Folate status of Ghanaian populations in London and Accra

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Migration to the UK is associated with higher incidence of stroke in African populations. A low folate status has been associated with increased risk of stroke, likely to be mediated through raised plasma homocysteine concentrations. We conducted a cross-sectional study to compare blood folate and homocysteine concentrations in eighty healthy Ghanaian migrants living in London matched by sex, age and occupation to 160 individuals from an urban population in Accra, Ghana. Folate intake was determined using three 24 h recalls. Fasting blood samples were collected for the determination of serum and erythrocyte folate and plasma homocysteine concentrations and the methylenetetrahydrofolate reductase (MTHFR) 677C → T polymorphism. Reported mean folate intake was 20% lower in London compared with Accra (P<0.001). However, serum folate was 44% higher, erythrocyte folate 30% higher and plasma homocysteine was 26% lower in subjects from London compared with those from Accra (P<0.001). These differences persisted after adjusting for confounders including the MTHFR 677C → T mutation, which was rare in both populations. Although there were no associations between dietary folate intake and blood folates (P>0.05), folic acid supplement use, which was more prevalent in London than Accra (25 and 10%, respectively, P=0.004) was associated with erythrocyte folate in both populations (P<0.01). The main predictors of plasma homocysteine concentrations were erythrocyte folate and male sex (P<0.001).

Findings from the present study suggest that migration from Ghana to the UK results in improvement of biomarkers of folate status despite the fact that reported dietary intake of folate was apparently lower in subjects from London.

Folate: Homocysteine: Ghana: United Kingdom

People of African origin living in the UK have a higher incidence and mortality from stroke than white people1,2. Changes in dietary patterns are likely to contribute to these differences, particularly since migration to the UK has been associated with increased incidence of stroke3. A low folate status increases the risk of stroke4 and a recent meta-analysis of randomised controlled trials has shown that folate acid supplementation reduces the risk in primary prevention5. Folic acid may prevent stroke by lowering raised plasma homocysteine concentrations, which have been linked to endothelial dysfunction6.

Folate status can be determined by assessing dietary intake of folate and measuring serum and erythrocyte folate (which reflect recent dietary intake and tissue levels, respectively). Plasma homocysteine is also used as a functional indicator of folate status since homocysteine concentrations rise even with marginal folate depletion7. Clinical trials have shown that moderately increasing folate intake from dietary sources and vitamin supplements can reduce homocysteine concentrations effectively8,9. In white populations, a common mutation in a critical enzyme in folate metabolism, methylenetetrahydrofolate reductase (MTHFR 677C → T), is a major determinant of plasma homocysteine concentrations, but the prevalence of this polymorphism is low in populations of African descent10–12; therefore folate status is likely to be the main determinant of homocysteine concentrations in these individuals.

There are no data comparing folate status in UK African migrants with populations from their country of origin (sedentary). A recent study has reported lower serum folate and higher plasma homocysteine in healthy Afro-Caribbeans in the UK compared with a matched population in Jamaica13. However, studies comparing folate and homocysteine concentrations in other ethnic populations in the UK have reported opposite effects; for example, UK Gujeratis had higher dietary folate intakes and serum folate and lower plasma homocysteine than those in India14. This indicates the need for ethnic group-specific studies to identify diet-related biomarkers that may influence risk of disease.

Ghanaians form the UK’s fifth largest group of African migrants, with an estimated 50 000–1 000 000 individuals living in London15. To our knowledge there is no available information on the folate status of Ghanaian migrants in the UK or in their country of origin. In the present study, we compared dietary folate intakes, serum and erythrocyte folate and plasma homocysteine concentrations in a Ghanaian migrant population in the UK matched with an urban population from Accra, Ghana, and determined the predictors of plasma homocysteine concentrations in this population. The MTHFR 677C → T polymorphism was determined

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Abbreviation: MTHFR, methylenetetrahydrofolate reductase.

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to confirm its low prevalence in this population and account for any confounding effects on plasma homocysteine concentrations.

Methods

Study population

Subjects in the present study were part of a cross-sectional study examining the relationship between diet and CVD risk factors in two Ghanaian populations: a first-generation Ghanaian migrant population in London, and an urban population in Accra, Ghana, their country of origin. A total of 240 healthy subjects (London, eighty subjects; Accra, 160 subjects) aged between 25 and 50 years were recruited from Ghanaian organisations in London and Accra. In order to provide robust comparisons, subjects in London were matched for age (± 2 years), sex, and occupation to participants from Accra. Matching was done prospectively. Two Accra-based participants were matched to one London-based participant. London-based participants were included if they were aged between 25 and 50 years and had resided in the UK for at least 5 years. Participants were ineligible if they were diabetic, pregnant or lactating or had any major underlying organ pathology.

Participants recruited into the study reported at the approved study centres in London and Accra and were assessed regarding demographic characteristics, folate intake, serum and erythrocyte folate, vitamin B12 and plasma homocysteine concentrations with questionnaires, interviews, physical examination and blood sampling. The study was carried out according to protocols approved by the King’s College London and University of Ghana Ethics Committees.

Anthropometric measurements

Participants underwent standard physical measurements including weight, height, waist circumference and blood pressure. Weight was measured with a Seca 762 scale (Vogel & Halke, Hamburg, Germany) to the nearest 0·1 kg in light clothing with subjects standing erect. Height was measured using a portable stadiometer and made to the nearest 0·1 cm. The BMI was calculated by dividing the body weight (in kg) by the square of the height (in metres).

Dietary assessment and general characteristics questionnaire

A self-administered questionnaire was used to obtain data about demographic characteristics, use of dietary supplements, health status, prescribed medications, and smoking and drinking habits. The questionnaire was pre-tested in a pilot study after which minor additions and changes were made to make it more presentable. Dietary intake was collected by a trained dietitian using the 24 h recall method for 3 d including a weekend day. The first interview was done on the day participants reported at the study centres and the further two were performed on the phone within the same week. Portion sizes were estimated using household measures common to the Ghanaian community.

Calculation of dietary folate intake

Nutrient intakes were estimated using the nutritional package Microdiet (version 1.2 software; Downlee Systems Ltd, Chapel-en-le-Frith, Derby, UK) with an added database incorporating McCance and Widdowson’s compositional data on immigrant groups and an updated database on the nutritional composition of commonly consumed Ghanaian dishes (M Owusu, unpublished results). Food folate and synthetic folic acid added to foods is handled as a single value in Microdiet. Folate content of the Ghanaian dishes was determined by microbiological assay using Lactobacillus rhamnosus (American Type Culture Collection (ATCC) 7469) after thermal extraction and tri-enzyme treatment.

Biochemical assessment

Fasting blood samples were obtained in the morning after a 12 h fast. For serum folate and vitamin B12 measurement, blood samples were collected into a vacutainer containing no anticoagulant, left to clot for 1 h at room temperature and then centrifuged at 2500 rpm for 15 min. Serum was transferred into plastic vials and frozen immediately. Blood samples for analysis of erythrocyte folate were collected into vacutainers containing EDTA and sent to the laboratory within 24 h of collection for full blood count and folate analyses. In Accra, it was not possible to run the assay for serum and erythrocyte folate; therefore, serum and whole blood samples were frozen at −80°C and shipped to London for analyses. However, full blood counts were performed in Accra within 24 h of blood collection. Fasting blood samples for homocysteine determination were collected into EDTA-containing vacutainer tubes, placed on ice and carried to the laboratory where the plasma was centrifuged at 3000 rpm for 10 min, transferred into plastic vials and frozen at −80°C until analysis. For DNA extraction, whole blood samples were collected into vacutainer tubes containing EDTA and frozen at −80°C until required for DNA extraction. Samples from Accra were transported on dry ice to King’s College London and refrigerated at −70°C until the analyses were performed.

Laboratory analysis

All samples were analysed at the Department of Clinical Biochemistry at King’s College Hospital, London. Serum folate, B12 and erythrocyte folate were analysed using the Bayer Advia Centaur folate kit (Bayer Diagnostics Europe Ltd, Swords, Co. Dublin, Republic of Ireland) and plasma homocysteine by the Bayer Advia Centaur homocysteine kit (Bayer Diagnostics Europe Ltd). For all assays, the within-run CV was < 2·2 % and between-run CV was < 5·2 %.

DNA extraction

The DNA for genotyping was extracted from whole blood using the Puregene DNA isolation kit (product number D5000; Gentra Systems, Inc., Minneapolis, MN, USA). DNA fragments were amplified from the genomic DNA with the PCR. The MTHFR 677C → T substitution was identified with the use of restriction enzyme digestion of the
PCR-amplified products, as previously described by Frosts et al. (17). The PCR product was digested using HinfI (GE Healthcare, Chalfont St Giles, Bucks, UK). The 677C → T polymorphism creates a HinfI recognition site in a 198 bp amplified DNA fragment. This was used to differentiate the various genotypes: CC, CT and TT.

Statistical analysis

Statistical analyses were performed using SPSS for Windows (version 14; SPSS, Inc., Chicago, IL, USA). Nutrient intakes were adjusted for energy using the method of residuals. This involved computing residuals from a regression model with energy intake as the independent variable and folate intake as the dependent variable (18). Residuals were then added to the mean folate intake for the group to obtain a score adjusted to the average energy intake. To compare differences in categorical variables between groups, χ² tests or Fisher’s exact tests were used. Independent-sample t tests were used to compare differences in age and BMI between locations, as these variables were normally distributed. Nutrient intakes (vitamin B₆, vitamin B₁₂ and folate) and biomarkers (serum and erythrocyte folate, serum vitamin B₁₂ and plasma homocysteine) were skewed and transformations failed to normalise the data; therefore Mann-Whitney U tests were used to compare differences between these variables. To test for the effect of potential confounders the analyses were repeated adjusted for age, sex, BMI, educational level, alcohol intake and the MTHFR genotype. Since the adjustment did not have any influence on the observed differences, only the unadjusted data are presented. Spearman’s correlation coefficients were used to investigate the relationship between nutrient intakes and biomarkers and between MTHFR genotype and blood folates and plasma homocysteine. Multiple linear regression analysis was performed to examine predictors of blood folate and homocysteine concentrations in each population. Differences were considered significant at P<0.05.

Results

Baseline characteristics and nutrient intakes of subjects

General characteristics and mean daily nutrient intakes of the study population are presented in Table 1. There were no significant differences between groups in any of the baseline variables measured, except that a greater proportion of London subjects had attained a higher education (tertiary) compared with subjects from Accra (57 v. 38 %; P=0.001). Allele frequencies for the MTHFR 677C → T mutation were 10 % in London and 8 % in Accra; overall, the prevalence of the TT genotype was 0.5 %. There was a greater proportion of supplement users in London compared with Accra (25 v. 10 %; P=0.004). Reported intakes of vitamins B₆, B₁₂ were higher in London compared with Accra (by 15 %, P=0.03 and 6 %, P=0.04, respectively). However, reported folate intakes (diet and supplements) were 20 % lower in London compared with Accra (P<0.001).

The main dietary sources of folate in London and Accra are shown in Table 2. The largest sources of folate in London were Ghanaian soups and bean dishes, accounting for 33 % of folate intake. This was followed by vegetable stews and fermented maize products (18 %). In addition, foods such as breakfast cereals, fruits and fruit juices and mixed vegetables also contributed to folate intake in London (accounting for 20 %). In Accra, the largest sources of folate were from Ghanaian soups, vegetable stews and bean dishes (61 %). Other dishes that contributed to folate intake were fermented maize products and fruits (20 %).

Biomarkers of folate and vitamin B₁₂ status

The mean serum folate, erythrocyte folate, plasma homocysteine and serum vitamin B₁₂ in subjects from London and Accra are shown in Table 3. Serum and erythrocyte folate was higher (by 44 and 30 %, respectively; P<0.001) and plasma homocysteine was lower (by 26 %; P<0.001) in participants from London compared with those from Accra. The differences between London and Accra were still significant (P=0.001) after controlling for age, sex, BMI, educational level, alcohol intake and the MTHFR 677C → T genotype.

There were no significant correlations between total folate intake (diet and supplements) and serum folate or erythrocyte folate in either London (r 0.208, P=0.068; r 0.189, P=0.103; respectively) or Accra (r 0.046 and r 0.071, respectively; P>0.05). There were no significant correlations between dietary folate intake in non-supplement users and serum or erythrocyte folate in either London (r 0.155 and r 0.028, respectively; P>0.05) or Accra (r 0.041 and r 0.032, respectively; P>0.05). There was, however, a positive correlation between supplemental folate intake and erythrocyte folate in London and Accra (r 0.419, P=0.001; r 0.225, P=0.005, respectively). There was a significant positive correlation between serum folate and erythrocyte folate in London (r 0.385; P=0.001) and Accra (r 0.352; P<0.001). Regression analysis confirmed that serum folate and folic acid supplement use were predictors of erythrocyte folate in both London and Accra (P<0.001). No significant association was found between MTHFR 677C → T genotype and serum folate or erythrocyte folate in either population (P>0.05).

There were negative correlations between serum folate and plasma homocysteine (London, r −0.192, P=0.09; Accra, r −0.325, P<0.001) and erythrocyte folate and plasma homocysteine (London, r −0.388, P=0.001; Accra, r −0.290, P<0.001). Regression analysis showed that erythrocyte folate was the main predictor of plasma homocysteine concentrations in London (P<0.001) while in Accra the main predictor was male sex (P=0.001). There were negative correlations between serum vitamin B₁₂ and plasma homocysteine in both London (r −0.289; P=0.01) and Accra (r −0.140; P=0.08). There was no association between the MTHFR 677C → T genotype and plasma homocysteine in either location (P>0.05).

Table 4 shows the biomarkers of folate status in folic acid supplement users and non-users in London and Accra. Compared with non-supplement users, folic acid supplement users had higher serum folate (by 24 % in London, P=0.09, and 67 % in Accra, P=0.008) and erythrocyte folate (by 33 % in London, P<0.001, and 32 % in Accra, P=0.006), and lower plasma homocysteine (by 16 % in London, P=0.150, and 21 % in Accra, P=0.01). Folic acid supplement
users in London had higher erythrocyte folate (28 %, $P=0.09$) and lower plasma homocysteine (22 %, $P=0.03$) compared with folic acid supplement users in Accra. However, non-users of folic acid supplements from London also had significantly higher serum and erythrocyte folate (by 48 %, $P=0.001$ and 27 %, $P=0.001$, respectively) and lower plasma homocysteine (by 27 %; $P=0.001$) compared with non-users from Accra.

**Discussion**

The present study compared folate status in a first-generation Ghanaian migrant population living in London with an urban population in Accra, Ghana. Results showed that reported folate intakes in London were significantly lower compared with those in Accra even though folate from supplements was added to daily intakes and there were more supplement users in London. The higher folate intakes in Accra resulted from large reported intakes of traditional Ghanaian vegetable stews and soups in addition to other foods processed from fermented maize products (Table 3). Reported median folate intakes in subjects from London (417 mg/d in men and 507 mg/d in women) were higher than median intakes of the average UK population (335 mg/d in men and 255 mg/d in women) (19).

Despite reporting lower intakes of folate, subjects from London had a more favourable folate status than subjects in Accra, with higher serum and erythrocyte folate ($P=0.004$) and lower plasma homocysteine concentrations ($P<0.001$). It is unclear why there were such marked differences between London and Accra.

### Table 1. Demographic characteristics and nutrient intakes of study participants

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<td>391</td>
<td>389†††</td>
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<td>Folate (diet and supplements) (µg/d)</td>
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Proportion was significantly different from that of the Accra group: * $P=0·004$, ** $P=0·001$ ($x^2$ test or Fisher’s exact test).

Median value was significantly different from that of the Accra group: † $P=0·04$, †† $P=0·03$, ††† $P<0·001$ (Mann–Whitney U test).

‡ London, n 77; Accra, n 145.
reported dietary intakes and biomarkers in these two populations. Although misreporting of dietary intake is known to occur with the 24 h recall\(^{20,21}\) and this method of dietary assessment has not been validated for B vitamin intake in this population, folate intakes in the present study were based on an average of three 24 h recalls, which included a weekend day, and there was a similar proportion of underreporters in both London and Accra (30 and 21\%, respectively). A possible explanation for the difference in the dietary folate intake observed between subjects from London and Accra is the use of slightly different databases to determine folate intakes. The folate data generated for subjects in Accra, who typically consumed traditional Ghanaian foods, came mainly from our recently generated database on the nutritional composition of commonly consumed Ghanaian dishes, whereas the folate data for the subjects in London were determined from the recently generated database in combination with McCance and Widdowson’s Food Composition Tables. However, the folate content of foods in the two databases has been determined by different analytical methods; the recently generated database used the microbiological assay with tri-enzyme extraction according to the method of Tamura\(^{16}\), whereas the Food Composition Tables used the microbiological assay with the traditional conjugase treatment alone. The application of tri-enzyme extraction has been reported to increase considerably the measurable folate in single foods and in composite meals\(^{16,22}\), and this could account for the higher folate intakes in the Accra population in the present study.

One possible explanation for the higher blood folate concentrations in the London population may be the higher consumption of folic acid from fortified foods and supplements. Folic acid has superior bioavailability compared with natural folates\(^{23}\) and it is well established that higher intakes of foods naturally rich in folates are a relatively ineffective means of increasing folate status compared with equivalent intakes of folic acid-fortified food\(^ {24}\). Breakfast cereals were rarely consumed in Accra but contributed to 5\% of the folate intake in London. Also, multivitamins containing folic acid were consumed by 25\% of participants from London compared with 10\% of participants from Accra and there was a strong association between reported use of folic acid supplements and erythrocyte folate concentrations in both

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**Table 2. Major foods contributing to folate intake in study participants**

<table>
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<tr>
<th></th>
<th>London (n 80)</th>
<th>Accra (n 160)</th>
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<tbody>
<tr>
<td>Light, palm nut or peanut soup</td>
<td>20</td>
<td>22</td>
</tr>
<tr>
<td>Bean dishes (bean stew, rice and beans)</td>
<td>13</td>
<td>19</td>
</tr>
<tr>
<td>Spinach*/garden egg stews, okro stews and soups</td>
<td>9</td>
<td>20</td>
</tr>
<tr>
<td>Fermented maize dishes (banku, kenkey, porridges)</td>
<td>9</td>
<td>11</td>
</tr>
<tr>
<td>Fruit juice</td>
<td>6</td>
<td>–</td>
</tr>
<tr>
<td>Ghanaian stew or gravy</td>
<td>6</td>
<td>–</td>
</tr>
<tr>
<td>Fruits</td>
<td>5</td>
<td>9</td>
</tr>
<tr>
<td>Breakfast cereals</td>
<td>5</td>
<td>–</td>
</tr>
<tr>
<td>Mixed vegetables</td>
<td>4</td>
<td>–</td>
</tr>
</tbody>
</table>

*Participants in Accra consumed 'kontomire' leaves, which are similar to spinach but not found commonly in the UK.

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**Table 3. Biomarkers of folate and vitamin B\(_\text{12}\) status in study participants**

<table>
<thead>
<tr>
<th></th>
<th>London (n 80)</th>
<th>Accra (n 160)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Men (n 43)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Median</td>
<td>32–25.9</td>
<td>12-0</td>
</tr>
<tr>
<td>Range</td>
<td>3.2–26.5</td>
<td>7.3–1310</td>
</tr>
<tr>
<td>Women (n 37)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Median</td>
<td>374–1086</td>
<td>7.3–1310</td>
</tr>
<tr>
<td>Range</td>
<td>5.9–17.8</td>
<td>150–1086</td>
</tr>
<tr>
<td>All (n 80)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Median</td>
<td>599</td>
<td>451</td>
</tr>
<tr>
<td>Range</td>
<td>286–880</td>
<td>135–1086</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th></th>
<th>Accra (n 160)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Men (n 86)</td>
<td></td>
</tr>
<tr>
<td>Median</td>
<td>3.2–26.5</td>
</tr>
<tr>
<td>Range</td>
<td>374–1310</td>
</tr>
<tr>
<td>Women (n 74)</td>
<td></td>
</tr>
<tr>
<td>Median</td>
<td>501</td>
</tr>
<tr>
<td>Range</td>
<td>181–1695</td>
</tr>
<tr>
<td>All (n 160)</td>
<td></td>
</tr>
<tr>
<td>Median</td>
<td>616</td>
</tr>
<tr>
<td>Range</td>
<td>229–2094</td>
</tr>
</tbody>
</table>

* Median value was significantly different from that of the Accra group (P<0.001; Mann–Whitney U test).
Table 4. Biomarkers of folate status in users v. non–users of folate acid supplements (Medians and ranges)

<table>
<thead>
<tr>
<th>Folic acid supplement users</th>
<th>Non-users of folic acid supplements</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>London (n = 20)</td>
</tr>
<tr>
<td></td>
<td>Median</td>
</tr>
<tr>
<td>Serum folate (nmol/l)</td>
<td>13·8</td>
</tr>
<tr>
<td>Plasma homocysteine (µmol/l)</td>
<td>8·6*</td>
</tr>
</tbody>
</table>

Median value was significantly different from that of the Accra group: *P<0·05, **P<0·001, ***P<0·001 (Mann–Whitney U test).

locations (P<0·001). However, it was evident that folic acid supplement use was not solely responsible for the differences in biomarkers observed between the two populations, since non-users of folic acid supplements in London had significantly higher serum and erythrocyte folate (P<0·001 and P<0·001, respectively) and significantly lower plasma homocysteine (P=0·001) than non-users in Accra. Similarly, supplement users in London had higher erythrocyte folate (P=0·09) and lower plasma homocysteine (P=0·03) than supplement users in Accra. This suggests that folic acid from fortified foods may have made a substantial contribution to the differences observed in these biomarkers between London and Accra.

Estimation of the contribution of folic acid from fortified foods to the total folate intake in the present study was not possible since it was difficult to obtain information on the brands of breakfast cereal consumed by participants. Different manufacturers fortify their cereals with different amounts of folic acid and cereals are fortified with folic acid at higher levels than the values given in the current nutrient database, which has not been updated for 5 years. Furthermore, the nutritional package used in the present study did not separate natural folate from synthetic folic acid used as fortificant. It is possible that participants from London were consuming some other foods that were fortified with folic acid, which were not accounted for in the nutritional database. Recently, certain brands of margarines and fruit juice have been fortified with folic acid.

Although we used an updated database (which included information on folate content of Ghanaian foods) to estimate folate intake we did not find a relationship between dietary folate intakes and serum or erythrocyte folate as reported in some studies (25–27). However, other studies have found no association between dietary folate intake assessed by FFQ and 7 day weighed record and serum or erythrocyte folate in healthy populations (28,29). Possible reasons for the lack of association in the present study are underestimation of folic acid intake from fortified foods and misreporting of food intake by participants. Nutrient intake in the present study was by the 24 h recall and a major limitation of this method is the inability to remember foods eaten and accurate description of food (30). It is possible that the lack of agreement between folate intake and blood folate values are related to the reporting of food intake in this group of individuals because we observed no relationship between dietary vitamin B12 intake and serum vitamin B12 concentrations in the present study (P>0·05). Furthermore, the relatively high reported dietary folate intakes in our population are not reflected in the serum folate concentrations (10·9 nmol/l in men and 12·0 nmol/l in women) which are lower than those reported in the average UK population (18·8 nmol/l in men and 21·0 nmol/l in women) (31), although they are comparable with serum folate concentrations in populations of African descent in the USA (12·6 nmol/l in men and 15·5 nmol/l in women) (32). However, erythrocyte folate concentrations (675 nmol/l in men and 757 nmol/l in women) in subjects from London were comparable with those of the average UK population (633 nmol/l in men and 610 nmol/l in women) (31). The present results suggest that further research should be conducted in how to obtain reliable dietary intake data from this type of population.

The significantly higher erythrocyte folate concentrations in London participants were accompanied by significantly lower plasma homocysteine concentrations. Plasma homocysteine was 26% lower in London compared with Accra. The median concentrations found in men (10·9 µmol/l) and women (8·4 µmol/l) in London were similar to previously reported measurements of plasma homocysteine in populations of African descent in the UK and USA (11,33). Mean plasma homocysteine concentrations in London participants were within the range considered normal (<15 µmol/l); however, men from Accra had a median value exceeding this (16·0 µmol/l). Plasma homocysteine concentrations above 12 µmol/l have been consistently associated with vascular disease (34). By this definition hyperhomocysteinemia is prevalent in the healthy Accra population and may have implications for stroke incidence.

The main predictors of plasma homocysteine in our population were male sex and erythrocyte folate concentrations. Population studies generally show that increasing age, male sex, smoking, coffee consumption, high blood pressure and the MTHFR 677C → T polymorphism are among the factors associated with plasma homocysteine concentrations (35–37). The present study did not find any association between plasma homocysteine and age, which may be due to the narrow age range in our population (25–50 years) or the relatively small sample size. Similar to other population studies (36,38,39), the present study showed plasma homocysteine concentrations to be correlated with blood folates and vitamin B12.

The MTHFR 677C → T mutation was not associated with plasma homocysteine concentrations in the present study, despite the fact that the mutation is a known determinant of homocysteine concentrations and plasma folate (17). However,
the prevalence of the MTHFR 677C → T mutation was low in our population, a finding consistent with other studies in popula-
tions of African descent\textsuperscript{(10–12)}. The lack of association of the MTHFR 677C → T mutation with plasma homocysteine may be due to the low prevalence of the TT genotype and the sample size, which may have been too small to detect such a relationship. The low prevalence of the MTHFR 677C → T mutation suggests that folate status in London and Accra participants is mainly a reflection of dietary factors.

Conclusion

Migration to the UK leads to an improvement in biomarkers of folate status in Ghanaian migrants from London compared with a matched population from Accra. Findings of the sample may be generalisable to a similar group with the same educational and socio-economic status in London and Accra. However, this sample may not be representative of the total Ghanaian population in the UK and Ghana. Since an improved folate status is associated with prevention of stroke as well as other conditions, there is a need to undertake large-scale epidemiological studies within the general adult Ghanaian population to confirm these findings.

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M. O. was involved in patient recruitment, collection of data, statistical analysis and manuscript writing. J. T. was involved in experiment design and manuscript writing. E. W. was involved in collection of data and manuscript writing. M. P. was involved in experiment design, statistical analysis and manuscript writing.

The authors do not have any conflicts of interest to declare.

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