Iron Deficiency Protects Against Severe *Plasmodium falciparum* Malaria and Death in Young Children

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(See the Editorial Commentary by Awah and Kaneko, on pages 1145–7.)

**Background.** Iron supplementation may increase malaria morbidity and mortality, but the effect of naturally occurring variation in iron status on malaria risk is not well studied.

**Methods.** A total of 785 Tanzanian children living in an area of intense malaria transmission were enrolled at birth, and intensively monitored for parasitemia and illness including malaria for up to 3 years, with an average of 47 blood smears. We assayed plasma samples collected at routine healthy-child visits, and evaluated the impact of iron deficiency (ID) on future malaria outcomes and mortality.

**Results.** ID at routine, well-child visits significantly decreased the odds of subsequent parasitemia (23% decrease, \( P < .001 \)) and subsequent severe malaria (38% decrease, \( P = .04 \)). ID was also associated with 60% lower all-cause mortality (\( P = .04 \)) and 66% lower malaria-associated mortality (\( P = .11 \)). When sick visits as well as routine healthy-child visits are included in analyses (average of 3 iron status assays/child), ID reduced the prevalence of parasitemia (6.6-fold), hyperparasitemia (24.0-fold), and severe malaria (4.0-fold) at the time of sample collection (all \( P < .001 \)).

**Conclusions.** Malaria risk is influenced by physiologic iron status, and therefore iron supplementation may have adverse effects even among children with ID. Future interventional studies should assess whether treatment for ID coupled with effective malaria control can mitigate the risks of iron supplementation for children in areas of malaria transmission.

Understanding the influence of iron status on the risk of malaria is necessary for planning and implementing iron deficiency (ID) control programs in sub-Saharan Africa. Both ID and malaria are common in this region and are major causes of anemia. ID anemia is associated with impaired cognitive and motor development, reduced growth velocity, and anorexia in children [1, 2]. International guidelines recommend iron and folic acid supplementation in children under 2 years of age in areas with a high prevalence of anemia. Consequently, iron supplementation programs are being implemented in many countries as a primary strategy for preventing ID and anemia in pregnant women and in children.

Although the detrimental effects of ID argue for aggressive intervention, the safety of universal routine iron supplementation remains unclear. Children in a malaria-endemic region of Tanzania who were randomized to receive iron supplements suffered from a 15% increased all-cause mortality [3]. Whereas results from several earlier studies also identified increased malarial risk in individuals treated with...
iron [4, 5], other studies have not identified risk associated with iron supplementation programs [6–8].

We recently observed that maternal ID is associated with a 5.5-fold reduced prevalence of placental malaria [9], but few studies have examined the role of iron status in modifying malaria risk in unsupplemented children [10]. In the current study, we evaluated the hypothesis that naturally occurring ID confers protection from malaria outcomes in a birth cohort of Tanzanian children. We report that ID protects children from malaria infection, malaria morbidity, and mortality.

MATERIALS AND METHODS

Ethical Approval
Ethical clearance was obtained from the Institutional Review Board of Seattle Biomedical Research Institute and the Medical Research Coordinating Committee of the National Institute for Medical Research, Tanzania.

Study Population
Subjects participated in the Mother–Offspring Malaria Studies (MOMS) project, which is based at Muheza Designated District Hospital (DDH) in northeastern Tanzania. Mothers presenting at Muheza DDH for delivery were enrolled and provided signed, informed consent for themselves and for their newborns before participation in the study. Details of the MOMS study design, enrollment methods, and exclusion criteria have been published elsewhere [9, 11].

Inclusion Criteria and Clinical Monitoring
We monitored 785 children for Plasmodium falciparum infection from birth up to 3 years of age. Children were evaluated at routine, well-child visits by a clinician every 2 weeks from birth to 1 year of age, then monthly thereafter, including blood smear analysis. Routine blood samples were collected at 3, 6, and 12 months of age, then once every 6 months in the second and third years of life. Blood smears and blood samples were also collected any time the child became sick. Sick children were examined by a medical officer upon presentation to the hospital or mobile clinic. Treatment outside the study was minimized by active, weekly surveillance by our mobile clinics.

Clinical malaria was defined as asexual P. falciparum parasitemia by blood smear coupled with symptoms suggestive of malaria such as temperature >37.5°C, nausea or vomiting, irritability, and poor feeding. Prompt treatment was provided to sick children according to the guidelines of the Tanzanian Ministry of Health, and study participants were instructed to obtain all medications including antimalarials through the project staff.

Sample Collection and Processing
Venous blood was collected and stored at 4°C until processing. After centrifugation, plasma was stored at −80°C. P. falciparum parasitemia was determined by Giemsa-stained thick blood smears prepared from capillary or venous blood. Parasite density was expressed as the number of asexual stage parasites/200 white blood cells (WBCs) in the thick smear. Sickle cell trait was determined by electrophoresis (Helena Laboratories, Beaumont, TX). Ferritin, C-reactive protein (CRP), and soluble transferrin receptor (sTfR) were assayed in plasma as described previously [12]. Hemograms were obtained on an impedance-based analyzer (Abbott Cell Dyne 1200).

Case Definitions
Severe malaria was defined according to the World Health Organization criteria [13] as a positive blood smear and one or more of the following: (1) respiratory distress defined by respiratory rate of >40/minute for children older than 2 months of age or a respiratory rate of >50/minute for children less than 2 months of age, in conjunction with clinical signs of respiratory distress; (2) a history of 2 or more convulsions in the 24 hours before or during hospitalization; (3) prostration defined by inability to sit unaided; (4) hypoglycemia defined by glucose <2.2 mmol/L; or (5) severe anemia defined by Hgb <5 g/dL. Hyperparasitemia was defined as parasitemia >2500/200 WBC.

Malaria-associated mortality was defined as death with a positive blood film obtained during the terminal illness. One child who died of bacterial meningitis but had a positive blood film was adjudicated as a nonmalarial death.

ID was defined as ferritin concentration <30 ng/mL when CRP was <8.2 µg/mL (ID in the absence of inflammation) or ferritin concentration <70 ng/mL when CRP was >8.2 µg/mL (ID in the presence of inflammation) [9, 14]. Secondary definitions of ID included the following: ferritin <12 ng/mL [15], ferritin <30 ng/mL [16], ferritin <12 ng/mL and CRP <6 µg/mL or ferritin ≥50 ng/mL and CRP ≥6 µg/mL [17], and sTfR (µg/mL)/log10 ferritin (ng/mL) > 1.5 [18] or 5.6 [19].

Statistical Analysis
Association between parasite density and concurrent body iron status was evaluated using generalized estimating equation (GEE)-based linear regression models to account for the correlation between measurements at multiple time points from the same child. Likewise, associations between prevalence data (ie, parasitemia, hyperparasitemia, or severe malaria) and concurrent iron status were estimated with GEE-based logistic regression models. Cross-sectional analyses considered outcomes concurrent with iron status measures. Longitudinal analyses examined the relationship between iron status measured at aparasitemic routine visits and subsequent risk of a malarial event. Odds ratios (ORs) and 95% confidence intervals (CIs) were calculated as estimates of risk. Cox proportional hazards analysis was used to evaluate the impact of iron status on...
mortality. In Cox models, iron status was coded as an unlagged time-varying covariate [20–22]. All GEE models were adjusted for Hemoglobin S, village, bed net use, low birth weight, and hemoglobin levels. In Cox models, these covariates were assessed and retained if $P < .1$. Age was considered as both a confounding covariate and as an effect modifier (interaction term) in our models.

**RESULTS**

**Baseline Characteristics of the Study Population**

A total of 785 children were included in this study with an average of 113 weeks of follow-up. Table 1 describes the baseline characteristics of the study population. Malaria parasitemia was frequent and intense: 16% (5082 of 32641) of routine blood smears were positive with a mean parasite density of 829 parasites per 200 WBCs.

**Prevalence of ID in the Study Area**

Because parasitemia can increase ferritin levels (resulting in underestimation of the prevalence of ID), we examined the prevalence of ID in samples obtained from aparasitemic children during routine visits (n = 1696). The prevalence of ID was low in neonates but increased rapidly during infancy (Figure 1). At routine, aparasitemic visits, children with ID had 6.6-fold lower odds of concurrent malaria parasitemia compared with children who were iron-replete after adjusting for potential confounders (OR [95% CI] = 0.15 [0.12, 0.19], $P < .001$; Figure 2A). This effect was significantly modified by age ($P < .001$).

At routine, parasitemic visits (n = 649), children with ID had 3.9-fold lower parasite density compared with children who were iron-replete after adjusting for potential confounders (adjusted mean parasite density [standard error] 390 per 200 WBC [59] versus 1526 per 200 WBC [127], $P < .001$; Figure 2B). This effect was also significantly modified by age ($P < .001$).

**ID Is Associated With Decreased Prevalence and Intensity of Parasitemia**

At routine visits (n = 2345), children with ID had 6.6-fold lower odds of concurrent malaria parasitemia compared with children who were iron-replete after adjusting for potential confounders (OR [95% CI] = 0.15 [0.12, 0.19], $P < .001$; Figure 2A). This effect was significantly modified by age ($P < .001$).

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**ID Predicts Resistance to Malaria Infection and Disease**

In cross-sectional analyses, inflammation accompanying malaria can lead to misclassification of children with ID as iron-replete (because ferritin is an acute phase reactant), and thus the conclusion that ID protects from concurrent malaria. To avoid this design concern, we prospectively examined the relationship between ID at each iron status determination and...
risk of future malaria outcomes. We adjusted for potential confounders, the number of iron status measures per child, the average age of the child at their iron status measurements, and the follow-up time interval after each iron status determination (ie, the time interval until next iron status determination or until completion of study). In GEE-based logistic regression models, children who were ID at routine aparasitemic visits had a 1.3-fold decreased odds of parasitemia (OR [95% CI] = 0.77 [0.66, 0.89], \( P < .001 \)) and a 1.6-fold decreased odds of severe malaria (OR [95% CI] = 0.62 (0.39, 0.98), \( P = .04 \)) during subsequent months compared with children with normal iron stores, whereas the risk of hyperparasitemia was not different in these groups (\( P = \) not significant [NS]). Concordant results were obtained when the proportion of future visits with these outcomes was analyzed in GEE-based linear regression models (Figure 4).

**DISCUSSION**

ID-associated morbidities support international guidelines for iron supplementation in children under 2 years of age in areas with a high prevalence of anemia [1, 2]. Unfortunately, iron plus folate supplementation given during a recent randomized, placebo-controlled trial involving \( N = 42,546 \)
children living in a high *P. falciparum* transmission area resulted in a 15% increase in all-cause mortality [3], and sub-
study analyses suggested that enhanced malarial morbidity was concentrated in children who were iron-replete at baseline. Although malaria morbidity and mortality were not increased in some other iron supplementation trials [6–8], this result has raised concern over routine iron supplementation programs in malarious areas.

Because iron supplementation may exacerbate malaria outcomes, we examined whether naturally occurring ID confers protection in children residing in a high transmission area of Tanzania. Specifically, we examined the influence of iron status on malaria risk in both cross-sectional and longitudinal analyses in a birth cohort of children who were receiving intensive, active health surveillance and prompt antimalarial treatment. We found that ID was highly prevalent and, despite intensive blood smear monitoring and prompt antimalarial treatment, children who were iron-replete had significantly more malaria infection, morbidity, and mortality.

We identified a high prevalence of ID in our birth cohort which far exceeds that reported from healthy European children [23] but is consistent with results from another study in rural Tanzania [24]. The proportion of mothers with ID at delivery in our cohort was 78.4% [9], which mirrors the ID frequency in their offspring after 1 year (Figure 1). This concordance may reflect a shared iron-poor diet, or inherited, potentially beneficial, mutations at any of several iron transport, storage, regulatory, or heme synthesis genes [25].

Children living in malaria-endemic areas acquire antidisease and antiparasite immunity as they age, and this process is accompanied by the acquisition of antiparasite antibodies [26]. We demonstrate for the first time that ID also provides strong protection against parasitemia and malaria-associated morbidity and mortality during childhood. Because ID modifies malaria risk, iron status should be assessed in studies that evaluate the impact of other factors, such as parasite exposure, genetic polymorphisms, or therapeutic interventions, on malaria outcomes.

Despite many iron supplementation trials in malarious areas, remarkably few studies have examined the relationship between iron status and malaria in unsupplemented populations. In a longitudinal study of children (n = 240) between 8 months and 8 years of age who were living in a low transmission area on the coast of Kenya, those with ID had a 30% lower incidence of mild clinical malaria (any parasitemia accompanied by axillary temperature >37.5°C) during 12 months of follow-up compared with children who were iron-replete [10]. In our study from Tanzania, ID reduced parasitemia (23%) and severe malaria (38%) to a roughly similar degree that it reduced mild clinical malaria risk in Kenya.

Several trials have demonstrated that iron supplementation exacerbates malaria. Iron supplementation increased the risk of malaria among children from Papua New Guinea [4] and the risk of fever associated with malarial parasitemia in Gambian children [27]. Recently, a community-based, randomized, controlled study in Zanzibar found that children who received iron plus folic acid supplementation had significantly increased mortality or hospital admission and had a trend toward increased mortality compared with children who received placebos [3]. In contrast, a recent meta-analysis of
iron supplementation trials in malarious areas did not find increased mortality in individuals who were given iron; however, the average length of follow-up in the pooled studies was less than 4 months, the analysis included children up to the age of 18 years, an age at which malaria-associated mortality is limited, and baseline iron status was rarely assessed [28].

The assessment of iron status in the context of infection and inflammation is daunting, and no ideal methods are available [9, 15–19, 29]. We defined iron status primarily based on serum ferritin levels [30]. Because serum ferritin is an acute phase protein and thus may increase during inflammation, some inflamed individuals with ID will be misclassified as iron-replete. We addressed this potential misclassification in several ways. First, we measured CRP concurrently as a marker of inflammation, and we adjusted the threshold ferritin level used to define ID cutoffs based on CRP levels [9, 14, 31]. We recognize that this approach remains susceptible to residual misclassification if the CRP elevation resolves before the ferritin response [32]. Second, we restricted our cross-sectional analyses of concurrent iron status and parasitemia to samples obtained during routine, nonsick visits (Figure 2). Third, we repeated our cross-sectional analyses of the impact of concurrent iron status on parasitemia, hyperparasitemia, and severe malaria (Figure 3) by using several secondary definitions of ID (see Supplementary Table 1). These alternative definitions of ID are reportedly less influenced by concurrent inflammation [15–19, 31], and all demonstrated similar relationships with malaria risk. Fourth, we examined the relationship between ID and future malaria risk, and we restricted our analyses to children who provided samples during routine, aparasitemic visits (Figures 4 and 5). This restriction significantly attenuates the possibility of misclassification of inflamed individuals with ID as iron-replete, and the longitudinal nature of the analysis allows directional inference.

Strikingly, ID measured at aparasitemic routine visits was a strong predictor of decreased mortality (Figure 5). During periods of ID, children had 60% reduced all-cause mortality and 66% reduced malaria-associated mortality over the 3 years of follow-up compared with iron-replete periods. These results may underestimate the decreased mortality attributable to ID because other macronutrient and micronutrient deficiencies that are frequently coincident with ID are causally linked to increased morbidity and mortality [33].

The mechanisms by which ID might limit parasite density are diverse and could involve both parasite and host-specific effects. Malaria parasites acquire iron in a transferrin-independent pathway [34], and chelation of intraerythrocytic iron reduces parasite growth [35]. In the host, chelation of iron increases cellular nitric oxide (NO) production and parasite killing in coculture systems [36] and similarly increases NO production in children being treated for cerebral malaria [37]. Iron inhibits the expression of inducible NO synthase (iNOS), and NO is a principal component of macrophage-mediated cytotoxicity towards *P. falciparum* [36]. Therefore, ID may amplify iNOS-mediated defenses against this pathogen, and iron chelation...
therapy has been associated with improved clinical course of cerebral malaria in some [38] but not all studies [39]. In addition, iNOS-mediated defenses control liver stage parasite development [40] and could contribute to the decreased frequency of parasitemia that we observed. Alternatively, ID may reflect concurrent hookworm infection; however, the relationship between hookworm infection and malaria outcomes remains unclear.

ID may pose a direct toxic insult to malarial parasites. Parasites detoxify heme by polymerizing it into crystals of hemozoin in a process that requires Fe$^{3+}$/carboxylate bonds [41]. During ID erythropoiesis, ferrochelatase inserts zinc rather than iron into protoporphyrin IX [42]. Zinc protoporphyrin (ZPP) binds hemozoin crystals and inhibits crystal elongation, resulting in heme toxicity and possibly reduced parasite growth [43]. Measures of ZPP and the iron regulatory peptide hepcidin may identify the role of iron availability and iron sequestration in mediating protection from malaria in future studies.

Our results in a large birth cohort among children living in an area with intense *P. falciparum* transmission indicate that ID confers significant protection from malaria morbidity and mortality despite active, intense health surveillance. These findings are consistent with experimental evidence that links ID to reduced parasite fitness and enhanced host resistance. Our results suggest that the relationship between iron status and malaria is a continuum of risk that increases as individuals progress from ID to normal iron stores. For this reason, efforts to focus iron supplementation among individuals with ID may nevertheless increase the risk of malaria morbidity and mortality. These data warrant additional interventional studies to ascertain the benefits and risks of iron supplementation for children living in malaria-endemic regions representing a broad range of transmission pressures, but only if coupled with effective malaria control measures.

**Supplementary Data**

Supplementary materials are available at *Clinical Infectious Diseases* online (http://cid.oxfordjournals.org). Supplementary materials consist of data provided by the author that are published to benefit the reader. The posted materials are not copyrighted. The contents of all supplementary data are the sole responsibility of the authors. Questions or messages regarding errors should be addressed to the author.

**Notes**

**Acknowledgments.** We thank the Mother–Offspring Malaria Studies Project staff for their efforts in collecting clinical data, sample processing, and interpreting malaria blood smears. We also thank Sunthorn Pond-Tor for assistance with iron status measures.

**Financial support.** This work was supported by grants from the US National Institutes of Health (grant AI52059) and the Bill & Melinda Gates Foundation (grant 1364; to P. E. D.). S. H. received grant support from Seattle BioMed.

**Potential conflicts of interest.** All authors: No reported conflicts. All authors have submitted the ICMJE Form for Disclosure of Potential Conflicts of Interest. Conflicts that the editors consider relevant to the content of the manuscript have been disclosed.

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