Changes in the total leukocyte and platelet counts in Papuan and non-Papuan adults from northeast Papua infected with acute Plasmodium vivax or uncomplicated Plasmodium falciparum malaria

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Abstract

Background: There are limited data on the evolution of the leukocyte and platelet counts in malaria patients.

Methods: In a clinical trial of chloroquine vs. chloroquine plus doxycycline vs. doxycycline alone against Plasmodium vivax (n = 64) or Plasmodium falciparum (n = 98) malaria, the total white cell (WCC) and platelet (PLT) counts were measured on Days 0, 3, 7 and 28 in 57 indigenous Papuans with lifelong malaria exposure and 105 non-Papuan immigrants from other parts of Indonesia with limited malaria exposure.

Results: The mean Day 0 WCC (n = 152) was 6.492 (range 2.1–13.4) × 10⁹/L and was significantly lower in the Papuans compared to the non-Papuans: 5.77 × 10⁹/L vs. 6.86 × 10⁹/L, difference = -1.09 [(95% CI -0.42 to -1.79 × 10⁹/L), P = 0.0018]. 14 (9.2%) and 9 (5.9%) patients had leukopaenia (<4.0 × 10⁹/L) and leukocytosis (>10.0 × 10⁹/L), respectively. By Day 28, the mean WCC increased significantly (P = 0.0003) from 6.37 to 7.47 × 10⁹/L (73 paired values) and was similar between the two groups. Ethnicity was the only WCC explanatory factor and only on Day 0.

The mean Day 0 platelet count (n = 151) was 113.0 (range 8.0–313.0) × 10⁹/L and rose significantly to 186.308 × 10⁹/L by Day 28 (P < 0.0001). There was a corresponding fall in patient proportions with thrombocytopenia (<150 × 10⁹/L): 119/151 (78.81%) vs. 16/73 (21.92%, P < 0.00001). Papuan and non-Papuan mean platelet counts were similar at all time points. Only malaria species on Day 0 was a significant platelet count explanatory factor. The mean D0 platelet counts were significantly lower (P = 0.025) in vivax (102.022 × 10⁹/L) vs. falciparum (122.125 × 10⁹/L) patients.

Conclusion: Changes in leukocytes and platelets were consistent with other malaria studies. The Papuan non-Papuan difference in the mean Day 0 WCC was small but might be related to the difference in malaria exposure.
Background
The malaria induced changes in the total white cell (WC) and platelet (PLT) counts have been documented in several clinical series of vivax and falciparum malaria in malaria immune and naïve patients of all ages [1-9], describing mostly the WC and PLT counts at presentation.

The majority of malaria patients at presentation have total WCCs in the normal range. Leukopenia may affect up to 15% (WCC < 4 × 10^9/L) of malaria infected adults [3,4,8,10] and was present in ~10% of 1,369 hospitalized Kenyan children with moderately severe or severe falciparum malaria (WCC < 6.1 × 10^9/L) [6]. Rates of leukopenia (>10 × 10^9/L in adults range between 1 and ~7% [1,3,4,9,11,12] and was 20% in the hospitalized Kenyan children (>16.5 × 10^9/L) [6]. Reported, malaria induced changes in the differential white cell counts include neutropenia, neutrophilia, immature neutrophils (left shift), neutrophil toxic granulation, lymphopenia, lymphocytosis, atypical lymphocytes, monocytosis, eosinopenia, post treatment eosinophilia, and leukemoid reactions [3-5,10,11,13-19]. The underlying mechanisms include a shift in neutrophils from the circulatory to the marginal pool to sites of inflammation, splenic localisation, serum lympho toxic factors, and intercurrent bacterial infections [13,15,20-23]. In experimentally induced falciparum malaria, Aotus monkeys had an increase in the absolute neutrophil counts seven days post infection that was followed by neutropenia by Day 14; the reverse was found for the lymphocyte counts [24]. Ethnicity, sex and immune status may also be relevant factors affecting the leukocyte count. Leukocytosis was more common in Plasmodium vivax infected African American soldiers (27%) compared to Caucasian soldiers (9%) but the latter had higher rates (47 vs. 26%) of relative lymphocytosis (lymphocyte count > 35%) [14]. Studies in temperate climates have found that individuals of African origin tend to have lower total leukocyte, neutrophil and platelet counts compared to Caucasians and women have higher leukocyte and neutrophil counts than men [25,26]. Malaria naive Sri Lankans infected with P. vivax had higher mean lymphocyte counts and significantly raised gamma/delta T cells during fever paroxysms compared to malaria immune vivax patients, suggesting a greater inflammatory response by the non immune patients [27]. Similarly, TNFα levels were greater in non immune Sri Lankans with P. vivax [28].

Studies examining trends in the total WCC are few. One study of experimentally induced falciparum malaria in non immune Americans showed an initial fall in the mean WCC reaching a nadir on Day 3 and rising thereafter to baseline values by study end [2]. Studies in African children have shown modest falls in the mean total WCC over 7 to 14 days whilst another study found that acutely ill falciparum infected children had a significant fall in the mean neutrophil count on Day 3 [29-31]. Malaria induced thrombocytopenia (platelet count < 150 × 10^9/L) is very common, affecting between 40–85% of patients [4,7,9,32-35]. Moderate (50–100 × 10^9/L) and severe (<50 × 10^9/L) thrombocytopenia occur in 30–50 and 6–16%, respectively [4,5,7,30,35]. Thrombocytopenia improves with disease resolution and the platelet count is generally normal within seven days [34,35] but ranged from 2–28 days in one series [35]. Considerable overlap exists between the platelet counts of the different malaria species and between uncomplicated and severe falciparum malaria. Two studies suggest that a lower platelet count may be an indicator of higher biomass falciparum and vivax infections [36,37] but Eriksson et al did not find this correlation [4]. Factors associated with malaria induced thrombocytopenia include splenomegaly, splenic sequestration and platelet removal by macrophages [38-40]. The role of antiplatelet antibodies is unclear [34,41,42].

In north east Papua, indigenous Papuans have life long exposure to malaria whereas Indonesians who immigrate to Papua usually contract malaria for the first time in Papua [43]. Whether this difference in malaria exposure affects the white cell and platelet counts in diseased patients is unknown. This paper presents data from a clinical trial on the evolution of the total white cell and platelet counts and explores possible Papuan non Papuan differences.

Materials and methods
Malarialometric data from this area of northeast Papua and study conduct are detailed elsewhere [44,45]. Briefly, indigenous adult Papuans (n = 57) and non Papuan (n = 105) adults from other parts of Indonesia were recruited into a hospital based, 28 days, clinical trial comparing chloroquine vs. chloroquine plus doxycycline vs. doxycycline for the treatment of parasitologically proven (Giemsa stained thick and thin blood films) acute uncomplicated falciparum and acute vivax malaria. There was no minimum parasitaemia for either species but a maximum of 150,000/μL asexual falciparum forms/μL for study entry. Recruited patients underwent a history, a detailed physical examination (including an abdominal examination to detect hepatosplenomegaly) and supervised treatment. Giemsa stained blood films were taken and read on Days 0–7, 14, 21, 28 and a blood count (Hb, total WCC, platelet count) was performed on Days 0, 3, 7, and 28; a manual differential white cell count was not done. Double entered, validated data were analysed using Epi Info 6.04b (Centers for Disease Control and Prevention, Atlanta, GA, USA) and Stata v 8 (Stata Corporation, USA). Normally distributed data were compared using the stu-
dent's 't' test or ANOVA or the Mann-Whitney U or Kruskall-Wallis for skewed data. Spearman's rank correlation was used to determine the relationship between skewed continuous data. Proportions were compared using uncorrected chi squared values. Multivariate analyses were performed to assess the independence of ethnicity, splenomegaly, gender, malaria species and drug regimen on the total white cell and platelet counts. A P value of ≤ 0.05 was considered significant; no adjustment was made for multiple comparisons. Written informed consent was obtained from all patients. The study was conducted according to the Indonesian Ministry of Health, the Indonesian Navy, and the United States Navy and Army regulations governing the protection of human subjects.

**Results**

A total of 162 malaria infected patients were recruited into the study: *P. vivax* (n = 63), *P. falciparum* (n = 89), and mixed infections (n = 10). Based on the dominant species, the 10 mixed infections were reclassified as vivax (n = 1) or falciparum (n = 9) malaria. Males (aged 15 to 44), numbered 146 (~90%), and women (aged 15 to 33) 16 (~10%). The Papuans numbered 57; the 105 non Papuans came from Java (n = 54), Sulawesi (n = 28), Ambon (n = 9), Bali (n = 7), Sumatera (n = 5), and other islands (n = 2). Median Papuan and non Papuan [interquartile (IQ) range] residential times in Papua were 21.5 (18 to 24) and three (1 to 5) years, respectively (P < 0.0001). 152/162 (93.8%) had blood results available for Day 0 and 74 patients completed the study to Day 28. Splenomegaly was more common in the Papuans, 26/57 (45.6%), compared to the non Papuans, 27/105 (25.7%), for a relative risk of 1.8 [95% confidence interval (CI) 1.15 to 2.7].

**Day 0 total white cell counts**

The Day 0 total WCCs ranged from 2.1–13.4 × 10⁹/L with a median and mean of 6.3 and 6.492 × 10⁹/L, respectively. Leukopenia (WCC < 4 × 10⁹/L) was present in 14 patients (9.2%); their WCCs ranged from 2.1–3.9 × 10⁹/L. Nine (5.9%) patients had a leukocytosis (WCC > 10 × 10⁹/L). Thus, the majority of patients, ~85%, had a normal total WCC at presentation. In the multivariate analysis, only ethnicity was a significant explanatory variable for the total WCC on Day 0. The mean Day 0 total WCC was significantly lower in the Papuans (5.7 × 10⁹/L) compared to the non Papuans (6.8 × 10⁹/L); difference = -1.09 [95% CI -0.42 to -1.79 × 10⁹/L], P = 0.0018. There was a positive correlation between the Day 0 WCC and the *P. vivax* parasitaemia (Sp rho = 0.43, P = 0.0003) but no correlation between the Day 0 WCCs and: (i) the Day 0 falciparum counts [Spearman’s (Sp) rho = 0.139, P = 0.19], (ii) the Day 0 temperature (Sp rho = -0.0066, P = 0.94), and (iii) the Day 0 platelet count (Sp rho = 0.125, P = 0.12).

**Evolution of the total white cell counts**

The changes in the mean white cell counts during follow up were similar for both malaria species and ethnic groups (Figures 1 and 2). Compared to baseline, the fall in the mean total WCC on Day 3 was significant in the non Papuans: (i) Pf: -0.898 (5.735 – 6.633) × 10⁹/L (P = 0.0015), (ii) Pf: -1.069 (6.132 – 7.201) × 10⁹/L (P = 0.0012). The mean Day 3 Day 0 difference between the non Papuans and Papuans was also significantly different.

![Figure 1](http://www.malariajournal.com/content/7/1/259)

**Figure 1**

The mean (95% CI) total white cell counts in falciparum infected Papuan and non Papuan patients.
By Day 28, the number of evaluable patients had fallen to 74. The mean total WCC was: (i) significantly higher compared to baseline (n = 73 pairs) 7.47 vs. 6.37 × 10^9/L: difference = 1.10 (0.52 – 1.69) × 10^9/L, P = 0.0003] and (ii) similar between the Papuans [7.79 × 10^9/L (n = 27)] and the non Papuans [7.30 × 10^9/L (n = 47)] but there was a trend towards a higher mean change in the Papuans compared to the non Papuans: 1.8 vs. 0.7 × 10^9/L (P = 0.07).

There was no Day 28 leukopaenia but five (6.7%) patients had leukocytosis. Their Day 28 Day 0 matching values (x 10^9/L) were: 10.4–3.4, 10.8–6, 12–7.2, 12.2–9.2, 16.4–7. They were all aparasitaemic, afebrile, and discharged clinically well from the hospital.

**Day 0 platelet counts**

The Day 0 platelet counts (PLT0) for all patients combined varied widely, ranging from 8–313 × 10^9/L (IQ range 76 to 141 × 10^9/L); the median and mean values were 102 and 113 × 10^9/μL, respectively. The majority of patients, 119/151 (78.8%), had thrombocytopaenia (<150 × 10^9/L); 43.7% (66/151) had moderate thrombocytopaenia (50–100 × 10^9/L). Five *P. falciparum* and eight *P. vivax* (8.6%) infected patients were severely thrombocytopaenic (<50 × 10^9/μL) but none had clinical signs of bleeding. Of the 88 patients who had urine dipstick tests, nine (10.2%) had microscopic haematuria (*vivax* = 4, *falciparum* = 5); the platelet counts in these nine ranged from 50–313 (median 101) × 10^9/L.

The multivariate analysis showed that only the malaria species was a significant independent variable to explain the Day 0 platelet count. The mean PLT0 was significantly lower (P = 0.025) in the *vivax* patients: 102.022 vs. 122.125 × 10^9/L. The PLT0 was negatively correlated with the Day 0 *falciparum* parasitaemia (Sp rho = -0.29, P = 0.005) but not with the Day 0 *vivax* parasitaemia (Sp rho = -0.0984, P = 0.4). There was also a significant negative correlation with the Day 0 temperature (Sp rho = -0.21, P = 0.009) and the PLT0 in the *vivax* patients (Sp rho = -0.26, P = 0.04) and a negative trend in *falciparum* patients (Sp rho = -0.18, P = 0.097).

**Evolution of the platelet counts**

The proportion of thrombocytopaenic patients declined over time: 78.8% (119/151) on D0, 74.26% (101/136) on D3, 23.62% (30/127) on D7 and 21.92% (16/73) on Day 28 (P < 0.0001, chi^2 for trend). By Day 28, the mean platelet count was significantly higher compared to baseline: 188.308 vs. 108.056 × 10^9/L (P < 0.0001), and ranged from 47.0 to 318.2 (IQ range 153 to 226) × 10^9/L (Figures 3, 4 and 5). The mean D28-D0 changes in platelet counts were not significantly different between the Papuans and non Papuans: 83.280 vs. 75.576 × 10^9/L (P = 0.6). The mean Day 7 platelet count was significantly higher than that of Day 28 only in the *vivax* infected non Papuans (n = 21): 175.819 vs. 251.238 × 10^9/L (P = 0.0036). The multivariate analyses were unremarkable, except for drug arm as a PLT7 explanatory factor. This was accounted for by the combination vs. the doxycycline arm: 235.463 vs 191.741 × 10^9/L (P = 0.029).
Discussion
These analyses have shown that at presentation the majority of malaria infected patients had total white cell counts within the normal range and most were thrombocytopenic. By Day 28, both mean counts had increased but thrombocytopenia remained in just over 20% of patients. These data complement and are broadly consistent with those of other clinical series.

Data from this study were from two dissimilar groups with respect to ethnicity and the degree of malaria acquired immunity. The Papuans had life long malaria exposure whilst the non Papuans had limited or no malaria exposure. The Papuans had lower mean Day 0 white counts than the non Papuans and there was a notable difference in the WCC changes over time. They rose steadily in the Papuans but fell significantly on Day 3 in the non Papuans, consistent with a study of experimen-
tally induced falciparum malaria in malaria naïve American volunteers [2]. By Day 28, the mean WCCs in the non Papuans had caught up and both groups had similar mean WCCs. This difference in the WCC progressions between the two ethnic groups is difficult to explain purely on the basis of a redistribution of white cells and/or splenic sequestration, given that the malaria naïve non Papuans had significantly less splenomegaly. These data might indicate that a lesser degree of malaria exposure, thus acquired immunity, in the non Papuans resulted in a more robust inflammatory response, reflected crudely by a higher, baseline total WCC [27]. The initial fall in the total WCC in the non Papuans might reflect a difference in response to dying parasitized red cells consequent to treatment.

Thrombocytopaenia was common at presentation in both groups of patients. There were no differences between the Papuans and non Papuans but the mean platelet count was lower in vivax compared to falciparum infected patients. There were negative correlations between the Day 0 platelet count and the falciparum parasite count and the Day 0 temperature in the vivax patients. These findings are consistent with some but not other reports and are of limited clinical value [2,4,8,31,34]. Over time, the mean platelet count increased, consistent with the findings of others but two reports document an initial fall in the platelet count followed by recovery [2,4]. The mean platelet count appeared higher on Day 7 compared to Day 28 but there are no intervening platelet counts, so the day of the mean peak platelet count cannot be determined. Some 20% of patients were still thrombocytopenic by study end, suggesting a number of patients need longer than four weeks for full platelet recovery. The multivariate analysis found drug arm to be significant factor for the Day 7 platelet count with the combination arm having a higher mean platelet count than the doxycycline arm. This is of little clinical significance.

Our study had limitations. The study sample was small and powered for a clinical trial. The data presented were secondary analyses to look for possible Papuan non Papuan differences. Many patients failed treatment before Day 28, given rescue treatment and withdrawn from the study; this may have introduced a statistical bias. Most patients were male, thus limiting the applicability of the findings. There were many statistical comparisons and some significant results may have occurred by chance. The differential white cell count was not measured and could have provided additional interesting data.

**Conclusion**

To conclude, these analyses have compared and contrasted the changes in the total white cell and platelet counts in Papuans and non Papuans with different degrees of malaria exposure that may partly explain the difference in the mean total white cell count at presentation between the two groups.
Authors’ contributions
WRJT, the principal investigator, developed the protocol, supervised study execution, analysed the data and wrote the first draft of the paper. CO conceived the study, developed the protocol and critically reviewed the manuscript. ET developed the protocol, was involved in study execution and critically reviewed the manuscript. SLH developed the protocol and critically reviewed the manuscript. HW played a key role in study execution and reviewed the manuscript. HLB developed the protocol and critically reviewed the manuscript. TLR developed the protocol and critically reviewed the manuscript. HW played a key role in study execution and reviewed the manuscript. TAfk was involved in study execution and reviewed the manuscript. SB was involved in study execution and reviewed the manuscript. DF was involved in data interpretation and critically reviewed the manuscript.

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References
1. Martelo OJ, Smoller M, Saladin TA: Malaria in American soldiers. Arch Intern Med 1969, 123:383-387.
2. Church LW, Le TP, Bryan JP, Gordon DM, Edelman R, Fries L, Davis JR, Herrington DA, Clyde DF, Shmuklarsky MJ, Schneider I, McGovern TW, Chubay JD, Ballou WR, Hoffman SL. Clinical manifestations of Plasmodium falciparum malaria experimentally induced by mosquito challenge. J Infect Dis 1997, 175:915-920.
3. Goldstein E: A clinical study of falciparum and vivax malaria in Vietnam servicemen. Mil Med 1968, 133:991-6.
4. Eriksson B, Hellgren U, Rombo L: Changes in erythrocyte sedimentation rate, C-reactive protein, and haematological parameters in patients with acute malaria. Scand J Infect Dis 1989, 21:435-41.
5. Stein CM, Gelfand: The clinical features and laboratory findings in acute Plasmodium falciparum malaria in Harare, Zimbabwe. Cent Afr J Med 1985, 31(9):166-170.
6. Ladhani S, Lowe B, Cole AO, Kowouno K, Newton CJ: Changes in white blood cells and platelets in children with falciparum malaria: relationship to disease outcome. Br J Haematol 2002, 119(3):839-847.
7. Lynk A, Gold R: Review of 40 children with imported malaria. Pediatr Infect Dis J 1989, 8:745-50.
8. Svenson JE, MacLean JD, Gyorkos TW, Keystone J: Imported malaria. Clinical presentation and examination of symptomatic travelers. Arch Intern Med 1995, 155:861-8.
9. Winters RA, Murray HW: Malaria – the mime revisited; fifteen years of experience at a New York city teaching hospital. Am J Med 1992, 243:243-6.
10. Jadhav UM, Singhvi R, Shah R: Prognostic implications of white cell differential count and white cell morphology in Malaria. J Postgrad Med 2003, 49:218-20.
11. Reiley CG, Barrett O: Leukocyte response in acute malaria. Am J Med Sci 1971, 262:153-8.
12. Sharma SK, Das RK, Das BK, Das PK: Haematological and coagulation profile in acute falciparum malaria. J Assoc Physicians India 1992, 40:581-3.
13. Dale DC, Wolf: Studies of the neutropenia of acute malaria. Blood 1973, 41:197-206.
14. Fisher GU, Gordon MH, Lobel HO, Runcik K: Malaria in soldiers returning from Vietnam. Am J Trop Med Hyg 1970, 19:27-39.
15. Hvidl, Kemp K: What is the cause of lymphopenia in malaria? Infect Immun 2000, 68:6087-9.
16. Kueh YK, Yeo KL: Haematological alterations in acute malaria. Scand J Haematol 1982, 29:147-52.
17. Shanks GD, Wilarataporn C: Eosinophilic response to falciparum malaria infections. Southeast Asian J Trop Med Public Health 1992, 23:795-7.
18. Irving KG, Kemp E, Olivier BJ, Mendelow BV: Unusual presentations of malaria in soldiers. Mil Med 1989, 154:319-21.
19. Bondoncourcelle D, Gaulard P, Mason DY: Preferential localisation of human lymphocytes bearing gamma delta T cell receptors to the red pulp of the spleen. J Clin Pathol 1990, 43:461-4.
20. Langhorne J, Simon-Haarsch B: Differential T cell responses to Plasmodium chabaudi chabaudi in peripheral blood and spleens of C57BL/6 mice during infection. J Immunol 1991, 146:2771-5.
21. Kakoma I, James MA, Whiteley HE, Montelegre F, Buese M, Fajfar-Whestone CJ, Clabaugh GW, Baek BK: Platelet kinetics and other hematological profiles in experimental Plasmodium falciparum infection: a comparative study between Saimiri and Aotus monkeys. Kisoenongchukhapchi 1992, 30:177-82.
22. Freedman DS, Gates L, Flanders WD, Van Assendelft OW, Barberiak JJ, Joesof MR, Byers T: Black/white differences in leucocyte subpopulations in men. Int J Epidemiol 1997, 26:577-64.
23. Bain BJ: Ethnic and sex differences in the total and differential white cell count and platelet count. J Clin Pathol 1996, 49:664-6.
24. Perera MK, Carter R, Gooenwardene R, Mendis KN: Transient increase in circulating gamma/delta T cells during Plasmodium vivax malarial paroxysms. J Exp Med 1994, 179:31-5.
25. Karunaweera ND, Carter R, Grau GE, Mendis KN: Demonstration of anti-disease immunity to Plasmodium vivax malaria in Sri Lanka using a quantitative method to assess clinical disease. Am J Trop Med Hyg 1998, 58:204-10.
26. Penali LK, Assi-Coulibaly L, Kapve B, Konan D, Ehouman A: Parasitological and clinical response to amodiaquine versus chloroquine in the treatment of Plasmodium falciparum malaria in children in an endemic area. Bull Soc Pathol Exot 1994, 87:244-7.
27. Staedke SG, Kanya MR, Dorsey G, Gasasira A, Ndegei G, Charlebois ED, Rosenthal PJ: Amodiaquine, sulfadoxine/pyrimethamine, and combination therapy for treatment of uncomplicated falciparum malaria in Kampala, Uganda: a randomised trial. Lancet 2001, 358:638-74.
28. Abdalla SH: Peripheral blood and bone marrow leucocytes in Gambian children with malaria: numerical changes and evaluation of phagocytosis. Ann Trop Paed 1988, 8(4):250-258.
29. Newton JA, Schepf GA, Wallace MR, Lobel HO, Kennedy CA, Oldfield EC: Malaria in US marines returning from Somalia. JAMA 1982, 247(5):397-399.
30. Oh MD, Shin H, Shin D, Kim U, Lee S, Kim N, Choi MH, Chai JY, Choe K: Clinical features of falciparum malaria. Am J Trop Med Hyg 2001, 65:143-6.
31. Looareesuwan S, Davis JG, Allen DL, Lee SH, Bunnag D, White NJ: Thrombocytopenia in malaria. Southeast Asian J Trop Med Public Health 1992, 23(1):44-50.
32. Moulin F, Lesage F, Legros A-H, Maroga C, Moussavou A, Guyon P, Marc E, Gentrel D: Thrombocytopenia and Plasmodium falciparum malaria in children with different exposures. Arch Dis Child 2003, 88:540-1.
36. Richards MW, Behrens RH, Doherty JF: Short report: hematologic changes in acute, imported Plasmodium falciparum malaria. Am J Trop Med Hyg 1998, 59:859.
37. Erhart LM, Yingyuen K, Chuanak N, Buathong N, Laoboontchai A, Miller RS, Meshnick SR, Gasser RA Jr, Wongsrichanalai C: Hematologic and clinical indices of malaria in a semi-immune population of western Thailand. Am J Trop Med Hyg 2004, 70:8-14.
38. Nazi GA: Haematological aspect of malaria in a population based hospital, Saudi Arabia. J Egypt Soc Parasitol 1995, 25:787-93.
39. Skudowitz RB, Katz J, Lurie A, Levin J, Metz J: Mechanisms of thrombocytopenia in malignant tertian malaria. BMJ 1973, 2:515-8.
40. Essien E: The circulating platelet in acute malaria infection. Br J Haematol 1989, 72:589-90.
41. Yamaguchi S, Kubota T, Yamagishi T, Okamoto K, Izumi T, Takada M, Kanou S, Suzuki M, Tsuchiya J, Naruse T: Severe thrombocytopenia suggesting immunological mechanisms in two cases of vivax malaria. Am J Hematol 1997, 56:183-6.
42. Beale PJ, Cormack JD, Oldrey TBN: Thrombocytopenia in malaria with immunoglobulin (IgM) changes. BMJ 1972, 1:34-9.
43. Baird JK, Jones TR, Danudrigo EW, Annis BA, Bangs MJ, Basri H, Purnomo, Masbar S: Age-dependent acquired protection against Plasmodium falciparum in people having two years exposure to hyperendemic malaria. Am J Trop Med Hyg 1991, 45:65-76.
44. Taylor WR, Widjaja H, Richie TL, Basri H, Obre C, Tjitra, Taufik E, Jones TR, Kain KC, Hoffman SL: Chloroquine/doxycycline combination versus chloroquine alone, and doxycycline alone for the treatment of Plasmodium falciparum and Plasmodium vivax malaria in northeastern Irian Jaya, Indonesia. Am J Trop Med Hyg 2001, 64:223-8.
45. Jones TR, Baird K, Bangs MJ, Annis BA, Purnomo, Basri H, Gunawan S, Harjosuwarno S, McElroy PD, Hoffman SL: Malaria vaccine study site in Irian Jaya, Indonesia: Plasmodium falciparum incidence measurements and epidemiological considerations in sample size estimation. Am J Trop Med Hyg 1994, 50:210-18.