Impact of type 2 Diabetes and Metformin use on Vitamin B12 Associated Biomarkers - an Observational Study

Authors
Corina Metaxas¹, Chantal Zurwerra¹, Gottfried Rudofsky², Kurt E. Hersberger¹, Philipp N. Walter², 3

Affiliations
1 Pharmaceutical Care Research Group, University of Basel, Switzerland
2 Department of Medicine, Kantonsspital Olten, Switzerland
3 Institute of Laboratory Medicine, Solothurn Hospitals, Switzerland

Key words
T2DM, metformin, vitamin B12 deficiency, vitamin B12, holotranscobalamin

ABSTRACT

Aims Assessment of the impact of type 2 diabetes (T2DM) and metformin use on vitamin B12 (VB12) associated biomarkers and their suitability to represent VB12 supply.

Methods Differences of VB12, holotranscobalamine (HoloTc), the biologically active fraction (%AB12) = HoloTc/VB12 * 100 and homocystein (Hcy) were analysed i) among diabetic outpatients with (DMMet+) and without metformin use (DMMet-) and ii) in comparison to an external non-diabetic reference group with low VB12 (< 200 pmol/L).

Results VB12 associated biomarkers were distributed equally between DMMet+ (n = 29, 58 %) and DMMet- (n = 21, 42 %). Significant differences in %AB12 in diabetic patients with low VB12 (n = 19) compared to the non-diabetic reference group (n = 31) were found. Higher %AB12 was associated with diabetes. Hcy levels were significantly associated with age, folic acid level, renal function and HoloTc but not with VB12.

Conclusions In T2DM patients with low VB12, %AB12 was confirmed as being higher in comparison to nondiabetic patients. The effect was not clearly attributable to metformin use. HoloTc was unaffected by the lowering of VB12 and significantly associated with the functional marker Hcy. Both findings support the use of HoloTc for the assessment of VB12 supply in diabetic patients.

Introduction

Recently, an increased frequency of VB12 deficiency among T2DM patients has been documented by several cross sectional studies and case reports [1]. Clinically, VB12 deficiency in adults may result in nonspecific symptoms such as tiredness, loss of appetite, hematologic manifestations (megaloblastic anemia), neurologic symptoms (e.g., polynuropathy, ataxia), as well as symptoms of a psychiatric nature (e.g., depression) [2, 3]. Additionally, cardiovascular manifestations associated with hyperhomocysteinemia were mentioned [4–7].

Clinical symptoms of VB12 deficiency in T2DM are comparable to those in the general population. Worsening of diabetic neuropathy has been described among patients with co-existing vitamin B12 deficiency. Furthermore, VB12 replacement has been shown to cause symptomatic improvement, reduction in pain, and paresthesia among patients with severe diabetic neuropathy [1], suggesting that functional VB12 deficiency in T2DM patients is clinically significant. Sensory polyneuropathy caused by VB12 deficiency mimics diabetic neuropathy [8]. Overlapping symptoms between T2DM and VB12 deficiency may complicate its clinical suspicion (or diagnosis). Biochemically, VB12 deficiency is charac-
characterized by subnormal to borderline VB12 values in serum (<148–221 pmol/L) [9]. Holotranscobalamin (HoloTc) is the bioactive form of VB12 and makes up to 20% of the total vitamin B12 concentration in the human body [10]. It has been discussed as a more specific and sensitive marker of VB12 deficiency [11–13]. Broad ranges of cut-off points for HoloTc have been described as <20–50 pmol/L [14]. Functional VB12 deficiency is characterized by elevated homocysteine (Hcy) and/or methylmalonic acid (MMA) levels [9].

Additionally, the clinical biochemistry of VB12 is influenced by diabetes and its treatment. Systematic observations in clinical trials as well as biochemical studies in animals raised questions on possible interactions between diabetes, metformin treatment, and VB12 metabolism. It has been proposed that the increased oxidative stress in diabetes is involved in the pathogenesis of functional VB12 deficiency [15]. Treatment of T2DM patients with metformin has been reported with reductions of 10–20% in plasma VB12 levels [16–20]. Metformin may impair VB12 absorption and thereby induce VB12 deficiency [20, 21]. One study described the correlation of cumulative metformin dose, low VB12 levels, and clinically more severe peripheral neuropathy [8]. Given the widespread use of metformin as first line treatment for patients with diabetes and normal kidney function, its effect on VB12 metabolism is remarkable [22]. Metformin is a biguanide. Its mechanism of action primarily involves decreasing hepatic glucose production and increasing glucose uptake [23]. Because VB12 deficiency is a reversible cause of demyelinating nervous system disease and bone marrow failure, its early detection and treatment are important [3]. Therefore, the identification of metformin as a risk factor for the development of VB12 deficiency is important and the evaluation of VB12 status is recommended [24]. However, there is reasonable doubt whether the reduction of VB12 levels reflects a true decrease in VB12 supply. A recent study in patients with diabetes (>65 years old) which were treated for 3 months with metformin showed that the inactive part of VB12 bound to haptocorrin is reduced in metformin-treated patients but not in the control group (non-significantly) [25]. Additionally, a study in rats found that metformin treatment increases liver accumulation of VB12 thereby resulting in decreased circulating VB12 and kidney accumulated VB12 [26]. The same authors found a significant reduction of serum VB12 but not HoloTc after 6 months of metformin treatment in women with polycystic ovarian syndrome (PCOS) [27]. These findings raise the question of whether low serum VB12 observed in patients treated with metformin actually reflects VB12 deficiency and rather supports the hypothesis of altered metabolism of VB12 in metformin-treated patients.

In this study, the effect of metformin treatment on VB12 status as reflected by total VB12 and HoloTc in T2DM patients is investigated. In particular, the %AB12 in T2DM patients with and without metformin treatment was compared and interactions between VB12, HoloTc, and homocysteine were analysed. Additionally, the impact of diabetes itself on VB12 associated biomarkers at low VB12 levels is investigated in a subgroup of patients with diabetes in comparison to an external reference group of non-diabetic patients.

Materials and Methods

This was an observational cross-sectional trial approved by the ethics committee of northwestern Switzerland (EKNZ), and was registered at ClinicalTrials.gov NCT02111967. The study was conducted in accordance with the Declaration of Helsinki and corresponding to International Conference on Harmonization Good Clinical Practice (ICH-GCP) guidelines. All patients gave written informed consent. Patients for the external reference group were included from a previous study (ClinicalTrials.gov NCT01832129). The primary hypothesis was that VB12 would be lower in T2DM patients treated with metformin while HoloTc levels would not differ compared to T2DM patients without metformin treatment.

Setting

The study included adults with type 2 diabetes and an external reference group of nondiabetic and metformin-naïve patients. Patients with T2DM were classified as metformin users (DMMet+) if they were treated with metformin for at least 6 months or as metformin-naive patients (DMMet-), when no prior use of metformin was reported (▶Fig. 1). A. Participants with newly initiated metformin (<6 months usage), concurrent intake of preparations containing VB12 (in the last 3 months), diagnosis of transcobalamin transporter defect, diagnosis of vitamin B6 deficiency, diagnosis of liver disease with CHILD-PUGH scores B and C, acute hepatitis, and diagnosis of alcohol abuse (defined as a diagnosed condition or by self-reported daily intake of alcohol), diagnosis of renal insufficiency station III, IV, and V (KDOQI), acute renal diseases or renal function below 60 ml/min according to the Cockroft–Gault equation were excluded from the study. For the reference group, data from patients from a previous study were used [28]. Out of 37 patients, 4 were excluded due to an alcohol use disorder (n = 4) (▶Fig. 1).

Patient recruiting

Patients with diabetes with (DMMet+) and without metformin treatment (DMMet-) were recruited during a routine visit at their diabetes specialist (▶Fig. 1; sample A). Patients with diabetes and serum VB12 levels below 200 pmol/L were classified as VB12 deficient (LVB12-DM). The external reference group of non-diabetic patients with low VB12 levels (LVB12-Ref) was established from a previously published study that recruited outpatients during a routine visit to their general practitioner. If patients had a diagnosis of diabetes, they were not excluded but assigned to the LVB12-DM Group (▶Fig. 1, sample B).

Questionnaires and physical examinations

Patients were asked to fill in a questionnaire about their nutrition, co-medication, and demographics. Patients with type 2 diabetes were screened for neuropathy signs using the Neuropathy Symptom Score (NSS) and the Neuropathy Disability Score (NDS) [29].

Pharmacological biomarker

All venous blood samples were analysed for VB12 (Beckman Coulter® Dxc 860i), HoloTc (Abbott Architect® i2000SR), and Hcy (Roche Cobas® 6000). The biologically active fraction (%AB12) was calculated by dividing HoloTc/VB12 * 100. Blood cell count was de-
determined on a Beckman Coulter DxH 800. Cut-off values defining VB12 deficiency were VB12 < 200 pmol/L, HoloTc < 37 pmol/L and Hcy > 15 µmol/L. Blood samples of patients with VB12 levels below 200 pmol/L were re-analyzed for VB12 on the Roche Cobas® 6000 to directly compare the results with the external reference group.

Study size

Based on published data, patients with diabetes were assumed to display VB12 levels of approximately 400 pmol/l, with an estimated standard deviation of 250 pmol/l (62.5 %) [30]. In this study, 40.1 % of patients had consumed VB12 supplements; therefore, we concluded that the standard deviation among patients without VB12 supplements was approximately half of the observed standard deviation (30 %). A reduction in plasma VB12 levels of 20–24 % is described in the scientific literature [25]. Since we expected a reduction in levels, a one-tailed test was applicable. Based on a reduction of VB12 levels of 24 %, a total of 50 samples had had to be analysed to ensure that the 80 % confidence interval includes the true difference of means with a α-significance level of 5 %.

Statistical analysis

Values are given as medians, with quartiles and percentages where appropriate. Frequencies were analyzed using Chi-square tests or Fisher’s test. The Mann-Whitney test was used to compare numerical variables between two groups. Bivariate correlation was performed when association between two scaled variables was assessed. A p-value ≤ 0.05 was considered significant. Multiple linear regression analyses were conducted to calculate i) the relative impact of VB12 and HoloTc on Hcy in relation to age, renal function assessed as estimated Glomerular filtration rate (eGFR) according to the Cockcroft-Gault equation, and folic acid levels; and ii) the relative impact of diabetes, age, BMI, renal function and folic acid levels on the biologically active fraction %AB12 = HoloTc/VB12 * 100 using a stepwise method to include the independent variables. The statistical procedures were performed with SPSS statistical software Version 24 (SPSS Inc., Chicago, Il, USA).

Results

Participants

Between March 2014 and August 2014, 50 outpatients with diabetes were recruited within the endocrinology unit. Recruitment and patient characteristics of the external reference group have been described elsewhere [28]. Two of the 37 recruited subjects in this study had diabetes and were thus integrated in the respective subgroup for the analysis of VB12 associated biomarkers in diabetes (▶ Fig. 1). Patient characteristics of T2DM patients (29 DMMet+ ; 21 DMMet-) were equally distributed (▶ Table 1).

VB12 associated biomarker levels in diabetes patients (sample A)

DMMet+ versus DMMet-

VB12 serum levels and HoloTc levels were slightly lower in DMMet+ patients compared to DMMet-, but not significantly (VB12: − 10.4 %; HoloTc: − 9.2 % ▶ Table 2). The %AB12, Hcy and blood count levels were equally distributed between the two groups (▶ Table 2). Out of 50 patients, 17 (34 %) had a VB12 value below 200 pmol/L (11 DMMet+ (39.3 %) and 6 DMMet- (28.6 %); Fishers-test: 0.55), 3 (6 %) had a HoloTc value below 37 pmol/L (3 DMMet+ (10.3 %) and 0 DMMet- (0 %); Fishers-test: 0.25) and 16 (32 %) had a Hcy level above 15 µmol/L (10 DMMet+ (34.5 %) and 6 DMMet- (28.6 %); Fishers-test: 0.75) (data not shown).

Correlations of VB12 associated biomarkers

Significant correlations (p < 0.05) of VB12 levels with age (r = − 0.42) and Hcy (r = − 0.37) were found for the entire study population, as well as with metformin dose (r = − 0.43), and renal function (r = − 0.41) in DMMet+ patients. For HoloTc, significant correlations

▶ Fig. 1 Sample A) Patients recruited by the diabetologist. Comparison of VB12, HoloTc and %AB12 between DMMet+ and DMMet- and assessment of associations between VB12, HoloTc, Hcy and severity of neuropathy. Sample B) Patients with low VB12 (< 200 pmol/L) recruited by diabetologist and general practitioners. Comparison of VB12, HoloTc and %AB12 between LVB12-DM and LVB12-Ref.
were limited to Hcy ($r = -0.44$) and metformin dose ($r = -0.47$) in DMMet+ patients. High Hcy levels were associated with high age ($r = 0.35$), low renal function ($r = -0.48$), and low folic acid levels ($r = -0.36$) in the entire study population. Mean daily metformin dose in DMMet+ group was $1.9 g \pm 0.6 g$ (range: $1.0 g–3.0 g$).

### Prediction of Hcy levels

In a multiple linear regression model, a significant association of Hcy with age ($β = 0.325$; $p = 0.029$), folic acid ($β = -0.359$; $p = 0.003$), HoloTc levels ($β = -0.300$; $p = 0.01$), and renal function expressed as eGFR by Cockroft-Gault ($β = -0.343$; $p = 0.014$) was found in all DM patients ($r^2 = 0.539$; $p = 0.022$), while no significant association of VB12 with Hcy was observed.

### VB12 associated biomarker levels according to the severity of neuropathy

Neither VB12, HoloTc, nor Hcy differed significantly between patients with mild, moderate, and severe symptoms of neuropathy (Fig. 2). Age and diabetes duration were significantly higher in patients with severe neuropathy compared to patients with mild neuropathy.

### Discussion

In our study, neither VB12, HoloTc, or %AB12 differed between T2DM patients regardless of metformin treatment. Median VB12 and HoloTc levels were within the normal range in both groups. Therefore, the primary hypothesis that VB12 levels would differ while HoloTc levels would be indifferent between T2DM patients with or without metformin treatment had to be rejected. Metformin treatment alone did not explain the altered VB12 metabolism as reflected by VB12 and HoloTc serum levels in all T2DM patients, as suggested by the literature [26, 27]. Nevertheless, the proportion of patients in the DMMet+ group with low VB12, low HoloTc, or high Hcy was higher compared to DMMet- group (not significant). Additionally, metformin dosage did negatively correlate with VB12 and HoloTc levels. These findings suggest that metformin may contribute to VB12 deficiency.

Further analysis focused on VB12-deficient subgroups and included non-diabetic patients. In this sample (B), a significant difference of the %AB12 was observed and confirmed by multiple regression analysis. However, the model explained only 9.2% of the variance observed. These results suggest that VB12 metabolism is affected by diabetes itself, as well as by other factors which were not included in the model. It has been proposed that duration of VB12 deficiency and causes of VB12 deficiency play a major role in the VB12/HoloTc ratio [31], such as liver diseases [32], in particular alcoholism [33]. In this study, patients with liver diseases and/or alcoholism were excluded. Duration of a VB12 deficiency can hardly be assessed in an observational study; therefore, interventional studies might be favorable to continue research on VB12 metabolism. Multiple regression analysis showed that T2DM in general had an impact on %AB12. Further observations regarding the impact of metformin in T2DM patients with low VB12 had a high variance (data not shown) due to the relatively small sample (17 T2DM patients; 11 DMMet+ and 6 DMMet-) and therefore were not taken into account. Thus, larger studies may be necessary to differ whether the altered VB12 metabolism is attributable to diabetes in general, or specifically to metformin use.
HoloTc has been proposed as a better marker to detect VB12 deficiency compared to serum VB12 [34, 35] in an aged population [36]. We found significant inverse correlations of VB12 and HoloTc with Hcy, a functional marker of VB12 deficiency. Although the effect was stronger between VB12 and Hcy compared to HoloTc and Hcy, stepwise multiple regression analysis included HoloTc as independent variable to explain variance in Hcy levels and not VB12. Thus, HoloTc seems favorable compared to VB12 to predict hyperhomocysteinemia caused by VB12 deficiency in T2DM patients. Therefore, our results support the finding that HoloTc might be a better marker than VB12 to detect VB12 deficiency. Furthermore, regression analysis showed that in our sample, elevated Hcy was also explained by age, folate deficiency or renal insufficiency, thus compromising its value as an independent reference for VB12 deficiency. Other studies found inverse correlations between GFR and Hcy within patients with normal renal function also [37, 38]. Metabolism of methylmalonic acid, another functional biomarker of VB12 deficiency, is not affected by vitamin B6 or folic acid. Therefore, measurement of methylmalonic acid might be more sensitive in detecting VB12 deficiency than Hcy. However, methylmalonic acid levels may be compromised in patients with reduced GFR, too [39].

Results from a previous study questioned whether low VB12 in metformin-treated patients with diabetes causes a true or functionally irrelevant cobalamin deficiency [27]. Interestingly, in our study, significant inverse correlations of Hcy with VB12 and HoloTc were exclusively found in DMMet+ patients. Even though VB12 levels were comparable within all DM patients, low VB12 and HoloTc levels did cause functional deficiency in DMMet+ but not in DMMet-. An explanation for this finding might be that no true VB12 deficient DMMet- patients were included in our study, supported by the fact that no HoloTc below 37 pmol/L was found in DMMet-group. Furthermore, our data suggest that metformin induces
VB12 deficiency in a dose-dependent manner. Other studies also found associations between metformin dose and VB12 levels [37, 40]. Thus, it is reasonable to screen patients treated with metformin for VB12 deficiency, as proposed [41]. In a future study, inclusion of anemic patients might help to clarify whether the elevation of functional markers results from VB12 deficiency or from impaired GFR, and whether low VB12 and/or HoloTc coincide with anemia.

VB12, HoloTc or Hcy levels did not differ in patients with mild, moderate or severe symptoms of neuropathy assessed with the NSS and NDS, while, well-established risk factors such as age and duration of diabetes differed between groups. While some studies showed associations between neuropathy, VB12 associated biomarkers, and metformin use [8, 42], others failed to find such correlations [43–45], making results controversial, overall. Controversy might exist because of a range of different assessments for neuropathy and differences in study designs exist. The observed incidence of a VB12 deficiency in this study was high (34 %). Irreversible neurologic damage caused by VB12 deficiency is preventable through treatment [3, 8], at relatively low costs, and has few side effects. Therefore, screening for VB12 deficiency independently from diabetic neuropathy seems reasonable.

We acknowledge some limitations. First, our sample size calculation was based on bigger difference in VB12 levels between T2DM patients with and without metformin. However, the prevalence of patients having VB12 deficiency defined as VB12 below 200 pmol/L was based on bigger difference in VB12 levels between T2DM patients with and without metformin. However, the prevalence of patients having VB12 deficiency defined as VB12 below 200 pmol/L (34 % (DMMet+ (39.3 %) and 6 DMMet- (28.6 %)) we observed is comparable to what is described for elderly users [46, 47]. Therefore, differences in VB12 associated biomarkers may also be observed within all DM patients when more patients were included. Second, the assessment if HoloTc or VB12 might be better to detect VB12 deficiency was only based on a laboratory marker (Hcy levels). In this study, non-anemic patients were included and therefore clinical symptoms where not taken into account. Further studies should include anemic patients and assess methylmalonic acid.

Conclusion

The clinical biochemistry of VB12 in T2DM patients with scarce VB12 supply is modified in comparison to nondiabetic patients. This results in higher %AB12 due to reduced VB12 levels. It needs to be clarified whether this effect is due to diabetes itself, metformin in treatment, and/or a combination of other health related situations. Assessment of HoloTc seems favorable compared to VB12 to predict hyperhomocysteinemia caused by VB12 deficiency in T2DM patients. This may be a direct consequence of the modified %AB12 in T2DM patients, which strengthens the recommendation to assess VB12 supply in clinical practice by measuring HoloTc. VB12, HoloTc and Hcy did not differ in patients with mild, moderate, or severe symptoms of neuropathy.

Acknowledgements

We would like to thank all participating patients and all participating general practitioners and their staff for patient recruitment. We thank Ursula Zoli (Aarelab, Olten) and the laboratory technicians at the Institute of Laboratory Medicine in Olten for their practical contribution. We thank William Caddy for proofreading the manuscript.

Other information: No grants from any external funding body were received to conduct this study.

Conflict of Interest

The authors declare no conflicts of interest.

References

[9] Schrier SL. Diagnosis and treatment of vitamin B 12 and folate deficiency. update (online, last update 09/2016) 2016;