Abstract

Background: Deficiencies of vitamin A and its precursors, the carotenoids are common problems in developing countries. Plasma levels of these components are used as biomarkers of their availability. The study was conducted to evaluate whether blood plasma obtained from capillaries can be compared with plasma obtained from venous blood with regard to its levels of retinol, carotenoids and \( \alpha \)-tocopherol and secondly to apply this technique to evaluate the levels of these components in children in a region with possible deficiencies.

Methods: The survey was conducted in a region of Laos in 81 children (age 35 to 59 months). Dietary intake was assessed by a questionnaire. Retinol, carotenoids and \( \alpha \)-tocopherol were determined by HPLC. Blood plasma was obtained either from capillary blood collected into microcapillaries and for reasons of methodological comparison in 14 adults from venous blood.

Results: The comparison between capillary and venous blood revealed that all components except zeaxanthin were 9 – 23 % higher in plasma obtained from capillaries with regard to its levels of retinol, carotenoids and \( \alpha \)-tocopherol and secondly to apply this technique to evaluate the levels of these components in children in a region with possible deficiencies.

Conclusions: Results show that in consideration of slightly lower levels than in venous blood, capillary blood can be used to evaluate retinol, carotenoids and \( \alpha \)-tocopherol as biomarkers of intake or status and to evaluate the possible effect of diet on absolute and relative carotenoid composition in children from Europe and Laos. Observed sex related differences might not be related to diet and would need further investigation.
Background
On a worldwide scale, more than 1/3rd of all children suffer from protein malnutrition or undernutrition, with the highest prevalence showing in the developing countries of Asia, Africa, Latin America and Oceania. In Asia 80%, in Africa 15% and in Latin America 5% of all children are suffering from consequences caused by protein malnutrition or undernutrition [1]. For more than 230 million children an inadequate vitamin A supply is shown, with 13 million of them being affected by night blindness. The main cause for the loss of their eye sight of currently world wide more than 2.8 million preschool children is dietary vitamin A deficiency [2]. Moreover, insufficient intake of vitamins and minerals is suggested to be a cause for the high prevalence of several sub-clinical diseases and growth retardation. This is especially true for Laos, where UNICEF noticed a growth retardation in all Laotian preschool children, which was classified by the WH0 to be the highest in the world. Concerning the widespread night blindness, the national vitamin A study established that the intake of mainly rice as dominant staple food together with recurrent infectious diseases are the main causes of the vitamin A deficiency [3-5].

Vitamin A deficiency is caused by either a deficiency of vitamin A itself or by a reduced availability of precursors of vitamin A, the carotenoids. The reduced availability is either caused by its limited presence in food or by a reduced absorption due to a reduced fat content of the diet or a reduced absorptive capacity of the gut [6]. In addition to their function as precursor of vitamin A [7], carotenoids together with vitamin E are important micronutrients to maintain health by protecting the body against free radical damage [8]. In this context different epidemiological studies show an association with a variety of degenerative diseases such as cancer, heart disease and age related macula degeneration [9].

Because carotenoids, tocopherol and retinol plasma levels are biomarkers of their uptake or status [10,11] these components are frequently investigated in populations with an affluent lifestyle especially with regard to factors affecting plasma levels of these components [12]. Only a few studies have addressed this question in developing countries especially with regard to infants. This is, however, of specific importance to understand firstly how the plasma levels of these components are influenced and secondly how these components might determine longevity and long-term health.

Major reasons for the limited number of studies available on the plasma concentrations in individuals from developing countries are on the one hand logistical problems and on the other hand the limited individual and public acceptance of such kind of investigations. Logistical problems include blood collection, serum preparation by centrifugation, transportation as well as short-term and long-term storage under cool or frozen conditions. Mothers are not very enthusiastic with regard to an invasive sampling of blood especially in their infants. Therefore methods that will avoid the invasive venous blood sampling and a majority of logistical problems as well would be of advantage to encourage more studies on the levels of vitamin A, α-tocopherol and carotenoids in populations of developing countries.

The purpose of this study was thus to answer two questions. Firstly, whether plasma levels of retinol, α-tocopherol and carotenoids determined in capillary blood will validly represent the respective venous blood levels and secondly, whether a method using capillary blood plasma is suitable to assess the possible relation of vitamin A, carotenoid and α-tocopherol status in infants with sex, growth failure and living conditions in children from the high- and lowland region of the Bolikhamsay province Laos.

Subjects and Methods

Subjects
Within a nutritional study of the Family Health Project in Laos in 2000/2001, a total 1514 Laotian preschool children of the Province of Bolikhamsay were examined with regard to their nutrition, health and socio-economic situation in order to determine different causal factors between infants having a stunted growth and those with a normal growth. All children aged 0 to 59 months were eligible for entry in this study. The regional ethic committee approved the study. Participation was voluntary and informed consent was obtained from all mothers of participating children.

Data and sample collection
Laotian field workers collected information by questionnaire that included social-economic-education status, nutritional habits and queries to diseases.

Anthropometric measurements comprising height and weight, were measured by standardized equipment.

The daily food intake by quantity and amount was investigated over three consecutive using an estimated food record. The results were analysed with a computer programme called INMUCAL, which were developed by the Institute of Nutrition of the Mahidol University in Bangkok. The food frequency questionnaire (FFQ) determined the regular nutrition habits of the sample children over a period of time. Afterwards the daily consumption were calculated. The mothers were asked in an interview to estimate how often (daily, weekly, monthly, yearly) the child eats given food items from a list. The amount of a
consumed food item the mother had to specify by using the cup and spoon set from Thailand. The food item from the given list were scaled with a digital scale according to the different sizes of the cups and spoon (SOEHNLE attache 8020, max. 2 kg, division 2 g, Germany).

Blood samples were obtained from a total of 81 children of different sex aged between 35 and 59 months. Data for the European control group (n = 101) has been published previously [13] and are included in this paper for reasons of comparison. Capillary blood was drawn into heparinized microcapillaries (60 µl) by puncturing the pad of the finger. To obtain plasma samples the microcapillaries were immediately centrifuged at the site for 4 minutes at 11500 g in a battery-driven miniature centrifuge (M 1101, Bayer Diagnostik GmbH, Germany). The plasma was transferred into opaque Eppendorf cups and stored in an ice-filled box. Within four hours they were transferred to the nearest hospital to be frozen at -15°C. From there they were shipped by air on dry ice to Germany where they were stored at -80°C. Analysis was performed within six months.

In order to compare the carotenoid and vitamin concentrations between capillary and venous blood, blood samples were obtained simultaneously from 14 individuals of different sex and age recruited from the institutional staff who gave informal consent to this sampling. Venous blood samples were taken from the antibrachial vein, capillary blood was drawn into heparinized microcapillaries (60 µl) by puncturing the pad of the finger. Samples were centrifuged and treated in a comparable way.

**Determination of retinol, carotenoids and α-tocopherol by HPLC**

Carotenoids (lutein, zeaxanthin, α-carotene, β-carotene, β-cryptoxanthin, lycopene), α-tocopherol and retinol and retinyl esters were separated and quantified by reversed-phased HPLC [14]. Due to the limited amount of sample available the method was modified. Briefly, 200 µl of ethanol were added to 20 µl plasma diluted with 180 µl H2O. After vortexing for 30 sec, the samples were extracted twice with n-hexane (1 ml each time stabilized with 0.05 % butylated hydroxytoluene (BHT)) and vortexed for 3 min. The supernatants were removed, pooled, and evaporated under nitrogen and reconstituted in 50 µl isopropanol and injected (40 µl) into the HPLC-system (Waters, Eschborn, Germany). Accuracy and precision of the analyses were verified using a standard reference material (SMR 968a fat-soluble vitamins in human serum; National Institute of Standards and Technology (NIST), Gaithersburg, MD, USA). Coefficient of variability over time using control plasma was less than 4 % for carotenoids, retinol, retinyl esters and α-tocopherol. The recovery rate was above 95% for all components.

**Statistical analyses**

Normally distributed data are expressed as means and standard deviation (SD), and non-normally distributed data are expressed as medians and ranges. Statistical analysis was performed with a paired t test. Probability values below 0.05 were considered significant.

**Results and Discussion**

Only a few studies have addressed the question of plasma carotenoid levels in children. These data however are important because nutritional deficiency has many detrimental effects with short and long-term consequences for children. These include growth retardation, reduced cognitive functions and vision as well as an increased incidence of infectious diseases. As single causes or in combination these factors increase the mortality rates in developing countries [15,16]. Therefore, monitoring the nutritional status is very important. Methods that facilitate the acquisition of such data are highly necessary because limited information still exists with regard to major determining influences on the levels of carotenoids, α-tocopherol and retinol for children in developing countries. This also concerns the interaction with factors of nutritional origin, or the important influence of the health status such as the acute phase response during infections [17,18]. One major advantage would be gained by reducing the invasiveness of blood sampling and the reduction of sample volume necessary to analyse key nutrients such as retinol, carotenoids and α-tocopherol [19,20].

**Comparison of plasma composition between samples obtained from venous or capillary blood**

Table 1 shows differences in plasma retinol, individual carotenoids (lutein, zeaxanthin, α-carotene, β-carotene, β-cryptoxanthin, lycopene) and α-tocopherol between plasma obtained from the same individuals either by puncturing of a vein or the capillaries of the fingertip. In general, all components were higher in plasma obtained from capillary blood. The most obvious differences were observed for α- and β-carotene. This observation is in accordance with a previous study in which, however, only β-carotene and α-tocopherol were investigated [21]. The increased levels have been attributed to the reduced fluid in capillaries due to differences in hydrostatic pressure and colloid osmotic pressure in the capillaries. In this case, however, one would expect that all components measured would show similar quantitative changes. Nevertheless, the differences between plasma obtained from capillaries or veins are in the same order of magnitude. Thus, capillary blood sampling is a valid procedure to determine micronutrients such as retinol, α-tocopherol and carotenoids under field condition not only in developing countries but also in neonatal and pediatric medicine. With regard to the situation in developing countries,
the procedure not only eliminates invasive blood sampling but also greatly reduces the equipment needed. Plasma can be obtained using a battery powered centrifuge and the small sample volumes can be stored and transported in well isolated storage containers even under less favourable conditions.

**Table 1: Differences in the concentration of retinol, carotenoids and α-tocopherol (µmol/l; mean ± SD) between plasma samples (n = 14) obtained from paired venous or capillary blood**

| Component       | Venous     | Capillary  |
|-----------------|------------|------------|
| α-Tocopherol    | 20.80 ± 10.02 | 23.35 ± 10.02* |
| Retinol         | 1.42 ± 0.31  | 1.69 ± 0.39*  |
| Lutein          | 0.21 ± 0.08  | 0.33 ± 0.09  |
| Zeaxanthin      | 0.02 ± 0.01  | 0.02 ± 0.01  |
| β-Cryptoxanthin | 0.11 ± 0.05  | 0.13 ± 0.06  |
| α-Carotene      | 0.11 ± 0.07  | 0.14 ± 0.09*  |
| β-Carotene      | 0.54 ± 0.28  | 0.66 ± 0.32*  |
| Lycopene        | 0.19 ± 0.12  | 0.25 ± 0.13***|

* P < 0.05 *** P < 0.001

**Table 2: Differences in the concentration of retinol, carotenoids and α-tocopherol (µmol/l; mean ± SD) between European and Laotian children and effect of sex**

| Component       | Europe all sexes | Laos girls | Laos boys |
|-----------------|------------------|------------|-----------|
| α-Tocopherol    | 15.84 ± 4.94     | 1.885 ± 1.755 | 0.469 ± 1.504* |
| Retinol         | 1.22 ± 0.27      | 0.967 ± 0.230 | 0.931 ± 0.178* |
| Lutein          | 0.31 ± 0.21      | 0.063 ± 0.050 | 0.003 ± 0.002***|
| Zeaxanthin      | 0.02 ± 0.02      | 0.004 ± 0.005 | 0.003 ± 0.002***|
| β-Cryptoxanthin | 0.16 ± 0.14      | 0.009 ± 0.020 | 0.005 ± 0.011***|
| α-Carotene      | 0.65 ± 0.43      | 0.003 ± 0.003 | 0.002 ± 0.007 |
| β-Carotene      | 1.57 ± 1.15      | 0.014 ± 0.018 | 0.006 ± 0.018***|
| Lycopene        | 0.26 ± 0.10      | traces       | traces     |

* P < 0.05 *** P < 0.001

**Concentration of retinol, carotenoids and α-tocopherol in plasma samples of children from Laos**

Table 2 summarizes the results on plasma concentrations of retinol, carotenoids and α-tocopherol and shows obvious differences between groups of children sampled in Europe and in Laos as well as sex differences in Laotian children. Studies that determined the levels of these components in plasma are limited and mostly restricted to developed countries. One reason for this, as has been pointed out earlier, are the difficulties to obtain appropriate samples. As reported for adults and children in Nepal and Africa the levels observed for carotenoids are substantially lower in children from Laos compared to slightly older children from Austria [13] or Germany [22] or from much older ones in the USA [23]. Similarly low levels were found in children from Nigeria and Pakistan [24,25]. The differences in dietary carotenoid pattern are further strengthened by the striking differences in the carotenoid pattern. While in European infants the dominant carotenoid was β-carotene representing 52% of total carotenoids, in infants from Laos the dominant carotenoid was lutein with 62% (P < 0.001) (Table 3). Both lower carotenoid plasma levels as well as differences in the plasma carotenoid pattern clearly reflects differences in the quantitative and qualitative availability of carotenoids. High lutein levels are indicative of a consumption of dark green leafy vegetable and are frequently found as dominant carotenoid in plasma samples obtained from populations in developing countries [24]. The relative contribution of individual carotenoids to total carotenoids in Laotian children was independent of origin or sex (data not presented).

The numerically smaller difference between the two groups was observed for plasma retinol, which can be
explained by the homeostatic regulation of its plasma levels [26]. In none of the children plasma levels were indicative of deficiency. Despite low vitamin A intakes, plasma retinol concentrations of an average of 0.91 – 1.07 µmol/l (95% C.I.) could be measured. Based on our protocol stunted children got 39 % and non-stunted children got 43 % of the recommended daily requirements. Other studies, however, show strong relationship between low vitamin A intake and low serum retinol concentrations [27]. The cause of this contradiction might be the homoeostatic regulation of the retinol and possible sufficient amounts present in the main storage organ for vitamin A, the liver. As shown in Table 4 the effect of growth retardation and the area in which the children lived was of less importance for the level of retinol, carotenoids and α-tocopherol in plasma.

Table 2 shows that in children from Laos substantial sex differences exist with regard to the levels of retinol, carotenoids and α-tocopherol. In boys from Laos all components were significantly lower (P < 0.01). This is in contrast to our results from Europe and studies from Pakistan in children and from Algeria in adults, in which no such sex differences were observed [13,22,24,28]. In the study conducted in Pakistan however, levels of lycopene and β-cryptoxanthine were below detection limit and β-carotene was in traces in only 8% of the children and in the study from Algeria only β-carotene and α-tocopherol were measured. The gender-specifically higher plasma concentrations of carotenoids in girls, however, have been reported by one study [29]. Another study found slightly higher levels of carotenoids only significant for lycopene in boys [23]. Based on the nutritional questionnaires taken in this study, this phenomenon cannot be explained by differences in nutrition. Additionally, the age of the children would exclude known effects of sex hormones on plasma levels of the investigated components.

### Conclusions

Our results show that first, there is the possibility of using capillary blood plasma to evaluate retinol, carotenoids and α-tocopherol as biomarkers of intake or status and the possible effect of diet on absolute and relative carotenoid composition. This modification including glass capillaries and a battery-driven centrifuge can substantially reduce the equipment necessary for plasma preparation,
transportation and storage. Second, that the application of this method in Laotian children shows important factors that influence plasma levels of retinol, carotenoids and α-tocopherol in Laotian infants such as sex and to a limited extent stunting and the region of origin.

**Competing interests**
None declared.

**Authors’ contributions**
FJS participated in the conception, design, data analysis and writing of the manuscript. JK participated in design, data analysis and writing of the manuscript. AH participated in the data analysis and the writing of parts of the manuscript. HJZ participated in the conception, design data analysis and critical revision of the manuscript. All authors have red and approved the last version of the manuscript.

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