Short Report

Functional predictors of treatment induced diabetic neuropathy (TIND): a prospective pilot study using clinical and neurophysiological functional tests

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Abstract
Background: A treatment-induced drop in HbA1c has been suggested to be a risk factor for TIND.
Methods: From 60 included patients with severe diabetes mellitus (HbA1c over 8.5) only 21 patients adhered to the study protocol over 1 year with a battery of autonomic nervous system tests scheduled before and after starting antidiabetic treatment.
Results: In patients with a drop of HbA1c greater than 2 per cent points only some neurophysiologic tests and lab values tended to deteriorate with a trend to improve at later time points along the study. None of these changes were statistically significant, most likely because the study failed to reach the planned number of patients.
Conclusion: Poor adherence to diabetes treatment and to following the study protocol were the assumed obstacles in our patient cohort selected for very high HbA1c levels. In future studies a multi-center trial and case numbers of up to 500 patients may be needed to account for drop outs in the range observed here. Moreover, the number of tests in each patient at each visit may have to be reduced and special educational group sessions are warranted to cope with the limited adherence.

Keywords: Treatment-induced neuropathy of diabetes (TIND), Diabetes mellitus, Heart rate variability, 30:15 ratio, Neuropathic pain, Autonomic neuropathies, HbA1c

Background
Treatment-induced neuropathy of diabetes (TIND) is a subacute type of diabetic neuropathy affecting small peripheral nerve fibers [1, 2]. TIND is characterized by acute neuropathic pain and autonomic dysfunction starting within 8 weeks of therapy initiation [3, 4]. Concomitant rapid decrease in HbA1c of more than 2 percent points over 3 months is usually seen [2, 5–7]. In a retrospective study with a 5-year observation period, Gibbons and Freeman [5] found that 11% of patients with diabetes developed TIND. The pathogenic relevance of a fast decline in HbA1c for the manifestation of TIND has been corroborated in a rodent diabetes model [8].

Originally, TIND was treated by reducing insulin doses tolerating a permissive hyperglycemic metabolic state [5, 9]. Reports of the efficacy of this treatment regimen were rarely documented [5]. Yet, maintaining chronic hyperglycemia may in itself augment the well-known long-term complications [10]. The present treatment
recommendations including insulin and other anti-diabetic compounds aim at a slower and gradual decline of HbA1c levels [11].

Under appropriate treatment TIND is a self-limiting disorder lasting over weeks or several months [3, 9, 11]. Symptomatic treatment of severe neuropathic pain includes antiepileptic drugs like pregabalin, tricyclic antidepressants, and in cardiac autonomic neuropathy angiotensin-converting enzyme inhibitors or antiarrhythmics [5, 7].

One of the potentially life-threatening complications of TIND-associated autonomic neuropathy is arrhythmia and cardiac failure that may lead to increased mortality [12].

Prospective studies are needed to investigate potential predictive factors for developing TIND including associated autonomic neuropathy. We therefore, initiated a single center, prospective pilot study in patients with diabetes and baseline HbA1c levels above 8.5%. At baseline and after receiving adequate treatments over a period of 1 year, we aimed at detecting abnormalities in various tests of autonomic dysfunction that might predict the risk for TIND by utilizing non-invasive neurophysiological functional test procedures.

Patients and methods
Sixty patients (23 women, 37 men) were screened when diagnosed with diabetes mellitus type 1 or 2 and HbA1c values greater than 8.5%. Out of these, 21 patients (5 women, 16 men) agreed to be repeatedly examined over a period of one year. Clinical and neurophysiological examinations were planned for all patients at baseline (T0) and after 3 (T1), 6 (T2), and 12 months (T3). In addition to standard blood analyses including HbA1c we performed measurements of C-reactive protein (CRP) to detect inflammatory pathology. A battery of non-invasive neurophysiological function tests was conducted including the following tests procedures: cardiovascular autonomic reflex tests (30:15-ratio; Valsalva-ratio, E/I-ratio); quantitative sudomotor axon reflex tests (QSART) measuring sweat production after stimulation with acetylcholine [13]; collectively, the group of cardiovascular autonomic reflex and the QSART examine parasympathetic function while PDD and the other autonomic tests examine sympathetic function [2, 14]. Abnormalities in any of these procedures indicate small fiber neuropathy. The study protocol was approved by the Ethics Committee at our institution (No. 241-2009-0911209). All participants gave written informed consent. The differences of the HbA1c values and the differences of the neurophysiological tests between T0 and T1 were calculated and used for Pearson’s correlation analyses. Patients were grouped according to the treatment-related reduction of HbA1c. Group A consisted of patients whose HbA1c dropped by 2 percentage points or more and group B dropped by less than 2 percentage points. Comparisons between the groups and between points in time were analyzed using unpaired (groups) and paired (time) t-tests and Wilcoxon–Mann–Whitney tests [16]. SPSS version 16 was used for the statistical analyses. We expected deterioration in neurophysiological test results at T1 as compared to T0, with further deterioration during the subsequent course. All participants gave written informed consent.

Results
Of the screened 60 Patients, only twenty-one (35%) agreed to participate in the planned further test procedures. Of the 21 only 13 achieved an HbA1c value reduction of more than 2 percent points in the first 3 months. Overall adherence was 22%.

The 21 patients participating over 1 year received a full battery of tests while receiving effective anti-diabetic medication and dietary recommendations. In the 13 patients of group A, mean reduction in HbA1c from T0 at T1 was 4.8 percent points (p = 0.001). Of these, only one patient suffered from neuropathic pain at T1 which later regressed. In the 8 patients of group B mean reduction of HbA1c was only 0.13 percent points on average (p = 0.53) suggesting an insufficient choice of anti-diabetic treatment or, more likely, poor treatment adherence (Table 1).

While values of functional tests were quite similar at baseline (T0) in both groups, patients of group A tended to later display abnormal test results in the 30:15 ratio, E/I-ratio, and CPT at T1. Subsequently, these abnormalities regressed. In contrast, in group B the test results tended to gradually deteriorate over 1 year. As an example, the course of the 30:15 ratio data is shown over 1 year (Fig. 1).

Furthermore, thirteen participants already showed clinical signs of diabetic sensory neuropathy at T0 (e.g., reduced vibration threshold), which was finally interpreted as a consequence of a hyperglycemic metabolic state. We followed these patients with overt neuropathy and compared the functional data over the individual study period (see Table 2, especially NES, CPT).

When looking at the duration of diabetes in the two groups, it became obvious that group A consisted of
newly diagnosed diabetics (0.23 years; one long-standing outlier not included). In contrast, in group B the average duration was 20 years (maximum 57 years). Two patients from group A had been diagnosed because of ketoacidotic coma. Due to the shorter duration of diabetes a lesser degree of chronic nerve damage was expected in group A. In group B with more expected chronic nerve damage we looked at the progression of neuropathy over the study period (Table 2).

**Discussion**

Our study aimed at testing the hypothesis that a rapid, treatment induced reduction of HbA1c may be associated with a more common and pronounced induction of TIND in patients with severe diabetes of type 1 or 2 (HbA1c > 8.5%). Deterioration of any one test parameter between T0 and T1 in patients with a rapidly lowered HbA1c was considered a candidate becoming a predictor of TIND. Against our expectation we found at most trends but no significant differences in any of the performed neurophysiological tests over time when comparing group A with group B. Within group A that was presumed to develop TIND, we could not reveal any major distinctive patterns of abnormality despite patients’ HbA1c rapidly declining upon treatment initiation. However, the low adherence to the planned study procedures prevented us from obtaining a sufficient number of observations to formally confirm or refute our hypothesis.

We are aware that the major difference, in the duration of diabetes between groups A, i.e. a few months vs. several years in group B may be an important factor...
influencing the observations reported here. This relates both to the underlying chronic neuropathy state and to the poor treatment efficacy over long periods of time in group B. Patients in this group may more commonly show poor adherence to adequate treatment and dietary behavior and may consequently also be candidates for poor study compliance as noted here.

Our screened 60 patients were all informed and trained how to comply with the treatments and tests. Before each visit, there was a telephone reminder (as effective means of increasing adherence) [17] of the examination date 1 week in advance. When asked for their reasons for discontinuing study participation, patients indicated that they rather wanted to focus on treating their diabetes than participating in study tests. Other reasons were concomitant illness, distance to the study site and that the many tests were too time consuming and even annoying. One additional reason for non-adherence may be the lack of having structured educational group sessions to support adherence. Still, it has not formally been shown in our German patient population that measures like group sessions would dependably improve adherence [18]. It remains to be tested if adherence would have been substantially better even with regular structured sessions.

One additional reason for non-adherence may be the inability to recruit and motivate participants to follow the test protocol which involved a number of visits. Given the observed standard deviation of 0.16 [20] at T1 in our present pooled data, a future TIND trial would need to enroll up to 500 patients to detect a minimal clinically important difference (MCID) at a power of 80%. Any future trial with this number of patients would warrant a multi-center design and should include educational training sessions and more incentives to increase adherence.

As a result of the reduced compliance demonstrated in this pilot study, especially in patients with long duration of diabetes, a follow-up study should include newly diagnosed diabetics who are also likely to have no or less pre-existing neuropathy.

Conclusions
In conclusion, the observations in Group A patients are consistent with involvement of parasympathetic and sympathetic C- and A-delta fibers as a potential indicator of treatment-related small fiber neuropathy. Because of the inability to recruit and motivate participants to follow the test protocol which involved a number of visits, a future TIND trial would need to enroll up to 500 patients to detect a minimal clinically important difference (MCID) at a power of 80%. Any future trial with this number of patients would warrant a multi-center design and should include educational training sessions and more incentives to increase adherence.

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**Table 2** Electrophysical parameters of group A and B and correlation analysis of all participants

|                  | T0               | T1               | p value | T3               | p value |
|------------------|------------------|------------------|---------|------------------|---------|
| 30:15 ratio      | 1 ± 0.12 (1.1 ± 0.3) | 0.95 ± 0.1 (1.2 ± 0.2) | 0.12 (0.8) | 1 ± 0.13 (0.9 ± 0.2) | 0.62 (0.3) |
| E/I ratio        | 2.91 ± 2.7 (2.6 ± 1) | 2 ± 0.9 (2.6 ± 0.7) | 0.18*** (0.8) | 2.4 ± 0.9 (2.7 ± 1) | 0.65 (0.3) |
| PDD (mm)         | 5.6 ± 0.5 (5.4 ± 1.7) | 5 ± 1.5 (5.3 ± 1.5) | **0.02** (0.6) | 5.4 ± 1 (5.4 ± 1.5) | 0.3 (0.6) |
| CPT (°C)         | 27.3 ± 3 (22 ± 9) | 25.1 ± 9 (24 ± 3.2) | 0.1*** (0.4***)) | 26.2 ± 5 (20 ± 10) | 0.4*** (0.6***)) |
| NES              | 2.8 ± 2.3 (4.1 ± 4) | 1.2 ± 1.3 (2.4 ± 2.3) | 0.04 (0.2) | 1.8 ± 1.9 (3.1 ± 2.9) | 0.2 (0.7) |
| NSS              | 4 ± 5 (4.5 ± 5) | 3.2 ± 5 (10.9 ± 11) | 0.3 (0.08) | 4.8 ± 8 (11.2 ± 15) | 0.5 (0.2) |

**Correlation analysis**

| Difference between T0 and T1 | Pearson correlation coefficient | p value |
|------------------------------|--------------------------------|---------|
| 30:15 ratio                  | 0.121                          | 0.6     |
| Valsalva ratio               | 0.02                           | 0.94    |
| E/I ratio                    | 0.4                            | 0.08    |
| PDD                          | 0.2                            | 0.45    |
| CPT                          | 0.21                           | 0.37    |
| WPT                          | 0.06                           | 0.81    |
| Latency of SSR right hand    | − 0.16                         | 0.5     |
| Sweat rate                   | − 0.034                        | 0.9     |

**Comparisons between T0 and T3**
p-value according to Wilcoxon-test, ***t-test
effort is encouraged for achieving a high number of patients.

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Authors’ contributions
PB, JC and YH planned the study, PB and YH collected and analyzed the patient data. PB, YH, KVT, and JC evaluated and interpreted the patient data and wrote the manuscript. PB, MB, KVT, and JC re-evaluated the data and co-edited the manuscript. All authors read and approved the final manuscript.

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Availability of data and materials
The sets of data generated during the current study are not publicly available due to privacy regulations but are available from the corresponding author on reasonable request.

Declarations

Ethics approval and consent to participate
The study protocol was approved by the Ethics Committee of University Leipzig (No. 241-2009-0911209). All participants gave written informed consent.

Competing interests
The authors declare that they have no competing interests.

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References
1. Chandler E, Brown M, Wintergerst K, Doll E. Treatment induced neuropathy of diabetes (TIND) in pediatric and adult populations: a literature review of the evidence. J Clin Endocrinol Metab. 2019;105:395–8. https://doi.org/10.1210/clinem/dgz067.
2. Gibbons CH, Freeman R. Treatment-induced diabetic neuropathy: a reversible painful autonomic neuropathy. Ann Neurol. 2010;67:534–41. https://doi.org/10.1002/ana.21952.
3. Knopp M, Srikantha M, Rajabally YA. Insulin neuritis and diabetic cachectic neuropathy: a review. Curr Diabetes Rev. 2015;9:267–74. https://doi.org/10.2174/1573399811309000007.
4. Dabby R, Sadeh M, Lampi Y, Gilad R, Watemberg N. Acute painful neuropathy induced by rapid correction of serum glucose levels in diabetic patients. Biomed Pharmacother. 2009;63:707–9. https://doi.org/10.1016/j.biopha.2008.08.011.
5. Gibbons CH, Freeman R. Treatment-induced neuropathy of diabetes: an acute, iatrogenic complication of diabetes. Brain. 2015;138:43–52. https://doi.org/10.1093/brain/awu307.
6. Gibbons CH. Treatment-induced neuropathy of diabetes. Curr Diab Rep. 2017;17:127. https://doi.org/10.1007/s11892-017-0960-6.
7. Hwang YT, Davies G. ‘Insulin neuritis’ to ‘treatment-induced neuropathy of diabetes’: a new name, same mystery. Pract Neurol. 2016;16:53–5. https://doi.org/10.1136/practneurol-2015-001215.
8. Baum P, Koj S, Klötting N, Blüher M, Classen J, Paeschke S, et al. Treatment-induced neuropathy in diabetes (TIND)-developing a disease model in type 1 diabetic rats. Int J Mol Sci. 2021;22:1571–85. https://doi.org/10.3390/ijms2204157-85.
9. Chantelau E, Meyer-Schwickerath R. Reversion of early worsenings of diabetic retinopathy by deliberate restoration of poor metabolic control. Ophthalmol Int J. Ophthalmol. 2003;21:373–7. https://doi.org/10.1159/000071355.
10. Campos C. Chronic hyperglycemia and glucose toxicity: pathology and clinical sequelae. Postgrad Med. 2012;124:90–7. https://doi.org/10.3810/pgm.2012.11.2615.
11. Siddique N, Durcan R, Smyth S, Tun T, Sreenan S, McDermott BH. Acute diabetic neuropathy following improved glycaemic control: a case series and review. Endocrinol Diabetes Metab Case Rep. 2020;1:1–4. https://doi.org/10.1530/EDM-19-0140.
12. Callaghan BC, Cheng HT, Stables CL, Smith AL, Feldman EL. Diabetic neuropathy: clinical manifestations and current treatments. Lancet Neurol. 2012;11:521–34. https://doi.org/10.1016/S1474-4422(12)70065-0.
13. Vinik A, Maser RE, Mitchell BD, Freeman R. Diabetic autonomic neuropathy. Diabetes Care. 2003;26:1553–79. https://doi.org/10.2337/diacare.26.5.1553.
14. Dutsch M, Markthol H, Michelson G, Neundörfer B, Hilz MJ. Pulpipigraphy refines the diagnosis of diabetic autonomic neuropathy. J Neurol Sci. 2004;222:75–81. https://doi.org/10.1016/j.jns.2004.04.008.
15. Chong PST, Cros DP. Technology literature review: quantitative sensory testing. Muscle Nerve. 2004;29:734–47. https://doi.org/10.10102/1002/mus.20053.
16. Fay M, Proshcan M, Wilcoxon–Mann–Whitney or t-test? On assumptions for hypothesis tests and multiple interpretations of decision rules. Stat Surv. 2010;4:1–39. https://doi.org/10.1214/09-SS051.
17. Capoccia K, Odegard P, Letassi Y. Medication adherence with diabetes medication: a systematic review of the literature. Diabetes Educ. 2016;42:34–71. https://doi.org/10.1177/0145721715619038.
18. Roter DL, Hall JA, Merisca R, Nordstrom B, Cretin D, Svarstad B. Effectiveness of interventions to improve patient compliance: a meta-analysis. Med Care. 1998;36:1138–61. https://doi.org/10.1097/00005650-199808000-00004.
19. Doggrell SA, Warot S. The association between the measurement of adherence to anti-diabetes medication and the HbA1c. Int J Clin Pharm. 2014;36:488–97. https://doi.org/10.1007/s11096-014-9929-6.
20. Chhabra SK, De S. Cardiovascular autonomic neuropathy in chronic obstructive pulmonary disease. Respir Med. 2005;99:126–33. https://doi.org/10.1016/j.rmed.2004.06.003.

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