Invited Commentary

Invited commentary in response to: ‘Identification of vitamin B$_{12}$ deficiency in vegetarian Indians’

The proportion of humans adopting a plant-based lifestyle is on the rise worldwide. Benefits on economics, climate and health have been documented and projected to continue over the next decades(1–3). Apart from its sustainability(4), well-planned plant-based diets are nutritionally complete for human beings of all ages(5), with vitamin B$_{12}$ being the only micronutrient not found in plants(6–8). Vitamin B$_{12}$ is only synthesised by a few bacteria and archaea(9–11); hence, humans adhering to a plant-based lifestyle can obtain the micronutrient from foods fermented with bacteria that naturally produce vitamin B$_{12}$, foods fortified with vitamin B$_{12}$ and over-the-counter supplements. However, the high prevalence of vitamin B$_{12}$ insufficiency in vegetarians across nations(12) suggests that food literacy programmes(13) are not yet meeting the needs of the current dietary shift. Vitamin B$_{12}$ deficiency, whether acquired or inherited, leads to the inactivation of the two vitamin B$_{12}$-dependent enzymes, cysteolic methionine synthase and mitochondrial methylmalonyl-CoA mutase(14,15). This manifests with elevated homocysteine (Hcy) and methymalonic acid (MMA), and in severe cases with inactivation of the two vitamin B$_{12}$-dependent enzymes, cyto-

The study by Naik et al.(16) examined vitamin B$_{12}$ status in a cohort of 119 young, healthy unsupplemented vegetarian Indian graduates. Using standard cut-off values for the assessment of vitamin B$_{12}$ status Naik et al. determined that 50% were vitamin B$_{12}$ deficient, 50–70% exhibited low plasma holo-TC and 70–90% presented elevated plasma total Hcy (tHcy). All participants in this study were asymptomatic of clinical vitamin B$_{12}$ deficiency(16). These findings are in good agreement with a previous study by Refsum et al. in a different Indian cohort(17). Interestingly, nineteen participants with plasma vitamin B$_{12}$ concentrations between 113 and 122 pmol/l had normal values of holo-TC (34–52 pmol/l), while exhibiting elevated tHcy according to standard cut-off reference values (discussed in Hannibah et al.(18) and references therein). Naik et al.(16) propose a new set of cut-off values to improve the diagnosis of vitamin B$_{12}$ deficiency in young vegetarian Indians. The authors propose the use of a combination of biomarkers and cut-off values of 100 and 19.6 pmol/l for plasma vitamin B$_{12}$ and holo-TC, respectively, and values of tHcy of 17·6 and 27 µmol/l for females and males, respectively(16). This proposal appears in line with previously documented data in a study performed on a cohort of American vegetarians by the late Dr Victor Herbert(19). Herbert divided vitamin B$_{12}$ status in vegetarians in four distinct stages, namely I, II, III and IV(19). Herbert and independent colleagues established that vegetarians can withstand long-term insufficient intake of vitamin B$_{12}$ due to up-regulation of enterohepatic circulation and intestinal reabsorption of traces of vitamin B$_{12}$(19–22). In healthy vegetarians, this represents a mechanism to optimise the recycling of the scarce micronutrient.

One strength of the study by Naik et al. is that the level of accuracy for identifying vitamin B$_{12}$ deficiency using Hcy as a metabolic biomarker did not depend on the concentration of vitamin B$_{12}$ chosen (see fig. 2, ROC, tHcy in Naik et al.(16)). Using the current standard cut-off of 150 pmol/l for plasma vitamin B$_{12}$, the sensitivity was 91·8% and the specificity 79·31%. Applying the new proposed cut-off value of 100 pmol/l for plasma vitamin B$_{12}$, the sensitivity was 82·72% and the specificity 89·47%(16).

An important consideration that emerges from this analysis concerns the origin of reference intervals used worldwide to diagnose vitamin B$_{12}$ deficiency. Reference ranges have been established by examining plasma vitamin B$_{12}$, Hcy, methylmalonic acid and holo-TC, in healthy individuals residing in Western industrialised nations of whom the vast majority (>95%) pursue an omnivorous lifestyle. An omnivorous diet provides sufficient vitamin B$_{12}$ to keep tHcy and MMA at a minimum, and holo-TC above the established cut-off for deficiency (<35 µg/ml). Are these reference ranges established in omnivores appropriate cut-offs to assess vitamin B$_{12}$ status in populations that adhere to vegetarian diets? More generally, what are the optimal intracellular concentrations of Cbl, Hcy and MMA required to support function? And how well do serum levels of Cbl, Hcy and MMA reflect cellular cobalamin status?

What follows is the question of whether slightly elevated tHcy and MMA as seen in asymptomatic vegetarians represent a prelude to clinical deficiency of vitamin B$_{12}$ or if instead, these are metabolically satisfactory levels of metabolites that will cause no harm in the long term, that is subclinical cobalamin deficiency. According to Carmel, subclinical cobalamin deficiency is a condition where mild biochemical changes are documented (elevated tHcy and MMA, low vitamin B$_{12}$ and holo-TC) but the patient is asymptomatic(20). According to this definition, asymptomatic individuals presenting with elevated tHcy and low vitamin B$_{12}$ and holo-TC in the study by Naik et al.(16) would classify as having subclinical cobalamin deficiency(20). Carmel described that subclinical deficiency of
Thionine rarely progresses into clinical deficiency, which brings us to the next issue: should vegetarian individuals with subclinical cobalamin deficiency receive treatment with cobalamin? If so, what should be the dose and form of administration? A possibility exists that what classifies as subclinical cobalamin deficiency in omnivores may not represent a status of abnormal vitamin B12 homeostasis in plant-based individuals. Conceivably, applying reference values and cut-offs established from studying omnivorous populations may lead to the over-diagnosis of vitamin B12 deficiency in vegetarians and vegans. More broadly, the meaning of elevated tHcy and the impact of hyperhomocysteinemia on health remain a debate. The finding that patients with classical homocystinuria (deficiency of the enzyme cystathionine β-synthase) sustain good health by staying at a target level of tHcy under 120 μmol/L, that is well beyond the accepted normal range, questions the suitability of the tHcy reference values in plasma whereby 15 μmol/L is defined as the upper limit. Vegetarian populations who naturally consume lower amounts of vitamin B12 may exhibit tHcy concentrations higher than 15 μmol/L, without it representing a health threat.

Should a new reference interval for assessing vitamin B12 status be considered when examining human populations that pursue a predominantly plant-based lifestyle as proposed by Naik et al.? In light of current knowledge this proposal is a reasonable one, yet the definite answer to this question demands further study in larger cohorts of humans who have adopted vegetarianism both short and long term. Further, a mathematical expression that combines two, three or four biomarkers of vitamin B12 status, namely the cB12 index, has been shown to be a more reliable indicator of vitamin B12 status. It would be valuable to examine the performance of the cB12 index with and without the introduction of new cut-off points for the accurate and timely diagnosis of vitamin B12 deficiency in vegetarians.

Another consideration that emerges from this analysis is what supplemental dose of cobalamin should be recommended to support good health in vegetarians? The most recent recommendation for strict vegetarians is a small oral dose of 2–6 μg daily. Higher doses were only recommended if absorption problems are confirmed in individual cases. Further, patients who recovered from a clinical cobalamin deficiency and have no absorption problems can be maintained safely with low-dose daily supplements in the range of 5–10 μg of cobalamin. Therapeutic doses of cobalamin (1 mg daily doses, 4 weeks) are recommended to individuals with subclinical cobalamin deficiency only when the patient exhibits sufficiently suspicious clinical findings.

Special populations, whether omnivore or vegetarian, where vitamin B12 demands may not be met satisfactorily should be considered with caution. This includes the elderly and women in reproductive age attempting to conceive or who are pregnant. Elders with inefficient absorption of vitamin B12, a natural condition that together with lower dietary intake may increase the risk of acquiring a deficiency in vegetarians more so than in omnivores. In the case of women attempting to conceive or already pregnant, it must be remembered that depletion of serum vitamin B12 occurs much later than cellular depletion and elevation of metabolites in serum, making serum vitamin B12 a poor standalone marker of cobalamin status. Therefore, a conceiving woman with depleted cellular vitamin B12 may seriously compromise embryogenesis and early development, as the fetus relies entirely on the maternal supply of vitamin B12 via the placenta. It is highly recommended that vegetarian and vegan women in reproductive age take a vitamin B12 supplement before conception as well as during pregnancy and breast-feeding to ensure sufficient supplies of the micronutrient to the baby.

With the worldwide increase of humans invoking a vegetarian lifestyle, a demand exists to address questions concerning the diagnosis, interpretation and management of vitamin B12 insufficiency in these populations. The study by Naik et al. sets the stage for further large-population studies both at the epidemiological and basic research fronts to reassess the criteria for diagnosing vitamin B12 deficiency in vegetarian populations. Careful examination of these criteria will allow clinicians to define the appropriate chemical form, mode of administration and dose of cobalamin supplementation required to prevent the onset of acute vitamin B12 deficiency, as well as the timely implementation of therapy.

Acknowledgements
The author is extremely grateful to Dr Donald W. Jacobsen for critical reading of this manuscript.

The author declares that there are no conflicts of interest.

Luciana Hannibal
Laboratory of Clinical Biochemistry and Metabolism, Department for Pediatrics, Medical Center, University of Freiburg, Mathildenhstr. 1, D-79106 Freiburg, Germany
email luciana.hannibal@uniklinik-freiburg.de
doi:10.1017/S000711451800048X

References