

The Malaria Concept in Pregnancy and the Mechanism of Evading the Immune System by the Malaria Parasite (Review)

Abstract

Malaria during pregnancy is a complex issue when considering the public health and an important contributor to maternal and infant morbidity and mortality in malaria-endemic countries. Malaria can be regarded as one of the leading cause of maternal deaths with regards to the sub-Saharan Africa. A minimum of 6 million women around the world stand the risk of being infected with malaria during pregnancy. Maternal deaths as a result of malaria occur at an approximate figure of 10, 000 per year while a minimum of 200, 00 babies also die on annual basis. Malaria remains a life threatening disease to the mother and her unborn child. The impact of the disease will depend on the strength of the mother, her immune system and the severity of the malaria. The people who are most at risk from malaria are women, who are experiencing their first pregnancies, and who are living in areas where stable malaria infections already exist. The protozoan parasites belong to the genus *Plasmodium*. Some relevant spp are *P. falciparum*, *P. vivax*, *P. malariae*, *P. ovale*, and extremely rarely *P. knowlesi* which causes malaria in macaques but can also infect humans. They are transmitted by the bite of a sporozoite-bearing female anopheline mosquito. After a period of pre-erythrocytic development in the liver, the blood stage infection, which causes the disease, begins. Parasitic invasion of the erythrocyte consumes haemoglobin and alters the red cell membrane. Malaria contributes to complications that can occur during pregnancy and these complications include anaemia, constant abortion, fetal deaths and prematurity. The first and second pregnancies experience the worst of this case. The World health Organization with governmental support over the years have put in great effort in tackling the menace of malaria in pregnancy. The major objective of the collaborative effort is the public sensitization on the use of insecticide treated mosquito nets (ITN), Intermittent preventive malaria treatment (IPT) and adequately treating acute malaria infections that occurs during pregnancy, while the combination of Sulfadoxine-Pyrimethamine as regarding the IPT has proven to be of a great importance in the prevention of chronic malaria cases that can occur during pregnancy. The introduction of the Artemisinin-Combination Therapy (ACT) by the World Health Organization serving as a first-line treatment less complicated cases of malaria occurring during pregnancy has also proven to be of high benefit. Recently, the administration of soluble of chondroitin sulphate A (CSA) to pregnant women has proven to drastically reduce parasite adhesion. The administration of chondroitinase can effectively reduce parasite by 90%, thereby reducing chances of the foetus being exposed to the parasite. It is very important to make confirmations before proceeding to the treatment of malaria and enforcing therapy completions ~~should also~~ should be encouraged.

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INTRODUCTION

Malaria can be linked to underdevelopment and the state of poverty. This has made the ~~disease~~ disease a serious public health challenge causing death rates that ranges in millions annually [31]. The sub-Saharan African region has been the most affected because the malaria infection in this region has been greatly linked to the most effective and dangerous malaria parasite which is the *Plasmodium falciparum* and the most effective malaria vector —

44 the mosquito *Anopheles Gambiae* — which spreads the most and the control also has proven
45 difficult[20]. Deaths as a result of malaria is most common among children and in regions
46 where the malaria infection has been known to be common, a semi- immunity protection is
47 acquired in the early stages of life and that ranges between 10 to 15 years [8]. However,
48 contrary to the low malaria prevalence in the older generation, pregnant women in endemic
49 regions are major victims of malaria infection, and the disease is most frequent and severe in
50 pregnant women [21, 27]. In the state of pregnancy, there is an occurrence of a transient
51 reduction in immunity, which is cell-mediated, and the responses of cells antigens of the *P.*
52 *falciparum* antigens experiences depression in pregnant women.

53 Pregnancy related malaria is therefore a major challenge in the sub-Saharan Africa, and this
54 affects approximately 24 million pregnant women on yearly basis. Although the infection by
55 *P. falciparum* in the state of pregnancy is usually ~~asymptomatic~~ asymptomatic, it is a constant
56 ~~contributor~~ contribute to unfavourable perinatal results with a great risk for child mortality,
57 particularly in regions where malaria infection cases are less common[4, 11]. Pregnancies in
58 women who reside in areas known for a high malaria infection rates usually have a high *P.*
59 *falciparum* parasitemia density and frequency, with high rates of maternal morbidity, which
60 includes abortion, fever, severe anaemia, and placental malaria [13, 17]. Annually, figures
61 ranging between 75,000 and 200,000 infant deaths are recorded to malaria infection in
62 pregnancy worldwide.

63 **PLASMODIUM CYCLE IN THE VECTOR**

64 Just like many protozoa, plasmodia pass through some basic stages in the course of their two-
65 host life cycle. The infective stage for humans is the uninucleate, lancet-shaped
66 sporozoite[23]. The sexual reproduction produces the Sporozoites in the midgut of vector
67 anopheles mosquitoes and move to the salivary gland. When an *Anopheles* mosquito that is
68 already infected gets to bites a human, it injects along with saliva into small blood vessels the
69 sporozoites. Sporozoites have been observed to enter the liver parenchymal cells within 30
70 minutes of inoculation. In the cells of the liver, the parasite develops into a spherical,
71 multinucleate liver-stage schizont containing 2,000 to 40,000 uninucleate merozoites. This
72 process fast multiplication is regarded as exoerythrocytic schizogony. This exoerythrocytic
73 liver phase of the infection is normally completed between 5 and 21 days, depending on the
74 *Plasmodium* species. However, in infections related to *P vivax* and *P ovale*, there might be a
75 delay in the maturation of liver-stage schizonts for close to 1 to 2 years [15]. These inactive

76 liver-phase parasites are regarded as hypnozoites. Irrespective of the required time for
77 development, the mature schizonts gets to rupture eventually, and it releases into the
78 bloodstream thousands of uninucleate merozoites. Each merozoite has the ability to infect a
79 red blood cell. The merozoite in the red blood cells develops to form either an erythrocytic-
80 stage (blood-stage) schizont (by the process of erythrocytic schizogony) or a spherical or
81 banana-shaped, uninucleate gametocyte. The mature erythrocytic-stage schizont is made up of
82 8 to 36 merozoites, which are being released into the bloodstream at the rupturing of the
83 schizont. These merozoites go ahead to infect another red blood cell generation [15].

84 The *Plasmodium* sexual stage which is the gametocyte, is infectious for mosquitoes that
85 ingest it in the process of feeding. Within the mosquito, the development of the gametocytes
86 into gametocytes into female and male gametes (macrogametes and microgametes,
87 respectively) is observed and their fertilization commences for their development
88 into sporozoites which has the human infection ability and this takes place within a time frame
89 of ~~2 of 2~~ 2 to 3 weeks [1].

90 PATHOGENESIS

91 The cause of clinical illness is linked to the erythrocytic stage of the malaria parasite [2]. The
92 earliest signs and symptoms of malaria has to do with erythrocytes rupturing when
93 erythrocytic-stage schizonts mature. The release of these materials from the parasites is
94 assumed to trigger the immune response from the host. The cytokines, reactive oxygen
95 intermediates, and other cellular products released during the immune response play a
96 important role in pathogenesis, and might be responsible for the feeling of fever, chills,
97 sweats, weakness, and other symptoms that can be linked to malaria [18]. In cases of
98 *falciparum* linked malaria (the form leading to most ~~mortality~~ mortality), erythrocytes that has
99 been infected gets to adhere to the endothelium of capillaries and post capillary venules,
100 which leads to a stoppage in the microcirculation and anoxia in local tissues. Cerebral
101 malaria is caused in the brain through this. It also leads to acute tubular necrosis and renal
102 failure in the kidney; and ~~can also~~ can cause ischemia and ulceration in the ~~intestine~~
103 ~~which~~ intestine, which can lead to gastrointestinal bleeding. The parasite experiences
104 protection from attack by the body's immune system because it lives in the liver and blood
105 cells for most part of its human life cycle, and is relatively not visible to surveillance by the
106 immune system [18]. However, the spleen gets to destroy circulating infected blood cells. To
107 prevent this eventual occurrence, the *P. falciparum* parasite shows proteins with adhesive

108 attributes on the surface of infected blood cells, which causes the sticking of the blood cells
109 to small blood vessels walls, thereby inhibiting the parasite from passing through the general
110 circulation and the spleen [5].

111 MALARIA IN PREGNANCY

112 Pregnant women stands ~~three~~ 3 times more likely chance to suffer more from severe ailment
113 resulting from malarial infection when compared with their counterparts who are not
114 pregnant, and their rate of death from chronic disease also approaches 50% [7, 30]. In areas
115 where malaria infection is common, the estimation is that a minimum of 25% of pregnant
116 women are infected with malaria, with the group with the highest infection and morbidity risk
117 are the ~~primigravida~~ prim-gravidas, children, and co-infected HIV patients [22]. The highest
118 infection rate occurs during the second trimester, giving credence to the importance of
119 antenatal care as part of the effort in the prevention and treatment of malaria. There are
120 hypothesis that the most of the pregnancy consequences are ~~as a result~~ because of ~~two~~ 2 main
121 factors: the immune-compromised pregnancy state and cases of having infected red blood
122 cells being sequestered in the placenta [22]. Immunosuppression in pregnancy poses special
123 problems. It makes malaria more common and more severe. ~~And~~ In addition, to add to the
124 woes, malaria itself suppresses immune response. Hormonal changes of pregnancy, reduced
125 function of ~~reticulo~~ reticula-endothelial system are the causes of immunosuppression in
126 pregnancy. This results in loss of acquired immunity to malaria, making the pregnant more
127 prone for malaria [7].

128 CONGENITAL MALARIA

129 Malaria during pregnancy may result in fetal exposure to malaria if parasites are transmitted
130 across the placenta and could result in congenital malaria [25]. The most dangerous type of
131 malaria, *P. falciparum*, also seems very able to infect cells in the placenta, leading to a higher
132 intensity infection, ~~and also~~ and reducing oxygen delivery to the baby. This, combined with
133 the mother's illness and anaemia, can lead to low birth weight, anaemia and other
134 complications in the child once it is born [14]. *P. falciparum* has the unique ability of
135 cytoadhesion. Chondroitin sulphate A (CSA) has been identified as the adhesion molecule
136 for parasite ~~attachement~~ attachment to placental cells. The administration of soluble of CSA to
137 pregnant women has proven to drastically reduce parasite adhesion. The administration of
138 chondroitinase can effectively reduce parasite by 90% [16].

139 Malaria can also pass through the placenta, or be transferred to the baby through blood during
140 childbirth. Most babies are thought to remain unharmed if the mum-to-be has malaria, as long
141 as the malaria is treated promptly and effectively [29].

142 **EFFECT OF MALARIA ON THE FOETUS**

143 1. INTRA-UTERINE GROWTH RETARDATION: This may occur as a result of pyrexia
144 and transplacental infection in susceptible woman. Erythrocytes infected with *P.*
145 *falciparum* congregate in the maternal placental vascular space, where the parasites replicate.
146 Malaria-infected placentas are frequently observed to carry antibodies, cytokines, and
147 macrophages, which are indicative of an active immune response. This immune response may
148 stimulate early labour, though the precise effect of malaria-parasitized placentas on
149 prematurity is not clear [26]. The IUGR effect appears to relate to nutrient transport to the
150 foetus. First, a high density of parasites and chronic parasite infection in the placental blood
151 and the associated cellular immune response may result in consumption of glucose and
152 oxygen that would have gone to the foetus. Malaria-associated maternal anaemia may also
153 contribute independently to IUGR, most likely through a reduction in oxygen transport to the
154 foetus. Until recently, the mechanism through which parasite sequestration occurs in the
155 placenta has been unclear [27].

156 2. LOW BIRTH WEIGHT (LBW): Low birth weight is the single greatest risk factor for
157 neonatal and infant mortality. Low birth weights due to malaria may result from IUGR or
158 premature delivery, it can also be influenced by many factors, including genetics, multiple
159 pregnancies, placental abnormalities, maternal nutrition, maternal age, gravidity, and history
160 of smoking, and a range of viral, bacterial, and parasitic infections. Infant mortality is three
161 times higher for LBW babies than for those of normal weight. This is due to placenta
162 ~~parasitisation which~~parasitisation, which interferes with placenta blood circulation and impairs
163 the growth of the foetus[9].

164 3. NEONATAL DEATH: Congenital malaria may cause death in the neonatal period of the
165 baby of a susceptible woman but it is very rare in endemic area because the ~~antibody which~~
166 ~~fights against malaria~~antibody that fights against malaria crosses the placenta and the infant
167 becomes passively immunized [28].

168

169 | **THE-MALARIA-PARASITE'S MECHANISM FOR EVADING THE IMMUNE**
170 **SYSTEM**

171 *Plasmodium*, the parasite responsible for malaria, infects red blood cells. It produces proteins
172 | in the red blood cells that bind to the surface of the host cell. These are known as adhesion
173 proteins [30]. They prevent the red blood cells from circulating correctly in the blood
174 capillaries, and trigger the symptoms of severe malaria. The parasite has 60 genes coding for
175 60 different adhesion proteins, only one of which appears on the surface of the red blood cell
176 | at any one time. In this wayway, the various adhesion proteins are presented in turn, and the
177 parasite keeps one step ahead of the host's immune system, which must learn to recognize
178 and then destroy infected cells [6].

179 **CONTROL OF MALARIA IN PREGNANCY**

180 Intermittent Preventive Treatment in Pregnancy. WHO recommends IPTp with sulfadoxine-
181 pyrimethamine (IPTp-SP). In September 2012, the WHO Malaria Policy Advisory Committee
182 reviewed the most recent evidence on efficacy and effectiveness of IPTp-SP and issued new
183 policy recommendations that promote the increased uptake of IPTp-SP in all areas of Africa
184 with moderate-to-high transmission of *Plasmodium falciparum* malaria.

185 Use of insecticide treated nets (ITNs). The second component of WHO's prevention
186 approach, the use of ITNs, benefits pregnant women and their families. In areas of stable
187 transmission, ITNs reduce the risk of malaria, which in turn produces significant protection
188 against maternal anaemia and low birth weight [10].

189 Prompt diagnosis and case management of malaria illness. Malaria case management is
190 another essential component of malaria control during pregnancy. Pregnant women with
191 symptomatic malaria are at higher risk of foetal loss, premature delivery, and death, and they
192 need urgent treatment. The goal in treatment of malaria during pregnancy is to cure the
193 infection completely; any level of parasitemia has consequences for mother and foetus [12].

194

195 **TREATMENT OF MALARIA**

196 It has been recommended by the World Health Organization (WHO) now recommends that
197 women in experiences subtle *P. falciparum* malaria in their second or third trimester period of
198 pregnancy should be treated with artemisinin-based combination therapy. In as much as the

199 | therapy seems to be ~~short-acting~~ short acting, it remains potent and effective due to
200 | the artemisinin component (i.e., artemether, artesunate, or dihydroartemisinin) which greatly
201 | reduces the parasite number during the first 3 days of administration [10]. The more
202 | sustainable partner drug (i.e., lumefantrine, piperaquine, amodiaquine, or mefloquine) which
203 | acts longer removes the remaining parasites, and as such preventing recrudescence malaria.
204 | The longer-acting partner drug is also has also been implicated as being responsible for the
205 | prophylactic effect that occurs after treatment, and this blocks the chances of new infections
206 | while concentration of the drug in the blood exceeds the minimum required for the inhibition
207 | of the parasite. Thus, the time range of the post-treatment prophylactic effect is as a result of
208 | the potency and half-life for the elimination of the drug [7]. The same mode of action is used
209 | in the preventive treatment, where repeated curative anti-malarial treatments removes
210 | potential asymptomatic infections ~~and also~~ and blocks the possibility of the occurrence of new
211 | infections. However, artemisinin-based combination therapy is not currently recommended in
212 | pregnancy for intermittent preventive treatment. The current recommendation from the WHO
213 | is for all women in regions with high risk of getting infected with malaria in Africa to receive
214 | intermittent preventive treatment with sulfadoxine-pyrimethamine as part of their antenatal
215 | care [30].

216

217 **CONCLUSION**

218 | Malaria is increasingly becoming one of the hardest infectious diseases to eradicate in Africa.
219 | The overall burden of the infection is devastating youth, women, and the health systems as a
220 | whole. It has affected human resources of Africa and lowered directly the economic growth
221 | that should have been experienced annually. It not only weakens the workforce, but also stops
222 | children from learning in school, prevents pregnant women from taking good care of their
223 | families effectively, and reduces the chances of having a health outcome after
224 | pregnancy. Malaria during pregnancy results in foetal exposure to malaria parasite. This
225 | combined with the mother's illness can lead to low birth weight, anaemia another
226 | complications in the child once it is born. Governments and donors have recognized this
227 | extraordinary toll and have put in more commitment towards the prevention, treatment, and
228 | eradication of the disease. The reduction in the ITN tariffs thereby making them cheaper and
229 | more affordable has also been successful, including the incorporation of programs to sensitize
230 | against infectious disease in reproductive health, and intermittent preventive treatment.

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