexual reproduction in red blood cells (12, 13). It is initially located in merozoite apical organelles, and after processing relocates to the surface of mature merozoite (14). Around the time of erythrocyte invasion, the pro-sequence of AMA-1 is processed proteolytically by two c-terminal cleavages (15). The complete P. vivax AMA-1 (comprised of 562) amino acids) contains an ectodomain. Sixteen conserved cysteines contribute to 8 disulfide bonds, and divide the ectodomain into three subdomains (DI, DII, and DIII). The amino acid residues 43-248 make domain I, 249-385 make domain II, and 386-487 correspond to domain III (16). The stage specificity, and location of AMA-1 suggest that this protein is associated with the process of erythrocyte invasion as shown by the inhibition of parasite invasion using antibodies against AMA-1 (17). Also, generation of the parasites with disrupted ama-1 gene by knockout technology suggests a critical role for AMA-1 in the development of the parasite during asexual blood-stage Vaccination with AMA-1 can provide protective immunity in mice and monkeys, and induce antibodies that inhibit parasite development in vitro. These evidences introduce AMA-1 as a promising blood-stage vaccine candidate. In addition, unlike most other blood-stage proteins, ama-1 gene, excluding some regions, is a conserved

protein among various *Plasmodium* species. Point mutations are responsible for limited diversity in this antigen, and most of them are in DI (19, 20). Antigenic diversity is supposed to be a major mechanism for the parasite to escape from the host immune system, but remains the greatest obstacle in designing a malaria vaccine. Some findings show that protective immunity against AMA-1 and MSP-1 (merozoite surface protein) has a high degree of strain specificity (22, 23). Thus, AMA-1 polymorphism is a major problem to the efficacy of this protein as a vaccine component. On the other hand, some epitopes of this antigen produced by different allelic forms may be involved in the manifestation of clinical symptoms in particular age groups. Such variants should be considered to prepare repertories of AMA-1 allelic forms for developing a universal malaria vaccine. In the current study, the P. vivax ama-1 gene was amplified and sequenced from 92 blood samples collected from two different malaria endemic regions of Iran. The study was conducted to analyze the population structure of *pvama-1* Iranian isolates as an important malaria vaccine candidate. Finally, this investigation introduces the common mutation sites that should be considered for vaccine design.

# Materials and methods

#### Study area and sample collection

A total of 92 blood samples were collected from two different malaria endemic regions of Iran (Hormozgan, and Sistan & Baluchestan provinces with the highest malaria incidence on the country (24)) during 2009-2011. The blood samples were obtained from patients with uncomplicated symptomatic malaria who have been referred to local health centers in these areas. The presence of *P. vivax* infection was diagnosed by the light microscopic analysis of Giemsa-stained thick, and thin blood smears. Two ml venous blood was collected from patients in tubes containing EDTA, and stored at -20 °C until used. Blood collections

were made after obtaining informed consent from the patients or the parents/legal guardians of children. This investigation was approved by the Ethics Review Committee of Pasteur Institute of Iran.

# DNA extraction and PCR amplification of the P. vivax ama-1 gene

Whole blood sample (200 µl) was used to isolate genomic parasite DNA using the QIAamp DNA Blood Mini Kit (Qiagen, Germany) according to the manufacturer's protocol. Two-step PCR method was performed to amplify the pvama-1 gene. The 1139 bp region of the pvama-1 gene was initially amplified using set of primers PvAMAF1 (5'-GTTGAGAGAAGCACACGAATG-3') PvAMA2 (5'-GAGATAAATATCCTCGGCAGG-3'), and then the primary PCR product was used as template in a second round of PCR to amplify the internal 1000 bp region for which DI and DII are available. using the PvAMAF2 (5'-TGCAGAAGTGGAAAATGCAAAG-3') and PvAMA4 (5'-CCATCAACACTGTACAGATTC-3') primers. PCR cycles were performed as described previously (25), and the data are available in the GenBank database under the accession numbers KF181626-KF181642, and KF422636-KF422681.

# Sequence and statistical analyzes

Nucleotide and amino acid sequences were aligned by MEGA program (26), with the Salvador-1 sequence (XM\_001615397) as the reference strain. The haplotypes were classified based on the nucleotide sequences using DnaSP ver. 5. 10. 01 (27). Nucleotide mutations and amino acid variable sites were detected by the MEGA program. Nucleotide BLAST search was done to compare the *pvama-1* sequences in this study with the previously reported sequences in the Gen Bank data base. Only the sequences for which DI and DII were available were included in analysis. Comparative analysis of the variable sites at *pvama-1* sequences was performed between 92

Iranian isolates, and the previously reported sequences from various geographic regions including India, Thailand, Venezuela and Sri lanka (28-32). The statistical analysis was performed using the two tailed Fisher's exact test in GraphPad program (http:// www.openepi.com/ oe2.3/ menu/openepimenu.htm).

#### Crystal structure of PvAMA-1

Amino acid substitutions were shown on the 3D structure of PvAMA-1. The crystal structure was generated with Weblab Viewer Lite 4.2 (http:// www. scalacs.org/ TecherResources), and PDB ID 1w8k (16) was applied as the starting model.

#### Results

#### pvama-1 mutations and haplotypes

Based on the sequencing data, a 911 bp region of the *ama-1* gene including the nucleotides 289 to 1199 was readable. This nucleotide sequence codes for amino acids 97 to 399 containing the major sequence of DI, complete DII, and partial DIII of AMA-1. The 92 amplicons sequenced were classified into 53 haplotypes (Table 1).

Nucleotide mutations were found in 49 sites in the first, the second and the third codon positions (15, 14, and 20, respectively). Variable amino acid sites 130N/K, 188K/N, and 242P/E are represented by third codon positions 390T/G, 564G/T, and 726 T/A that are also responsible for the generation of new parasite variants (Table 1).

# Haplotypes association with age of infection

The number of malaria cases was highest in the 20-30 as well as 10-20 age groups (Fig.1; Table2). Nonetheless, the percentage distribution of variants among age was not significantly different from that of the whole population. The most frequent haplotype was H18 (8.7%) distributed in 10-30 and 50-60 age categories, in a statistically non-significant manner (Table 2).

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 Table 2. Distribution of malaria cases and haplotype frequency by different age categories among 92 Iranian
 PvAMA-1 isolates 1-10 9.78 H5 (×3), H6, H7, H11, H14, H48, H52 10-20 27.17 H3, H5 (×4), H7, H8, H9, H10, H11, H17, H19, H23, H24 (×5), H25, H28, H30, H36 (×2), H44, H51 H1(×2), H4 (×3), H9, H10, H12, H13, H16, H18 (×3), H20, H21, H26, H31, H32, H34, H35, H37, 20-30 28.26 H42, H43, H45, H46, H47 30-40 15.21 H9, H12, H15 (×2), H18 (×3), H27 (×3), H29, H32, H34, H53 40-50 4.34 H4 (×2), H33, H40 50-60 9.78 H2, H13, H18 (×2), H22, H27, H39, H41, H49 60-70 5.43 H4 (×2), H38, H39, H50 Total (%) 100

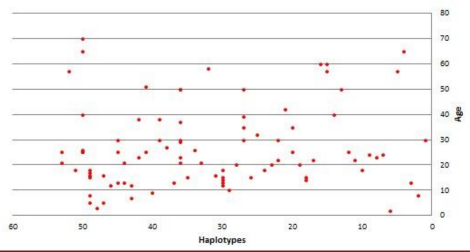


Fig. 1. Distribution of haplotype frequency by age among 92 Iranian P. vivax isolates.

#### **PvAMA-1** variants

Based on the amino acid sequences, 30 variable sites were detected, 23 of them were dimorphic, 6 were trimorphic and the position 112 showed four alternative amino acid residues (R, T, N, K, tetramorphic). The positions 112, 189, 190, 228, 382, 384 and 385 at DI and DII of the PvAMA-1 included substitutions with more than two possible amino acid residues (Fig. 2). Some of the amino acid substitutions that occurred in the segment caused changes in the charge of the protein. These consisted of Q189K, K190Q substitutions in DI, and Q277K, K352Q substitutions in DII of PvAMA-1 where they could affect the protein structure. Although, some amino acid substitutions with no effect on the protein

charge were K120R, and D242E in AMA-1 DI. Identified patterns of the mutation are one of the most important strategies used by the malaria parasites to escape from the host immune system and should be considered for vaccine design.

Comparative analysis of the polymorphic sites in PvAMA-1 between Iranian isolates and those previously reported from the other malaria endemic areas (India, Thailand, Venezuela and Sri lanka) (28-32) revealed that 9 of these variable sites were common (Table 3) including 107 D/A, 112R/T/N/K, 189E/N/K, 190K/Q/E, 210P/S in DI, 288G/E, 352K/E, 380Q/K in DII, and 384L/P/R in DIII of the PvAMA-1.

Association of PvAMA-1 variants with age of infection

In order to evaluate the effect of polymorphic amino acid positions in the AMA-1 on age of infection, the rate of amino acid residues at mutation sites shared between Iran, India, Thailand, Venezuela and Sri Lanka was assessed. The results showed a strong association between the rate of

amino acid residues Q and P in position 384, and clinical attacks in 30-40 age group. Similarly, P384, D107, K189, K190, and L384 positions were associated with malaria incidence in over 40 years (P<0.05, Table 4).

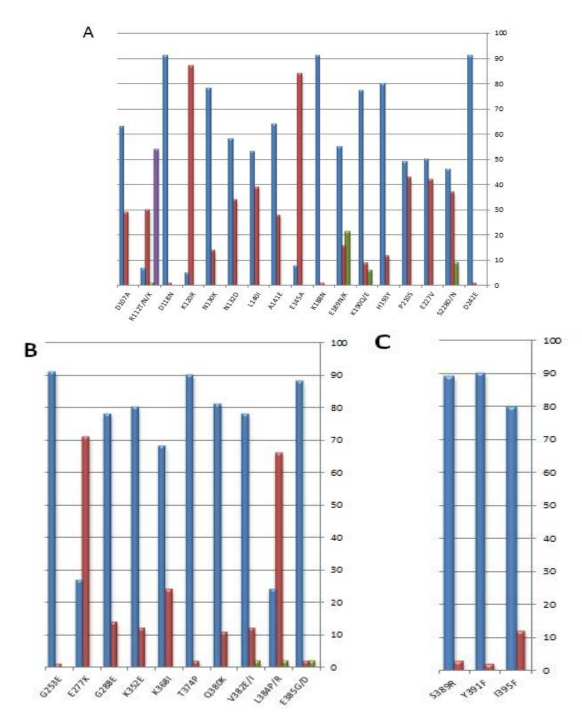


Fig. 2. The frequency of amino acid residues of each polymorphic position at PvAMA-1 isolates from Iran (n=92). A: Domain I; B: Domain II; C: Domain III..

**Table 3**. Comparative analyzes of variable sites at PvAMA-1 between Iranian isolates and the previously reported sequences in Genbank database.

reported	sequences	in Genbank	database,							
	Iran (n=92)		India (n=72)	a	Thailand (n=	=60) <sup>b</sup>	Venezuela (1	n=40) <sup>c</sup>	Sri Lanka (r	n=23) <sup>d</sup>
Position	mutations	f	mutations	f	mutations	f	mutations	f	mutations	f
25	_	_	_	_	Q/H	43/17	Q	_	Q	_
42	_	_	_	_	G	_	G/V	33/7	G	_
44	_	_	_	_	T	_	T/S	30/10	T	_
66	_	_	_	_	R	_	R/K	14/26	R	_
107	D/A	63/29	D/A/N	41/30/1	D/A	1/59	D/A	8/32	D/A/N	19/3/1
112	R/T/N/K	7/30/1/54	R/T/N/K	10/27/1/34	T/K	42/18	R/T/K/S	6/28/5/1	T/K	1/22
118	D/N	91/1	D/N	71/1	D	_	D	_	D	_
120	R/K	87/5	K/R/S	2/69/1	K/R	42/18	R/K	33/7	R	_
130	N/K	78/14	N/K	64/8	N	_	N/K	26/14	N	_
132	N/D	58/34	N/D	36/36	N	_	N/D	26/14	N/D	22/1
140	L/I	53/39	L/I	40/32	I	_	L/I	29/11	I/L	18/5
141	A/E	64/28	A/E	58/14	A	_	A/E	18/22	A/E	20/3
145	A/E	84/8	A/E	54/18	A	_	A/E	25/15	A	_
188	K/N	91/1	K/N	70/2	K	_	K	_	K/N	22/1
189	E/N/K	55/16/21	E/N/K	31/7/34	E/K	17/43	E/K	24/16	E/K	17/6
190	K/Q/E	77/9/6	K/Q/E	53/5/14	K/E	17/43	K/Q	24/16	K/E	20/3
193	H/Y	80/12	H/Y	6/66	Н	_	H/Y	35/5	H/Y	21/2
210	P/S	49/43	P/S	35/37	P/S	42/18	P/S	34/6	P/S	9/14
218	V	_	V	_	V/L	43/17	V	_	V/L	22/1
227	E/V	50/42	E/V	44/28	E	_	E	_	E/V	16/7
228	S/D/N	46/37/9	S/D	44/28	S	_	S	_	S/D	16/7
242	D/E	91/1	D	_	D	_	D	_	D	_
253	G/E	91/1	G/E	71/1	G	_	G	_	G/E	19/4
277	K/E	71/27	K/E	61/11	K	_	K/E	39/1	K/E	14/9
288	G/E	78/14	G/E	63/9	G/E	18/42	G/E	35/5	G/E	18/5
352	K/E	80/12	K/E/N	63/6/3	K/N	43/17	K/N	32/8	K/E/N	17/2/4
368	K/I	68/24	K/I	60/12	K	_	K/I	33/7	K/I	11/17
370	F	_	F	_	F	_	F	_	F/L	1/44
374	T/P	90/2	T	_	T	_	T/S	_	T	_
380	Q/K	81/11	Q/K/R	69/2/1	Q/R	59/1	Q/K	33/7	Q/K/R	17/3/3
382	V/E/I	78/12/2	V/E	64/8	V	_	V	_	V	_
384	L/P/R	24/66/2	L/P/R	38/19/15	L/R	1/59	L/P	18/22	L/P/R	9/7/7
385	E/G/D	88/2/2	E/Q/D	61/1/10	E	_	E	_	E/D	16/7
389	S/R	89/3	_	_	S	_	S	_	S	_
391	Y/F	90/2	_	_	Y	_	Y	_	Y	_
395	I/F	80/12	_	_	I	_	I	_	I	_
400	_	_	_	_	K	_	K	_	K/R	20/3
438	_	_	_	_	Н	_	H/R	12/28	H/R	20/3
445	_	_	_	_	N/D	43/17	N	_	N	_

f: frequency; n: number of isolates. The polymorphic positions common in the five comparative regions are boldfaced. <sup>a</sup> Thakur et al. (2008) (28); Rajesh et al. (2007) (29). <sup>b</sup> Putaporntip et al. (2009) (30). <sup>c</sup> Ord et al. (2008) (31). <sup>d</sup> Gunasekera et al. (2007) (32).

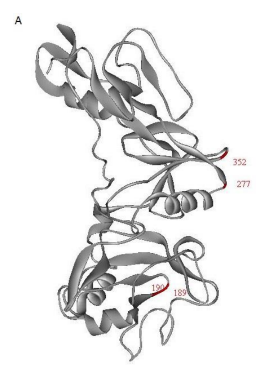
Mapping of the amino acid substitutions on 3D structure of PvAMA-1 showed that mutations with changes in the charge of the protein were exclusively located in loop and turn secondary structures whereas, other mutation sites had been spread in the molecule (Fig. 3).

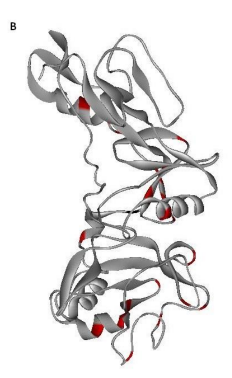
# Discussion

The present study was aimed to investigate the associations between the incidence of PvAMA-1 variants and age of infection.

The data show that the incidence of malaria reached the peak in the late teenage years, and declined with age in adulthood (Table 2; Fig. 1). This age distribution confirms that in populations with low malaria transmission, the incidence increases with age in teenagers and then declines; whereas in areas exposed to a very low level of transmission or to

epidemic malaria, the clinical attack occurs across all ages (33). The decrease in malaria intensity with age in adults in the present study is in agreement with the study on P. falciparum in Kwazulu Natalin South Africa (34), and another report on P. vivax infection by Denis-lozano et al. (35) in Mexico that can be explained by the low level of transmission intensity in these areas. Besides, adults may have had immunity early in life due to previous exposure to the malaria parasite. Our findings are in contrast with the report of Baird et al. (36) in Irian Jaya which showed an increase in malaria infection with age in a population of adults from Java after migration to a hyper endemic area. However, the examined population in the present study has been exposed to a lower level of transmission than the population in the Irian Java study. The scattering plot shows an increasing incidence rate with age





**Fig. 3.** The surface representation of the mutation sites. Data was generated with ribbon model in Weblab Viewer Lite 4.2 program. Red color shows the amino acid substitutions and all other positions are gray color. A: mutations that changed the charge of the antigen (E189K, K190E, E277K and K352E); B: other polymorphic sites.

Tal	ble 4	<b>4.</b> Age	- spec	cific	patte	erns	of ar	nino	acio	d subs	tituti	ons a	t PvAl	MA-1	in Ira	anian	isola	ites.										
			107		112					189			190		2	10	2:	88		352			380			384		
Age	f	D	A	N	R	T	N	K	S	Е	N	K	K	Q	Е	P	S	G	Е	K	Е	N	Q	K	R	L	P	R
Iran	92	68.4	31.5	-	7.6	32.6	1	58.6	-	59.7	17.3	22.8	83.6	9.7	6.5	53.2	46.7	84.7	15.2	86.9	13	-	88	11.9	-	26	71.7	2.1
0- 10	9	71.4	28.5	-	-	28.5	-	71.4	-	71.4	14.2	14.2	100*	-	-	42.8	57.1	85.7	14.2	85.7	14.2	-	57.1*	42.8*	-	14.2*	85.7*	-
10- 20	25	62.5	37.5	-	6.2	43.7	6.2	43.7	-	56.2	25	18.7	93.7*	6.2	-	62.5	37.5	87.5	12.5	93.7	6.2	-	81.2	18.7	-	12.5*	87.5*	-
20- 30	26	71.4	28.5	-	4.7	23.8	-	71.4	-	61.9	23.8	14.2	95.2*	-	4.7	57.1	42.8	90.4	9.5	90.4	9.5	-	100*	-	-	28.5	66.6	4.7
30- 40	14	88.8*	11.1*	-	11.1	33.3	-	55.5	-	44.4*	11.1	44.4*	66.6*	11.1	22.2*	55.5	44.4	88.8	11.1	88.8	11.1	-	1.0**	-	-	55.5*	33.3**	11.1
40- 50	4	100*	-	-	-	33.3	-	66.6	-	66.6	-	33.3	66.6*	33.3*	-	66.6	33.3	66.6*	33.3*	100*	-	-	100*	-	-	-	100**	-
50- 60	9	37.5**	62.5*	_	25*	50*	-	25*	_	25*	_	75**	37.5**	37.5*	25*	75*	25*	87.5	12.5	87.5	12.5	-	100*	-	_	37.5	62.5	-
60- 70	5	75	25	_	25*	25	_	50	_	50	_	50*	75	25*	_	50	50	100*	_	100*	_	_	100*	_	_	_	100**	_

The polymorphic amino acid residues of each mutation site in the studied age groups were compared with that of the whole population by the two tailed Fisher exact test in Graph pad program.\*: significant values with P<0.05;\*\*: significant values with P<0.0001

during childhood, whereas incidence is the lowest in patients under five years old. One possible explanation is that the children under 5 years old are commonly indoor at night, and are protected against mosquito bites. As children grow older, they may be more at risk of infective mosquito bites. Though, it is improbable that adults are at lower risk of infective bites than teenagers. The decline in malaria incidence in adults might explain that the infective bites in early childhood can be responsible for clinical protection in adults (33, 37). Therefore, in areas with high malaria transmission, some adults are less at risk for the disease due to the immunity acquisition occurrence prior in the life. In addition, it is possible that misdiagnosis has led to the decline in incidence recorded in young children. Misdiagnosed children may possess a measure of clinical tolerance by antimalarial therapy which has unlikely increased the malaria incidence with age in children. The existence of real differences in the reported malaria cases in different age categories would therefore support the concept of the clinical immunity hypothesis in some age categories (34).

Some amino acid substitutions found in AMA-1 are thought to be associated with disrupting epitopic targets of the immune system. Changes in amino acid charge from positive to negative or vice versa have a drastic effect on epitope detection by antibodies (21). In the present study, such replacements have occurred in four amino acid positions including Q189K, K190Q substitutions at Domain I, and Q277K, K352Q G substitutions at Domain II (Fig. 2 A and B) which were also found in India, Venezuela and Sri Lanka (Table 3).

Despite this, the Q277K replacement has not been found in Thai population that may be explained by the lower host immune selection pressure due to very low malaria incidence rate (30) and/or insufficient time to generate new alleles bearing this mutation. Moreover, some mutations involved in the amino acid changes with the same charge may be associated with functional constraints of the protein (K120R, D242E in AMA-1 Domain I) (Fig. 2 A).

Comparison of the polymorphic positions between five different populations showed that 9 positions were common (Table 3). The conservation of the common mutations suggests that these particular substitutions are necessary for the parasite to escape from the host immune system (21). If other mutations could play a similar role in the destruction of epitopes targeted by the immune system, different mutations would probably be expected between separate populations as shown in various haplotype repertoires.

The distribution of polymorphic amino acid residues (common between 5 comparative regions) by age categories (Table 4) indicated a strong age dependence between the frequency of Q and P residues in position 384, and clinical disease in 30-40 age group. Additionally, a similar situation was found in P384, D107, K189, K190, and L384 positions in age 40 and over. In another study performed on *P. falciparum* AMA-1, Cortés et al. showed that residues E187 and E243 were strongly associated with malaria incidence in children under 10 years of age (21).

This strain-dependent incidence suggests that certain residues at particular positions decrease the AMA-1 immunogenicity, and a long exposure is necessary for inducing an effective immune response. These positions were assumed to be effective in controlling growth of the malaria parasite (21).

Furthermore, these findings show that particular amino acid substitutions could be responsible for the clinical attacks in different age groups, and there is an age-related selection in the immune system against certain amino acids at particular positions of the parasite's AMA-1. In addition, the clinical presentation of the malaria infections is associated with different AMA-1 strains, suggesting that parasite factors are involved. Correspondingly, it was proposed that the higher morbidity in P. falciparum species could be related to the presence of certain residues in AMA-1(21).

Concerning the effect of certain amino acid residues on malaria infection rates, we cannot determine if these residues affect directly or their cooperation with other AMA-1 residues involved. yet, as the amino acid residues associated with the clinical malaria cases occur in multiple AMA-1 sequences, few parasite transmission can be ruled out. Thus, it is improbable that the clinical attacks are due to a combination of particular residues in other loci.

Knowledge of the amino acid substitutions, particularly those with charge conversion, could help to recognize the main epitopes that elicit protective antibodies. In the present study, these substitutions were exclusively observed in turn and loop secondary structures of the antigen (Fig. 3). Turns and loops generally lie on the surface of the proteins where they can participate in interactions between proteins and other molecules such as antibodies (38, 39).

The present study provides the necessary information to design a malaria vaccine to be effective in different age categories, and suggests that some of the epitopes in AMA-1 providing protective immunity response are strain specific. This problem is one of the difficulties in designing efficient vaccine based on only one allelic form of the protein, and was confirmed by previous reports (17, 19, 22, 40). One possible strategy to develop a universal malaria vaccine would be the inclusion of a large number of variants of the protein. Nonethe less, this strategy might be difficult for vaccine design. Therefore, knowledge of the sequences observed in the population to detect which of the variants may induce the protective immunity, could help to design an AMA-1-based vaccine. This study introduces 9 variable sites common between Iran, Thailand, Venezuela and Sri lanka including 107 D/A, 112R/T/N/K, 189E/N/K, 190K/Q/E, 210P/S in DI, 288G/E, 352K/E, 380Q/K in DII, and 384L/P/R in DIII of the PvAMA-1, that could be used as components of a polyvalent malaria vaccine based on PvAMA-1.

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# **Conflict of interest**

The authors declared no conflict of interest.

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