The content of African diets is adequate to achieve optimal efficacy with fixed-dose artemether-lumefantrine: a review of the evidence

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Abstract

A fixed-dose combination of artemether-lumefantrine (AL, Coartem®) has shown high efficacy, good tolerability and cost-effectiveness in adults and children with uncomplicated malaria caused by Plasmodium falciparum. Lumefantrine bioavailability is enhanced by food, particularly fat.

As the fat content of sub-Saharan African meals is approximately a third that of Western countries, it raises the question of whether fat consumption by African patients is sufficient for good efficacy. Data from healthy volunteers have indicated that drinking 36 mL soya milk (containing only 1.2 g of fat) results in 90% of the lumefantrine absorption obtained with 500 mL milk (16 g fat). African diets are typically based on a carbohydrate staple (starchy root vegetables, fruit [plantain] or cereals) supplemented by soups, relishes and sauces derived from vegetables, pulses, nuts or fish. The most important sources of dietary fat in African countries are oil crops (e.g. peanuts, soya beans) and cooking oils as red palm, peanut, coconut and sesame oils. Total fat intake in the majority of sub-Saharan countries is estimated to be in the range 30–60 g/person/day across the whole population (average 43 g/person/day). Breast-feeding of infants up to two years of age is standard, with one study estimating a fat intake of 15–30 g fat/day from breast milk up to the age of 18 months. Weaning foods typically contain low levels of fat, and the transition from breast milk to complete weaning is associated with a marked reduction in dietary fat. Nevertheless, fat intake >10 g/day has been reported in young children post-weaning. A randomized trial in Uganda reported no difference in the efficacy of AL between patients receiving supervised meals with a fixed fat content (~23 g fat) or taking AL unsupervised, suggesting that fat intake at home was sufficient for optimal efficacy. Moreover, randomized trials in African children aged 5–59 months have shown similar high cure rates to those observed in older populations, indicating that food consumption is adequate post-weaning. In conclusion, it appears that only a very small amount of dietary fat is necessary to ensure optimal efficacy with AL and that the fat content of standard meals or breast milk in sub-Saharan Africa is adequate.
Background
Progress in the control and treatment of malaria in Africa has been threatened by growing resistance to previously effective therapies, notably chloroquine and sulphadoxine/pyrimethamine. Combination therapy is now widely regarded as the most effective strategy for limiting the emergence and spread of resistance and combination treatment based on artemisinin is now recommended by the World Health Organization (WHO) [1].

Artemether-lumefantrine (AL, Coartem®, Novartis Pharma AG, Basel, Switzerland) combines the artemisinin-derivative artemether, which has a prompt onset of action and provides fast symptomatic relief, with lumefantrine, which has a slower onset of action and clears residual parasites [2]. AL has consistently shown high efficacy and tolerability in African adults and children with uncomplicated malaria caused by Plasmodium falciparum, achieving 28-day polymerase chain reaction (PCR)-corrected parasitological cure rates of >95% [3-7]. AL is the first and only fixed-dose artemisinin-based combination treatment (ACT) pre-qualified by the WHO, and has now been adopted as first-line treatment for uncomplicated malaria due to P. falciparum in more than 20 African countries including Kenya, Malawi, Uganda, Tanzania, Mozambique, Nigeria, South Africa and Ethiopia. Many other countries have also switched to AL as first-line therapy. The manufacturer introduced price reductions in 2006 and 2008, cutting the public sector price by approximately 50% since 2001.

Cost-effectiveness assessments, one based on data from a randomized trial in Tanzania [8] and another based on public health data collected in the field in Zambia [9], have confirmed that AL offers a cost-effective treatment option compared to other antimalarial drugs.

Artemether and lumefantrine are lipophilic molecules. Food, especially dietary fat, enhances the bioavailability of both agents, but the effect is more pronounced with lumefantrine [2]. The fat content of African meals tends to be relatively low; indeed, total fat consumption in sub-Saharan Africa is approximately a third that of Western countries [10]. This raises the important question of whether adults or children prescribed AL in sub-Saharan Africa consume sufficient fat to ensure parasite clearance. This article reviews the typical components and fat content of African diets and consider the adequacy of fat intake in terms of achieving optimal efficacy.

Characteristics of lumefantrine absorption
Lumefantrine, like several other lipophilic antimalarial agents, is absorbed slowly after oral administration, and bioavailability varies between individuals [2,11-13]. Concomitant food increases lumefantrine bioavailability [14]. In a double-blind trial of 260 patients with uncomplicated malaria in Thailand, Ezzet et al showed that both the extent and variability of lumefantrine absorption improved with clinical recovery as normal food intake was resumed [14]. Oral bioavailability increased by 108% when a normal meal was eaten close to AL dosing, compared to the fasting condition. Adequate lumefantrine absorption is important in order to avoid low plasma concentrations with risk of treatment failure [2].

The key question for the healthcare team is how much dietary fat is necessary to achieve complete parasite clearance with AL. This question has been investigated by Ashley et al in a cross-over pharmacokinetic study in which lumefantrine exposure (area under the curve, AUC) was measured in healthy volunteers after a single dose of AL was administered with different volumes of soya milk or with no milk [15]. Based on these data, the authors constructed a population model which indicated that in this population of healthy volunteers, drinking 36 mL of soya milk (containing only 1.2 g of fat) resulted in 90% of the lumefantrine exposure obtained with 500 mL milk (16 g fat); 74 mL of soya milk (2.4 g of fat) achieved 99% of the exposure seen with 500 mL (Figure 1). Ashley and colleagues pointed out that 1.2 g of fat can be obtained in a remarkably small amount of food: for example, a single teaspoon of vegetable oil contains 5 g of fat.

Typical components of African diets
Diets in Africa vary widely according to regional, economic, environmental and cultural factors. Nevertheless, there are certain typical patterns of food consumption (Table 1). African diets are traditionally based on a carbohydrate staple that is supplemented by soups, relishes and sauces derived from a wide range of other foodstuffs. The carbohydrate component may be primarily cereals (e.g. maize, sorghum, millet, rice), starchy roots (e.g. potatoes, sweet potatoes, yams, cassava) or fruit (plantain), depending on the country and local area. In recent years there has been a trend away from the consumption of roots towards maize, rice and wheat, particularly in cities [16]. Accompaniments to the carbohydrate staple are usually composed of vegetables (e.g. eggplant, cabbage, carrots, spinach, kales, French beans, onions), pulses, nuts (particularly peanuts), meat (chicken, beef or pork) or fish, depending on local availability, household income, season and community habits, with the greatest variety seen in coastal regions and the fertile highlands [16].

The Food and Agriculture Organization (FAO) of the United Nations compiles data on various aspects of food production and availability [17]. These statistics offer a valuable source of information, although it should be borne in mind that estimates are based on the amount of food available for human consumption, rather than
actual amounts eaten, and do not account for wastage. Table 2 summarizes the consumption of selected food groups in sub-Saharan countries based on FAO data, and highlight the predominance of cereals and starchy root vegetables [18].

The most important sources of dietary fat in African countries are oil crops (e.g. peanuts, soy beans) and oils obtained from vegetables or plants such as red palm oil, groundnut oil, coconut oil and sesame oil. These are widely used in food preparation, for frying or as additions to sauces and stews. Additional contributions to fat intake are made by other commonly-used foodstuffs such as nuts and pulses. Table 3 describes the typical fat content of some staple foodstuffs [19]. While food composition inevitably varies according to crop variety, growing conditions and freshness, these illustrate that relatively small amounts of certain widely-consumed items such as soya beans and ground nuts contain high levels of fat. A further contribution comes from whole-grain cereals if the cereal germ is not separated out prior to milling: for example, maize contains 4.8 g/100 g [19] and is a staple food in many countries.

**Total fat intake in African countries**

Total energy intake per person across sub-Saharan Africa is currently estimated to be 2220 kcal/day, compared to 3340 kcal/day in developed countries [10]. Regional differences exist, most notably a higher average energy intake in West Africa compared to other areas (Table 4). Estimates of total fat intake from the FAO for the period 2001–2003 suggest that the average total amount of fat consumed per person per day vary considerably between sub-Saharan African countries (Table 5) [10]. The majority of countries, however, fall into the range 30–60 g/person/day. The daily intake of fat across sub-Saharan Africa is estimated to be 43 g/person/day, with the highest fat intake in West Africa (Table 1); for comparison, people in developed countries consume 123 g/person/day, on average [10]. The per capita daily intake of fat has generally remained stable or shown an upward trend in the majority of African countries over the last three decades [10], with exceptions being due to major conflicts or famines.

**Fat content of breast milk and weaning foods**

It is standard to breast-feed infants up to 4–6 months in all regions of Africa, although supplementary cow’s milk may be given. The fat content of breast milk in rural African areas varies with the season, stage of lactation and between individual mothers, but an investigation in a farming community in The Gambia recorded an average fat content of 40 g/L breast milk [20], similar to that reported in other parts of the world [21-23]. Breast-fed infants in The Gambia have been shown to receive 15–30 g fat/day from breast milk up to the age of 18 months, with breast milk continuing to make a major contribution

| Region      | Adults                                                                 | Infants                                                                 |
|-------------|------------------------------------------------------------------------|------------------------------------------------------------------------|
| East Africa | Starchy foods (e.g. cooked bananas, potatoes, cassava, corn, wheat products, rice). Food consumed to accompany starchy includes beef, lamb, poultry, stews made of legumes, soups, vegetables (mostly fried in cooking oil or fat or cream) and peanut/groundnut-based sauces | Exclusively 4–6 months unless contraindicated. Weaning may start from month 3 onwards Supplementary milk from cows Early food: porridge containing milk or coconut oil. Exclusively breast-fed up to 4 to 6 months unless contraindicated Supplementary milk (cow milk or baby formula) Early foods include chocolate drinks, mashed yam, beans Exclusively breast-fed up to 6 months Supplementary milk from cows Early food: porridge from maize meal and milk Exclusively breast-fed up to 6 months unless contraindicated Supplementary milk from cows Early food: porridge maize meal |
| West Africa | Bean balls fried with oil, maize fortified with palm oil, groundnuts, bread, eggs, rice, yam, beef or fish stew, yams, vegetables (mostly fried in cooking oil) | Supplementary milk from cows Early food: porridge containing milk or coconut oil. Exclusively breast-fed up to 4 to 6 months unless contraindicated Supplementary milk (cow milk or baby formula) Early foods include chocolate drinks, mashed yam, beans Supplementary milk from cows Early food: porridge maize meal |
| Central Africa | Rice, cassava, sorghum, millet, vegetables (fried in peanut oil), palm oil, fish, meat | Early food: porridge maize meal |
| Southern Africa | Maize, mashed potatoes, vegetables, beef, rice, bread, cassava | Early food: porridge maize meal |
to fat intake even after partial weaning [24]. Comparing infants in The Gambia and the United Kingdom, Prentice et al found that fat intake from breast milk was at least as high in the Gambian children while being breast-fed exclusively [24].

Infants usually continue to receive breast milk until two years of age, with some food introduced from three months or later. The first weaning food is usually porridge made from a cereal such as rice, corn or millet mixed with water, or possibly cow's milk, with salt and sugar. Such porridges have a relatively low fat content (e.g. 0.1–0.3 g/100 g); although addition of soya milk or peanuts in some regions increases the fat content (1.2–1.9 g/100 g) [25]. In West Africa there tends to be a greater diversity of early

| Table 2: Average consumption of selected food groups in sub-Saharan African countries during 2001–2003 [18] |
|---------------------------------------------------------------|
| **Average intake (g/person/day)** |
| Cereals | Fish, seafood | Fruit | Meat | Milk | Oilcrops* | Starchy roots | Sugars, sweeteners | Vegetable oil | Vegetables |
|----------|---------------|-------|------|------|----------|--------------|----------------|---------------|------------|
| Burkina Faso | 601 | 6 | 14 | 30 | 56 | 42 | 14 | 16 | 15 | 13 | 47 |
| Burundi | 84 | 5 | 284 | 10 | 13 | 2 | 103 | 632 | 8 | 2 | 93 |
| Cameroon | 288 | 38 | 213 | 41 | 40 | 19 | 39 | 400 | 28 | 21 | 217 |
| Congo, Dem Republic of | 102 | 16 | 82 | 13 | 4 | 12 | 10 | 842 | 8 | 15 | 25 |
| Côte d'Ivoire | 331 | 40 | 201 | 31 | 21 | 20 | 1 | 619 | 30 | 37 | 103 |
| Ethiopia | 384 | 1 | 28 | 23 | 61 | 2 | 35 | 168 | 12 | 4 | 31 |
| Ghana | 249 | 80 | 319 | 27 | 21 | 35 | 2 | 1107 | 19 | 16 | 88 |
| Kenya | 337 | 12 | 154 | 40 | 263 | 4 | 42 | 164 | 55 | 20 | 102 |
| Malawi | 403 | 10 | 107 | 14 | 12 | 13 | 34 | 504 | 31 | 9 | 54 |
| Mali | 502 | 21 | 8 | 51 | 126 | 5 | 25 | 30 | 30 | 18 | 66 |
| Mozambique | 289 | 6 | 48 | 15 | 12 | 5 | 26 | 665 | 31 | 19 | 17 |
| Nigeria | 399 | 20 | 185 | 23 | 20 | 21 | 26 | 593 | 31 | 38 | 167 |
| Sudan | 370 | 5 | 83 | 59 | 410 | 11 | 22 | 13 | 55 | 21 | 135 |
| Uganda | 170 | 20 | 556 | 31 | 67 | 35 | 65 | 561 | 24 | 5 | 55 |
| United Republic of Tanzania | 307 | 19 | 81 | 27 | 70 | 13 | 28 | 518 | 21 | 14 | 76 |
| Zimbabwe | 355 | 4 | 32 | 42 | 54 | 17 | 13 | 43 | 97 | 26 | 30 |

* Excluding products

| Table 3: Fat content of typical foodstuffs [19] |
|-----------------------------------------------|
| **Type** | **Food** | **Fat content (g/100 g)** |
|-----------------|----------|---------------------------|
| Carbohydrates and legumes | Maize | 4.8 |
| | Millet | 3.5 |
| | Rice | 0.5 |
| | Sorghum | 3.2 |
| | Wheat flour | 2.0 |
| | Cassava meal | 0.5 |
| | Beans | 1.5 |
| | Lentil | 1.2 |
| | Pigeon peas | 1.3 |
| | Soya beans | 18.0 |
| | Yam | 0.1 |
| Meat/fish | Beef | 18.0 |
| | Chicken | 6.5 |
| | Eggs | 10.0 |
| | Fish | 7.4 |
| | Peanuts | 45.0 |
| | Coconut | 35.0 |
| | Sunflower | 46.0 |
| | Avocado | 18.0 |
foodstuffs, including chocolate, mashed potatoes or beans (Table 1).

The transition from breast milk to weaning foods and then to adult foods is associated with a marked reduction in the proportion of energy that is obtained from fat. A study in Kenya was undertaken in infants during the second year of life (mean 14–20 months) to assess fat intake over the period of weaning [26]. In these young children, the mean quantity of breast milk ingested was calculated to be 502 ± 239 g per 24 hours, although this varied widely between individuals [26]. The authors calculated the average fat intake from milk to be approximately 17 g/day, which was supplemented by 10.5 g/day from weaning foods. In the period prior to weaning, fat intake from breast milk decreased to approximately 11 g/day but after complete weaning the increased food consumption provided only an additional 3.5 g/day. A further detailed study in The Gambia has shown that total fat intake remained relatively stable over the first 17 months of life, but as the percentage of other foods (notably cereals and peanuts) increased in the diet, the percentage of energy from fat decreased from 50% during breast-feeding to 30% during mixed feeding and to 15% after total weaning [24]. Nevertheless, total fat intake, which was ~25 g/day until complete weaning, was >10 g/day subsequently.

Cow’s milk is only given to infants infrequently in many African countries because of the low milk yield per animal and scarcity of preservation equipment, although it is more common in some countries (e.g. Kenya, Somalia and Sudan). Where cow’s milk is used, however, it is a good source of fat with a fat content of 30 g/L [19], only slightly lower than breast milk.

Relevant experience from clinical trials
In most trials of AL in African populations, AL has been administered under supervision with food or drink given in the hospital or clinic [4,7,27-29]. Two studies, however, have assessed ‘real-life effectiveness’ of AL, in which the drug was taken unsupervised at home [6,30]. These are helpful in providing an indication of whether normal, unsupervised food consumption during AL administration ensures good efficacy. Piola et al undertook a randomized trial in 957 patients in Uganda to compare the effectiveness of AL when administered under supervision in hospital with fat intake ensured by a meal comprising ~300 mL milk (10 g fat) with 30 g peanuts (13 g fat) versus AL taken unsupervised at home [6]. In the unsupervised group, only the first dose was taken in the clinic, and patients were then discharged and advised to take subsequent doses with a meal that contained fat (with no fat content specified), or breast milk in breastfed infants. The

| Region                      | Dietary energy consumption (kCal/person/day) | Dietary fat consumption (kCal/person/day) |
|-----------------------------|---------------------------------------------|------------------------------------------|
| Sub-Saharan Africa          | 2220                                        | 43                                       |
| Central Africa              | 1830                                        | 36                                       |
| East Africa                 | 2040                                        | 35                                       |
| Southern Africa             | 2080                                        | 36                                       |
| West Africa                 | 2580                                        | 57                                       |

Table 4: Average dietary energy and fat consumption in regions of Africa. [10]

| Total fat consumption (g/person/day) |
|--------------------------------------|
| Burkina Faso                        | 56                                        |
| Burundi                              | 10                                        |
| Cameroon                             | 46                                        |
| Congo, Democratic Republic of        | 26                                        |
| Côte d’Ivoire                       | 59                                        |
| Ethiopia                             | 20                                        |
| Ghana                                | 38                                        |
| Kenya                                | 49                                        |
| Malawi                               | 33                                        |
| Mali                                 | 46                                        |
| Mozambique                           | 33                                        |
| Nigeria                              | 63                                        |
| Sudan                                | 69                                        |
| Uganda                               | 32                                        |
| United Republic of Tanzania          | 31                                        |
| Zimbabwe                             | 55                                        |

Table 5: Average dietary fat consumption in sub-Saharan African countries during 2001–2003 [10]
28-day PCR-corrected cure rate in the evaluable population was 100% in both cohorts (95% CI 98–100 in the supervised group and 99–100 in the unsupervised group). The type of meal eaten at home was not recorded, so it cannot be determined if patients ate especially fatty meals or standard food. Lumezantrine plasma concentration was measured on days 3 and 7 in a random selection of 433 patients taking part in the study [31], and multivariate modelling of these data confirmed that supervised treatment was significantly associated with higher lumezantrine concentrations (p < 0.0001) [31]. In another trial, undertaken in an area of Tanzania with a high level of resistance to sulfadoxine-pyrimethamine and amodiaquine, Mutabingwa and colleagues randomized 1292 children aged 4–59 months with uncomplicated malaria to AL or to conventional drugs [30]. All agents were taken at home, without supervision, and no specific advice was given regarding concomitant food consumption. At 28 days, 97.3% of patients receiving AL showed parasitological cure after adjustment for recrudescence.

Together, these findings suggest that food consumption during unsupervised AL dosing in African patients, including children, provides sufficient fat to ensure effective parasitological clearance.

As discussed above, the fat consumption of infants shortly after weaning can be relatively low. It is highly relevant, therefore, that randomized trials in African children aged from five months [32,33] or 12 months [34] up to 59 months have consistently shown 28-day PCR-corrected parasitological cure rates of greater than 95% i.e. similar to cure rates observed in older populations.

It should be noted that data are scarce concerning lumezantrine absorption, and the effect of food consumption, in patients with HIV/AIDS or other coinfections such as tuberculosis. Pharmacokinetic and pharmacodynamic studies are required in the HIV/AIDS population, particularly in the light of increased antimalarial failure rates in such patients due to increased parasite burden and reduced host immunity associated with HIV infection (1).

Conclusion

Although lumezantrine bioavailability is enhanced by fat, pharmacokinetic evidence suggests that only a very small amount of dietary fat is required [15]. In the event of varying dietary fat intake, such as during unsupervised dosing where fat consumption was not controlled [6] or was not specifically advised [30], the efficacy of AL does not appear to have been impaired. This is consistent with published statistics concerning the dietary fat intake of African populations, which indicate that average fat consumption is generally in the range of 30–60 g/day in sub-Saharan African countries. Even in young children shortly after weaning, in whom fat intake tends to be low, it has been estimated that fat intake exceeds 10 g/day. In summary, there is good evidence to suggest that the content of standard African diets or breast milk is adequate to ensure excellent efficacy for AL.

Abbreviations

ACT: artemisinin-based combination treatment; AL: artemether-lumezantrine; AUC: area under the curve; FAO: Food and Agriculture Organization; PCR: polymerase chain reaction.

Competing interests

The authors declare that they have no competing interests other than COF, who received a research grant from Novartis in 2002, and NM and ON, who are employees of Novartis Pharma AG.

Authors’ contributions

All authors provided critical review of the text and approved the final version.

References

1. World Health Organization: Guidelines for treatment of malaria Geneva: World Health Organization; 2006.
2. White NJ, van Vugt M, Ezet F: Clinical pharmacokinetics and pharmacodynamics and pharmacodynamics of artemether-lumezantrine. Clin Pharmacokinet 1999, 37:105-125.
3. Makanga M, Premji Z, Falade C, Karbwang J, Mueller EA, Andriano K, Hunt P. De Palacios P: Efficacy and safety of the six-dose regimen of artemether-lumezantrine in pediatrics with uncomplicated Plasmodium falciparum malaria: a pooled analysis of individual patient data. Am J Trop Med Hyg 2006, 74:991-998.
4. Falade C, Makanga M, Premji Z, Omgmann CE, Stockmeyer M, de Palacios P: Efficacy and safety of artemether-lumezantrine (Coartem®) tablets (six-dose regimen) in African infants and children with acute, uncomplicated falciparum malaria. Trans R Soc Trop Med Hyg 2005, 99:459-467.
5. Dorsey G, Stadke S, Clark TD, Njama-Meya D, Nzuzubara B, Maiteki-Sebuguzi C, Dokomajilar C, Kamya MR, Rosenhall P: Combination therapy for uncomplicated falciparum malaria in Ugandan children: a randomized trial. JAMA 2007, 297:2210-2219.
6. Piola P, Fog C, Bajunirwe F, Biraro S, Grandesso F, Ruzagira E, Babin-gumira J, Kigozi I, Kuguli J, Kyomuhendo J, Ferradini L, Taylor W, Checchi F, Guthmann P: Supervised versus unsupervised intake of six-dose artemether-lumezantrine for treatment of acute, uncomplicated Plasmodium falciparum malaria in Mbarara, Uganda: a randomized trial. Lancet 2005, 365:1467-1473.
7. Abdullah S, Sagara I, Borrmann S, D’Alessandro U, Gonzalez R, Hamel M, Ogutu B, Märtensson A, Lyimo J, Miga H, Sasi P, Nahum A, Bassat Q, Juma E, Otieno L, Bjorkman A, Beck HP, Andriano K, Cousin M, Leefèvre G, Ubben D, Premji Z: Efficacy and safety of artemether-lumezantrine dispersible tablet in African infants and children with uncomplicated malaria: a randomised, investigator-blinded, multi-centre comparison with the crushed commercial tablet. Lancet 2008 in press.
8. Wiseman V, Kim M, Mutabingwa TK, Whitty CJ: Cost-effectiveness study of three antimalarial drug combinations in Tanzania. PLoS Med 2006, 3:e373.
9. Chanda P, Maijye F, Chitam BM, Spilayanambe N, Hawela N, Banda P, Okorosobo T: A cost-effectiveness analysis of artemether-lumezantrine for treatment of uncomplicated malaria in Zambia. Malar J 2007, 6:21.
10. Food and Agriculture Organization of the United Nations Statistics Division [http://www.fao.org/docrep/foodsecurity/files/ FoodConsumptionNutrients_en.xls]. Accessed April 2008.
11. Lefèvre G, Thomsen MS: Clinical pharmacokinetics of artemether and lumefantrine (Riamet®). Clin Drug Invest 1999, 18:467-480.
12. Ezet F, van Vugt M, Nosten F, Looareesuwan S, White NJ: Pharmacokinetics and pharmacodynamics of lumefantrine (benflumetol) in acute falciparum malaria. Antimicrob Agents Chemother 2000, 44:697-704.
13. Schmidt LK: Food-drug interactions. Drugs 2002, 62:1481-1502.
14. Ezet F, Mull R, Karbwang J: Population pharmacokinetics and therapeutic response of CGP 56697 (artemether + benflumetol) in malaria patients. Br J Clin Pharmacol 1998, 46:553-561.
15. Ashley EA, Stepniewska K, Lindegårdh N, Annerberg A, Kham A, Brockman A, Singhavison P, White NJ, Nosten F: How much fat is necessary to optimize lumefantrine oral bioavailability? Trop Med Int Health 2007, 12:195-200.
16. Food and Agriculture Organization Corporate Document Repository. Agriculture, Food and Nutrition for Africa: A Resource Book for Teachers of Africa. Chapter 7: Food, nutrients and diets [http://www.fao.org/docrep/W0078E/w0078e08.htm#P7404_499006]. Accessed April 2008
17. Food and Agriculture Organization of the United Nations [http://www.fao.org/]. Accessed April 2008
18. Food and Agriculture Organization of the United Nations, Food Consumption Food Groups [http://www.fao.org/hoststat/foodsecurity/Files/FoodConsumptionFoodGroups_en.xls]. Accessed April 2008
19. Food and Agriculture Organization of the United Nations/US Department of Health, Education and Welfare Food Composition Table for Use in Africa [http://www.fao.org/docrep/003/X6877E/X6877E00.htm]. Accessed April 2008
20. Prentice A, Prentice AM, Whitehead RG: Breast-milk fat concentrations of rural African women. 2. Long-term variations within a community. Br J Nutr 1981, 45:495-503.
21. Kent JC, Miotoulas LR, Cregan MD, Ramsay DT, Doherty DA, Hartmann PE: Volume and frequency of breastfeedings and fat content of breast milk throughout the day. Pediatrics 2006, 117:e387-395.
22. Larnkjaer A, Schack-Nielsen L, Michaelsen KF: Fat content in human milk according to duration of lactation. Pediatrics 2006, 117:988-989.
23. Liyanage C, Hettiarachchi M, Mangalalaweewa P, Malawipathirana S: Adequacy of vitamin A and fat in the breast milk of lactating women in south Sri Lanka. Public Health Nutr 2008 in press.
24. Prentice AM, Paul AA: Fat and energy needs of children in developing countries. Am J Clin Nutr 2000, 72(Suppl):1253S-1265S.
25. Hudson GJ, John PMV, Paul AA: Variation in the composition of Gambian foods; the importance of water in relation to energy and protein content. Ecol Food Nutr 1980, 10:9-17.
26. Onyango AW, Rezevezre O, Esrey SA: The contribution of breast milk to toddler diets in western Kenya. Bull World Health Organ 2002, 80:292-299.
27. von Seidlein L, Jaffar S, Pinder M, Haywood M, Snounou G, Gemetery B, Gathmann I, Royce C, Greenwood B: Treatment of African children with uncomplicated falciparum. J Infect Dis 1997, 176:1113-1116.
28. von Seidlein L, Bojang K, Jones P, Jaffar S, Pinder M, Obare S, Doherty T, Haywood M, Snounou G, Gemetery B, Gathmann I, Royce C, McAdam K, Greenwood B: A randomized controlled trial of artemether/benflumetol, a new anti-malarial and pyrimethamine/sulfadoxine in the treatment of uncomplicated falciparum malaria in African children. Am J Trop Med Hyg 1998, 58:638-644.
29. Hatz C, Abdulla S, Mull R, Schellenberg D, Gathmann I, Kibata P, Beck HP, Tanner M, Royce C: Efficacy and safety of CGP 56697 (artemether and benflumetol) compared with chloroquine to treat acute falciparum malaria in Tanzanian children aged 1-5 years. Trop Med Int Health 1998, 3:498-504.
30. Mutabingwa T, Anthony D, Heller A, Hallett R, Ahmed J, Drakeley C, Greenwood BM, Whitty CJ: Amodiaquine alone, amodiaquine + sulfadoxine-pyrimethamine, amodiaquine +artesunate, and artemether-lumefantrine for outpatient treatment of malaria in Tanzanian children: a four-arm randomised effectiveness trial. Lancet 2005, 365:1474-1480.
31. Checchi F, Piola P, Fogg P, Bajunirwe F, Biraro S, Grandesso F, Ruanggira E, Babigumira J, Kigozi I, Kiguli J, Kyomuhendo J, Ferradini L, Tay lor WR: Supervised versus unsupervised antimalarial treatment with six-dose artemether-lumefantrine: pharmacokinetics and dosage-related findings from a clinical trial in Uganda. Malaria J 2006, 5:59.
32. Kabanywanyi AM, Mwita A, Sumari D, Mandike R, Mugitu K, Abdulla S: Efficacy and safety of artemisinin-based antimalarial in the treatment of uncomplicated malaria in children in southern Tanzania. Malar J 2007, 6:146.
33. Koram KA, Abukai B, Duah N, Quashie N: Comparative efficacy of antimalarial drugs including ACTs in the treatment of uncomplicated malaria among children under 5 years in Ghana. Acta Trop 2005, 95:194-203.
34. Fanello CI, Karena C, van Doren W, Van Overmeir C, Ngamije D, D’Alessandro U: A randomised trial to assess the safety and efficacy of artemether-lumefantrine (Coartem) for the treatment of uncomplicated Plasmodium falciparum malaria in Randa. Trans R Soc Trop Med Hyg 2007, 101:344-350.