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Functional vitamin B12 markers were compared with serum vitamin B12 levels in 350 individuals, to see if there was a correlation between markers of vitamin B12 deficiency, and the actual levels of vitamin B12 in serum. No correlation was found between normal to elevated levels of vitamin B12 and urinary methyl malonic acid, or vitamin B12 deficiency associated neurometabolites. Hence, whilst serum vitamin B12 levels were normal or elevated these individuals had Paradoxical B12 deficiency. Vitamin B12 deficiency markers were, however, correlated with markers of vitamin B12 deficiency, suggesting that the vitamin B2 deficiency had led to elevated levels of vitamin B12, which was largely inactive. Given the role in energy metabolism, correct diagnosis of vitamin B12 deficiency is essential. The mechanism is discussed.

Keywords: Paradoxical vitamin B12 • Organic acids test • Vitamin B2 • Creatine

Introduction

The current dogma on vitamin B12 deficiency implies a direct link between levels of serum vitamin B12, and sufficiency. Hence as vitamin B12 levels reduce, markers such as MMA and homocysteine increase in an inverse relationship to serum B12 levels [1-4]. A consequence of this “Dogma” is that if a person does not have vitamin B12 levels lower than a highly variable amount, dependent upon study (148 pmol/L, 179 pg/ml, 230 pmol/L), they cannot be B12 deficient [1,5,6]. Hence vitamin B12 deficiency would be mainly due to poor diet, or poor absorption or lack of Intrinsic Factor [7]? Many people, however, experience symptoms of vitamin B12 deficiency yet their serum levels of vitamin B12 may be normal or much higher than normal. Subsequent examination of biochemical markers such as MMA or homocysteine may show that these markers are moderate to highly elevate. As such the symptoms and biochemical markers are indicative of vitamin B12 deficiency, yet the serum vitamin B12 levels are paradoxically high. Such persons are deemed to have “Paradoxical Vitamin B12 Deficiency”. Generally, however, the reason(s) for “Paradoxical B12 Deficiency” are not known, even despite its association with a greater increase in “all-cause mortality”, chronic viral liver disease, Anorexia Nervosa, and death from COVID-19 [8-10].

It has been known for over 40 years that measurements for serum vitamin B12 levels can be greatly distorted by the presence of non-functional cobalamin analogues, and in many cases measurement of serum B12 levels is not predictive of the levels of functionally active B12 [11-20]. In addition, low intracellular folate levels can often be associated with the presence of inactive analogues of cobalamin in serum [21]. Alternatively, it can be associated with inflammatory conditions such as Rheumatoid arthritis due to the overproduction of the B12 binding proteins Tran’s cobalamin and haptocorrin, Cystic fibrosis, and patients treated with nitrous oxide poisoning [22-28].

More recently elevated B12 levels have been associated with a poorer prognosis following treatment for cancer [26-29]. The higher the B12, the poorer the prognosis. We have not been able to find any study where the authors have “looked at” metabolic markers that one would ascribe to functional B2 deficiency, and compared it to the elevated B12 levels in these cancer studies. Studies on gastric cancer have suggested that the cancers themselves over-produce haptocorrin (R Binder), and hence elevated serum Hc-B12 may be indicative of prognosis [30-34]. Elevations of serum B12 have also been found in cats with neoplasms [35]. Extreme levels of Hc-B12 (>18,000 pg/m B12) have been found in some metastatic cancers [36].

Despite the almost completely non-predictive nature of serum vitamin B12 levels it is still the measurement of choice for vitamin B12 sufficiency. Its utility is arguably restricted to determining conditions such as dietary insufficiency, poor absorption or Pernicious Anaemia, where serum levels are routinely low, and as such many physicians miss functional vitamin B12 deficiency in patients.

In previous studies we have shown that functional vitamin B12 deficiency is often associated with functional vitamin B2 deficiency. In the current study, we have compared the levels of serum vitamin B12, with various biochemical markers associated with vitamin B12 deficiency, and have compared the later markers with markers of functional vitamin B2 deficiency.

Materials and Methods

Study sample

A retrospective analysis was performed upon data submitted to us for interpretation from a cohort of 350 children and adults who had submitted their Organic Acids Test data (Great Plains Laboratories, Lenexa, KS, USA) for analysis and who had also supplied their serum vitamin B12 levels. Subjects were from a range of countries including USA, Canada, United Kingdom, Ireland, Germany, Spain, France, Italy, Bulgaria, India, Sweden, Bulgaria, Serbia, Dubai, Croatia and Australia. Age distribution of the 350 individuals was 1-5 (30); 6-10 (30); 11-15 (40); 16-20 (19); >21 (216). Data was collected over a period of 5 years. No selection was made in the acceptance of data, with no data being rejected. Individual data is plotted as Scattergrams (see Graphs 1 to 7). Data is represented as mmol/mol creatine for NT.

Results

Absolute vitamin B12 levels in serum were compared to Methyl Malonic Acid (MMA) levels measured in urinary Organic Acids (OAT). Representation of the data in a scatter gram (Figure 1) showed that there was no obvious relationship between levels of vitamin B12 in serum, and the standard marker of vitamin B12 deficiency, MMA (Methyl Malonic Acid). This is in contrast to
the comparison of absolute vitamin B12 deficiency in serum, where MMA starts to increase as vitamin B12 drops below 221 pmol/L (Figure 1) [6, 37].

Previous studies by us have shown that deficiency of active vitamin B2 is correlated with the vitamin B12 deficiency markers HVA, VMA, QA, KA and 5HIAA. The level of MMA was compared to the vitamin B2 deficiency marker, glutaric acid (Figure 2), as too were the markers, HVA, VMA, QA, KA and 5HIAA (Figures 3-6).

As can be seen there was good correlation between both MMA and glutaric acid with VMA, HVA and QA, however, the comparison was less strong for KA. This presumably is because in order to make the metabolite KA, there is a Pyridoxal-Phosphate-dependent enzyme (kynurenine-aminotransferase) step. Formation of the active form of vitamin B6, pyridoxal phosphate (P5P), requires FMN, hence in low Iodine, or Selenium the amount of FMN would be reduced, as too P5P, and so too the amount of KA (Figure 7).

![Figure 2. Comparison of urinary MMA with glutaric acid.](image2.png)

![Figure 3. Comparison of urinary HVA with MMA (left panel) with glutaric acid (right panel).](image3.png)

![Figure 4. Comparison of urinary VMA with MMA (left panel) with glutaric acid (right panel).](image4.png)

![Figure 5. Comparison of urinary KA with MMA (left panel) with glutaric acid (right panel).](image5.png)
Formation of SHiAA was less well correlated with either MMA or glutaric acid, however, it is interesting to note that SHiAA levels tended to increase when glutaric acid levels were lower, a situation that occurs when functional B2 activity is reduced and so presumably the activity of the FAD-dependent enzyme, monoamine oxidase, which is responsible for the inactivation of serotonin to SHiAA, would be higher. In extreme B2 deficiency, MAO would have reduced activity and so serotonin break-down would be reduced.

**Discussion**

In a previous study we compared compared markers of functional vitamin B2 deficiency with those of functional vitamin B12 deficiency, and found a correlation supporting the hypothesis that lack of functional vitamin B2 eventually causes a lack of functional vitamin B12. In this study we have attempted to see if levels of serum vitamin B12 are predictive of functional vitamin B12 levels, and found that they are not at all predictive of functional B12, however, there was a strong correlation between MMA, a marker for Adenosyl B12 deficiency and VMA/HVA/QA (markers of methyl B12 deficiency) and MMA and glutaric acid (a marker of functional B2 deficiency).

The data supports the hypothesis that in functional B2 deficiency, the activity of the two vitamin B12 house-keeper enzymes methylenetetrahydrofolate-reductase (MTHFR-FAD/NAD dependent enzyme) and methionine synthase reductase (MTRR–FMN/FAD/NAD dependent enzyme), is greatly reduced and so the repeated cycling of methylCo(III)B12 ⇌ Co(I)B12 is greatly compromised with the result that Co(I)B12 is rapidly oxidized to inactive Co(II)B12, which cannot be rescued by MTRR in the reaction Co(II)B12+SAM ≥ MethylCo(III)B12+SAH. This inactive cobalamin is then secreted from the cell and accumulates in serum, thereby raising the apparent level of vitamin B12 in serum and yet the B12 is inactive. Subjects with these high levels of inactive B12, therefore have “Paradoxical vitamin B12 deficiency”. This functional B12 deficiency cannot be resolved by additional vitamin B12 until the functional vitamin B2 deficiency is resolved, and the activities of MTHFR and MTRR are restored. The major cause of Paradoxical Vitamin B12 Deficiency appears to lack of functional vitamin B2, which may occur due to overt vitamin B2 deficiency in a person’s diet, Hypothyroidism [38], or due to lack of adequate intake of Iodine, Selenium and or Molybdenum, which in turn leads to insufficient production of the two active forms of vitamin B2, namely FMN and FAD. FMN and FAD both have critical roles in cycling and maintenance of activity of vitamin B12, particularly methyl B12.

Methyl-Co (III) B12 has a major role in the body in the removal of homocysteine, and in the regeneration of methionine in the methylation cycling using the enzyme methionine synthase reductase (MTR).

Functional Methyl-Co (III) B12 is essential for the formation of creatine, the cytoplasmic energy transfer shuttle; hence lack of functional Methyl-Co (III) B12 has important implications for organs with high energy demand, such as the heart, muscles and nervous system. Potentially this explains the observation that “Paradoxical B12 deficiency” is common in conditions such as Chronic Fatigue Syndrome (CFS) and is almost universal in Autism Spectrum Disorder, and in dementia.

**Conclusion**

Dietary deficiency of Iodine, Selenium and/or Molybdenum can lead to an inability to activate dietary or supplemental vitamin B2. This in turn leads to an inability to maintain the functional activity of vitamin B12. Prolonged lack of active vitamin B2 eventually leads to an accumulation of inactive vitamin B12 in serum, with a gradual increase in the absolute amount of serum B12. Thus, elevated vitamin B12 is found in serum, yet the B12 is largely inactive, leading to a condition of Paradoxical vitamin B12 deficiency.

**Compliance with Ethical Standards**

We declare that there are no potential conflicts of interest.

Research did not involve human participants and/or animals.

No Informed consent is required, all data is “blinded” and as such is anonymous.

Data analysis was carried out under the Australian National Health and Medical Research Council guidelines (NHMRC). Under these guidelines, all data was deidentified and steps were taken to ensure the anonymity and confidentiality of the data. Deidentification has consisted of absolute anonymity and confidentiality of the data, such that no specifics such as gender, ethnicity, Country of Origin, etc. are associated with any data point in the study.

As such per the NHMRC guidelines:

- The research does not carry any risk to the participants
- The benefits of the research are many and will be of considerable benefit to any past, current or future participants, and as such represent no harm.
- The data is from over 600 participants collected over 6 years and as such it would be impracticable to obtain consent from the participants. Further the participants had been notified at the time of analysis that data presented for analysis might potentially be used in research – but would be totally de-identified (which it has been).
References


