Hemoglobin A1c modifies the association between triglyceride and time in hypoglycemia determined by flash glucose monitoring in adults with type 1 diabetes: implications for individualized therapy and decision-making

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Background: We aimed to investigate the associations of flash glucose monitoring (FGM)-derived metrics with lipid profiles and identify potential modifiers of these associations among adults with type 1 diabetes (T1D).

Methods: A cross-sectional study was conducted among 108 Chinese adults with T1D who used FGM for 14 consecutive days. The relationship between FGM-derived metrics and lipid variables and potential modifiers were identified using interaction and subgroup analysis.

Results: Serum triglyceride level inversely correlated with time below range (glucose <3.9 mmol/L) and time in range (glucose 3.9–10.0 mmol/L) and positively correlated with time above range (glucose >10.0 mmol/L) (Spearman’s r=−0.34, −0.25, 0.34, respectively, all P<0.01). Additionally, triglyceride levels had positive correlation with absolute measures of glycemic variability (GV) but not with the coefficient of variation for glucose (Spearman’s r=0.12, P>0.05), a relative measure. Multivariate linear regression analysis adjusting for potential confounders including gender, age, disease duration, body mass index (BMI), daily insulin dose, fasting C-peptide, and dyslipidemia medication use showed that higher triglyceride level independently predicted decrease in time below range and time in range and increase in time above range (all P<0.01). Furthermore, interaction analysis found that the interaction between HbA1c and triglyceride was significant in the time below range (P for interaction =0.034). The association between triglyceride and time below range differed substantially after stratification by HbA1c, which was significant in those with HbA1c <7.0% whereas inconsequential among those with HbA1c ≥7.0%. In those with HbA1c <7.0% (n=44), the area under receiver operating characteristic curve of triglyceride predicting achievement of targets of time below range (<4%) was 0.856 (95% confidence interval 0.688–1.000, P=0.042) with an optimal cutoff value of 0.50 mmol/L (sensitivity 100%, specificity 66.7%, positive predictive value 94.4%).

Conclusions: In adults with T1D, HbA1c may be a potential modifier of the association between triglyceride and time below range, suggesting it might be necessary for those with HbA1c <7.0% accompanied by lower triglyceride levels to set a less intensive glycemic target to minimize risk of hypoglycemia.

Keywords: Flash glucose monitoring (FGM); glucose variability; time in ranges; triglyceride; type 1 diabetes (T1D)

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Introduction

Type 1 diabetes (T1D) is characterized by absolute deficiency of insulin secretion caused by immune-mediated pancreatic β-cell destruction, resulting in lifelong dependence on exogenous insulin treatments. Drastic glycemic variability (GV) and recurrent hypoglycemia, a hallmark of T1D, pose an immense challenge in maintaining optimal glycemic control in T1D. Moreover, both have been associated with long-term diabetes complications as well as mortality (1,2), apart from the well-established risk factor hemoglobin A1c (HbA1c). Although, HbA1c represents overall glycemic control over a period of 2 to 3 months, it cannot provide specific information about the amplitude of GV and the occurrence of acute glycemic events including hypoglycemia and hyperglycemia. HbA1c thus fails to optimize individualized therapy and decision-making in diabetes management (2-6), especially for T1D. Even though HbA1c is the most canonical standard for glycemic control, other metrics such as GV, glucose patterns, frequency of hypoglycemia, and time in target are equally important components of glycemic management of T1D, which could facilitate individualized therapy and target setting (6). Continuous glucose monitoring (CGM) has garnered increasing attention for its capability in providing comprehensive information on GV, time in target, glucose patterns, as well as frequency and severity of hypoglycemia and hyperglycemia. More recently, there has been a spike in interest in flash glucose monitoring (FGM) in routine clinical practice for its easy access to glucose readings without need for calibration with finger prick glucose tests and non-inferiority to other well-established CGM sensors in terms of device accuracy (7). Mounting evidence from randomized clinical trials and real-world studies demonstrated that patients with T1D benefited from FGM use in various aspects including improvement of HbA1c (8), reduction of GV and hypoglycemia (9), as well as enhancement in treatment satisfaction and mental well-being (10). Similar results were seen among individuals with type 2 diabetes, showing that FGM could lower HbA1c and improve treatment satisfaction without increasing hypoglycemia (11). With substantial rise in FGM use in clinical settings, GV and time in range metrics derived from FGM have exerted increasing impact on individualized therapy and decision-making in T1D; however, this effect has been largely undefined and warrants further investigation. It has been well-established that lipid profiles are closely related to glycemic control determined by HbA1c in T1D (12,13). Nevertheless, it remains unexplored how GV and time in range metrics derived from FGM are associated with alteration in lipids in T1D and whether there exist variables that modify these associations. Therefore, we sought to conduct a cross-sectional study in adults with T1D to investigate the associations of GV and time in range metrics derived from FGM with lipid profiles and identify the potential modifiers of these associations. We present the following article in accordance with the STROBE reporting checklist (available at http://dx.doi.org/10.21037/atm-20-6344).

Methods

Study population

We performed a cross-sectional study among Chinese adults with T1D who visited the endocrinology department of Peking Union Medical College Hospital from November 2018 to June 2019. The inclusion criteria were as follows: (I) age ≥18 years; (II) clinical diagnosis of T1D, which was defined as diabetes with fasting C-peptide level ≤0.8 ng/ml at diabetes onset; (III) body mass index (BMI) ≤35 kg/m²; (IV) continuous insulin therapy for at least 3 months prior to enrollment in the study; (V) no history of diabetic ketoacidosis or hyperglycemic hyperosmolar syndrome in the past 3 months. The study was conducted in accordance with the Declaration of Helsinki (as revised in 2013). The study was conducted under a protocol approved by the Ethics Committee of Peking Union Medical College Hospital (Approval No. B275). All participants voluntarily signed the written informed consent form. A total of 114 adults were enrolled in the study. All participants were assigned to wear FGM sensors (Freestyle Libre, Abbott, Witney, UK) for 14 consecutive days. The participants were blinded to their FGM data to avoid any bias with respect to GV from the participants. To guarantee the accuracy and reliability of the data, those with more than 70% FGM data available within the 14-day period were finally included in the study (14). Consequently, data of 108 individuals were included in the
final analyses.

**Laboratory analysis**

The FGM sensors were inserted on day 0 and removed on day 14 at the same time of day. The laboratory analysis of blood samples was performed at baseline (study day 0). After an overnight fast for at least 10 hours, venous blood samples were drawn in the morning. The level of plasma glucose was determined by the glucose oxidase assay, and the level of C-peptide was detected using chemiluminescence immunoassay with a detection limit of 0.01 ng/mL. HbA1c was assessed using a high-performance liquid chromatography assay. Total cholesterol (TC), total triglyceride (TG), high-density lipoprotein cholesterol (HDL-C), and low-density lipoprotein cholesterol (LDL-C) as well as level of uric acid (UA) were measured using an automatic analyzer.

**Analysis of data from FGM and calculations of FGM-derived metrics**

The data recorded on the day the glucose sensor for FGM was inserted were excluded to avoid any bias due to insufficient glucose stabilization between the sensor and the interstitial fluid during the first 24 hours after insertion of the sensor. Thirteen validated 24-h glycemic profiles (calculated from glucose readings obtained at 15-min intervals), were used for calculations of FGM-derived metrics, including mean of glucose (Mean), coefficient of variation for glucose (CV) (15), standard deviation of glucose (SD), mean amplitude of glucose excursion (MAGE) (16), mean of daily differences (MODD) (17), interquartile range of glucose (IQR), time below range (glucose <3.9 mmol/L) (TBR), time in range (glucose 3.9–10.0 mmol/L) (TIR), time above range (glucose >10.0 mmol/L) (TAR).

**Statistical analyses**

Statistical analyses were implemented using RStudio version 1.2.1335 (https://www.rstudio.com). Continuous data were presented as mean ± SD (normal distribution) or median (IQR) (skewed distribution), whereas categorical data were presented as percentages. Spearman’s correlation analysis was performed to explore bivariate correlations. Stepwise multivariate linear regression analyses were conducted to determine the independent predictors for the FGM-derived variables after adjusting for potential confounders. To determine performances of the lipid variables for predicting the FGM-derived variables, receiver operating characteristic (ROC) curves were established. Statistical significance was set at a two-sided P value <0.05.

**Results**

**Characteristics of the study population**

As presented in Table 1, data from a total of 108 participants (39 males and 79 females) aged between 18 and 77 years were included in the final analyses. The median of disease duration and HbA1c was 7.7 years and 7.4%, respectively. The fasting C-peptide level of the participants was quite low with a median of 0.01 ng/mL. The lipid profiles were within normal ranges in most participants, and the percentage of dyslipidemia medication use was as low as 8.3% (8/108), probably because of the relatively young age and normal-to-low BMI of the participants. The average CV was 40%, indicating that more than half of the patients suffered from frequent hypoglycemic episodes because a CV higher than 36% suggests labile diabetes. According to the recommendations from the international consensus on time in range (14), the targets for glycemic control in adults with T1D were as follows: TIR >70%, TBR <4%, and TAR <25%. The average TIR was 56.68%, whereas the median of TBR and TAR was 7.77% and 27.84%, respectively. The percentage of patients with TIR >70%, TBR <4%, or TAR <25% was 18.5%, 30.6%, and 43.5%, respectively, suggesting that targets of glycemic control were not achieved in most patients with T1D.

**Correlations between FGM-derived metrics and clinical variables**

As shown in Figure 1, positive correlations of HbA1c with SD, MAGE, MODD, IQR, TIR, and TAR (Spearman’s r=0.69, 0.49, 0.34, 0.43, 0.46, 0.56, and 0.62, respectively, all parameters had a P<0.01) were observed. Interestingly, TG level was inversely correlated with TBR and TIR, and positively correlated with TAR (Spearman’s r=-0.34, -0.25, and 0.34, respectively, P<0.01 in all correlations). Additionally, TG level was not correlated with CV (Spearman’s r=0.12, P=0.05), a relative measure of GV, but was positively correlated with absolute measures of GV including SD (Spearman’s r=0.24, P<0.01), MAGE (Spearman’s r=0.25, P<0.01), MODD (Spearman’s r=0.20,
Independent predictive role of TG for time in ranges determined by FGM

To investigate the predictive role of TG for time in ranges determined by FGM, stepwise multivariate linear regression analysis was performed to adjust for potential confounders. Gender and age were adjusted in model 1. Subsequently, based on model 1, disease duration, BMI, daily insulin dose, and fasting C-peptide were further adjusted in model 2. Moreover, based on model 2, dyslipidemia medication use was further adjusted in model 3. As indicated in Table 2, TG remained significantly associated with TBR, TIR, and TAR after stepwise adjustment for potential confounders in models 1, 2, and 3, suggesting its independent predictive role in time in ranges determined by FGM among adults with T1D.

Differed association between TG and TBR depending on HbA1c

To investigate the impact of clinical variables on the relationship between TG and time in ranges derived from FGM, interaction and subgroup analyses were performed. Intriguingly, interaction was observed between HbA1c and TG in TBR (P for interaction =0.034), with subgroup analysis stratified according to HbA1c; HbA1c <7.0% showed a borderline statistical significance (P for interaction =0.083), whereas no interactions were found between other clinical variables and TG in TIR and TAR (P for interaction >0.10 in all evaluated parameters). Furthermore, the association between TG and TBR differed substantially when patients were stratified by HbA1c. A moderate-to-strong negative correlation between TG and TBR was observed among patients with HbA1c <7.0% (r=−0.43, P=0.006), whereas such an association was not observed among those with HbA1c ≥7.0% (r=−0.13, P=0.320) (Figure 2).

Assessment of predictive performance of TG for the target achievement of time in ranges

To assess the performance of TG in predicting whether the recommended targets of time in ranges (TBR <4%, TIR >70%, TAR <25%) could be achieved, ROC curves were established and area under the curves were calculated. Figure 3 indicated significant predictive performance

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Table 1 Baseline characteristics of the study population

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>Participants (n=108)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Male, n (%)</td>
<td>39 (36.1%)</td>
</tr>
<tr>
<td>Onset (year)</td>
<td>27.84 (21.21–38.94)</td>
</tr>
<tr>
<td>Duration (years)</td>
<td>7.70 (3.02–13.28)</td>
</tr>
<tr>
<td>Age (years)</td>
<td>35.91 (30.33–49.39)</td>
</tr>
<tr>
<td>BMI (kg/m²)</td>
<td>21.07 (19.71–23.12)</td>
</tr>
<tr>
<td>SBP (mmHg)</td>
<td>114.00 (103.00–124.00)</td>
</tr>
<tr>
<td>DBP (mmHg)</td>
<td>69.51±9.57</td>
</tr>
<tr>
<td>WC (cm)</td>
<td>76.61±8.96</td>
</tr>
<tr>
<td>HC (cm)</td>
<td>94.42±5.88</td>
</tr>
<tr>
<td>WHR</td>
<td>0.81±0.07</td>
</tr>
<tr>
<td>Daily insulin dose (U/d)</td>
<td>35.69±13.22</td>
</tr>
<tr>
<td>Daily insulin dose (U/kg/d)</td>
<td>0.60±0.20</td>
</tr>
<tr>
<td>Basal insulin (U/d)</td>
<td>14.25±6.71</td>
</tr>
<tr>
<td>Meal insulin (U/d)</td>
<td>21.44±9.32</td>
</tr>
<tr>
<td>HbA1c (%)</td>
<td>7.40 (6.55–8.40)</td>
</tr>
<tr>
<td>FPG (mmol/L)</td>
<td>8.80 (5.90–12.50)</td>
</tr>
<tr>
<td>FCP (ng/ml)</td>
<td>0.01 (0.01–0.28)</td>
</tr>
<tr>
<td>UA (mmol/L)</td>
<td>252.00 (201.00–287.00)</td>
</tr>
<tr>
<td>TC (mmol/L)</td>
<td>4.68±1.01</td>
</tr>
<tr>
<td>TG (mmol/L)</td>
<td>0.56 (0.44–0.76)</td>
</tr>
<tr>
<td>HDL-C (mmol/L)</td>
<td>2.00 (1.60–2.54)</td>
</tr>
<tr>
<td>LDL-C (mmol/L)</td>
<td>1.94 (1.44–2.44)</td>
</tr>
<tr>
<td>Dyslipidemia medication (%)</td>
<td>8.3% (9/108)</td>
</tr>
<tr>
<td>Mean (mmol/L)</td>
<td>7.92 (6.72–9.52)</td>
</tr>
<tr>
<td>SD</td>
<td>3.34±0.96</td>
</tr>
<tr>
<td>CV (%)</td>
<td>40.09±7.91</td>
</tr>
<tr>
<td>MODD</td>
<td>3.22 (2.50–3.88)</td>
</tr>
<tr>
<td>MAGE</td>
<td>6.87±1.77</td>
</tr>
<tr>
<td>IQR (mmol/L)</td>
<td>4.76±1.53</td>
</tr>
<tr>
<td>TAR (%)</td>
<td>27.84 (14.72–48.76)</td>
</tr>
<tr>
<td>TBR (%)</td>
<td>7.77 (3.59–18.07)</td>
</tr>
<tr>
<td>TAR (%)</td>
<td>58.68±16.89</td>
</tr>
</tbody>
</table>

BMI, body mass index; SBP, systolic blood pressure; DBP, diastolic blood pressure; WC, waist circumference; HC, hip circumference; WHR, waist-to-hip ratio; HbA1c, hemoglobin A1c; FPG, fasting plasma glucose; FCP, fasting C-peptide; UA, uric acid; TC, total cholesterol; TG, triglyceride; HDL-C, high-density lipoprotein cholesterol; LDL-C, low-density lipoprotein cholesterol; Mean, mean of glucose; CV, coefficient of variation for glucose; SD, standard deviation of glucose; MAGE, mean amplitude of glucose excursion; MODD, mean of daily differences; IQR, interquartile range of glucose; TBR, time below range (glucose <3.9 mmol/L); TIR, time in range (glucose 3.9–10.0 mmol/L); TAR, time above range (glucose >10.0 mmol/L).
Figure 1 Spearman correlation heat map between FGM-derived metrics and clinical variables. The values of the color bar represent the Spearman’s r. The red color means a positive correlation whereas the purple color means negative correlation. Darker color indicates increased correlation. *P<0.05; **P<0.01. BMI, body mass index; SBP, systolic blood pressure; DBP, diastolic blood pressure; WC, waist circumference; HbA1c, hemoglobin A1c; FCP, fasting C-peptide; UA, uric acid; TC, total cholesterol; TG, triglyceride; HDL-C, high-density lipoprotein cholesterol; LDL-C, low-density lipoprotein cholesterol; Mean, mean of glucose; CV, coefficient of variation for glucose; SD, standard deviation of glucose; MAGE, mean amplitude of glucose excursion; MODD, mean of daily differences; IQR, interquartile range of glucose; TBR, time below range (glucose <3.9 mmol/L); TIR, time in range (glucose 3.9–10.0 mmol/L); TAR, time above range (glucose >10.0 mmol/L).

Table 2 Stepwise multivariate linear regression models with level of serum TG predicting time in ranges determined by FGM

<table>
<thead>
<tr>
<th>Model</th>
<th>TBR</th>
<th>P value</th>
<th>TIR</th>
<th>P value</th>
<th>TAR</th>
<th>P value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Model 1</td>
<td>β: -6.69</td>
<td>0.012</td>
<td>β: -13.61</td>
<td>0.003</td>
<td>β: 19.90</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Model 2</td>
<td>β: -7.01</td>
<td>0.006</td>
<td>β: -13.75</td>
<td>0.002</td>
<td>β: 20.21</td>
<td>0.001</td>
</tr>
<tr>
<td>Model 3</td>
<td>β: -7.46</td>
<td>0.004</td>
<td>β: -15.05</td>
<td>&lt;0.001</td>
<td>β: 21.31</td>
<td>&lt;0.001</td>
</tr>
</tbody>
</table>

Model 1: adjusted for gender and age; Model 2: adjusted for model 1 + duration + BMI + daily insulin dose + FCP; Model 3: adjusted for model 2 + dyslipidemia medication. TBR, time below range (glucose <3.9 mmol/L); TIR, time in range (glucose 3.9–10.0 mmol/L); TAR, time above range (glucose >10.0 mmol/L).
for TBR <4% (AUC = 0.704, 95% CI: 0.600–0.807, P=0.001) (Figure 3A) and TAR <25% (AUC = 0.693, 95% CI: 0.588–0.798, P=0.001) (Figure 3C), whereas no significant predictive role of TG for TIR >70% was observed (AUC = 0.576, 95% CI: 0.449–0.704, P=0.296) (Figure 3D). Given that the association between TG and TBR differed substantially after stratification by HbA1c and was significant in those with HbA1c <7.0%, we further conducted ROC analysis in those with HbA1c <7.0% (n=44) with an AUC 0.856 (95% CI: 0.688–1.000, P=0.042) and an optimal cutoff value of TG of 0.50 mmol/L (sensitivity 100%, specificity 66.7%, positive predictive value 94.4%) using Youden’s index (Figure 3B).

Discussion

In the current study, we revealed, for the first time, the relationships of serum TG with measures of GV and time in range metrics derived from FGM. We found correlations between TG and absolute measures of GV, and identified the independent predictive role of TG for time in ranges determined by FGM. Further, we identified the effect of modification by HbA1c on the association between TG and TBR, indicating that serum TG testing may provide an adjunctive marker for assessment of GV and time in ranges in T1D. HbA1c may act as a potential modifier of associations between TG and TBR. It is noteworthy that modification by HbA1c could support and facilitate individualized therapy and decision-making for clinicians in glycemic management of T1D.

GV metrics determined by CGM could be assessed by CV, a relative measure, and absolute measures including SD, MAGE, MODD, and IQR. CV with a recognized cutoff value of 36% has been used as a threshold to separate the diabetes from labile diabetes in previous studies (2,18). We did not find any correlation of CV with HbA1c or lipid profiles, whereas absolute measures of GV were positively correlated with HbA1c and TG. HDL-C was previously reported to correlate with CGM-derived CV in T1D, while TG level was positively correlated with CGM-derived SD in T2D without insulin therapy instead of T1D (19). This discrepancy might be explained by the small sample size of the previous study, different CGM devices, or different populations. Large-scale investigations are warranted to validate further the associations between the FGM-derived metrics and lipid profiles.

Accumulating evidence has shown that time in ranges including TIR, TAR, and TBR could be appropriate metrics of glycemic control in research and clinical settings (20). In the current study, a high TG level was independently associated with shorter TBR and TIR, and longer TAR, suggesting its potential value as an adjunctive marker of the quality of glycemic control. Previous studies demonstrated that elevated TG was closely related to chronic hyperglycemia indicated by high HbA1c in T1D (12,13,21). Our findings further contributed that TG is also associated with short-term hyperglycemia detected by FGM. We found that the serum TG level was also positively correlated with a series of indicators of insulin sensitivity including BMI, waist circumference, waist to hip ratio, and HbA1c (Spearman’s r=0.22, 0.22, 0.31, and 0.36, respectively, all P<0.05). Thus, the possible explanation for the association of TG with TBR and TAR could be that higher serum TG level was recognized as a marker of lower insulin sensitivity (22,23). Moreover, TG has been validated as one of the vital elements of insulin sensitivity clinical predictive models for T1D with favorable predictive performances (24,25). TIR has been recognized as another key indicator of effective and safe glycemic control in research and clinical settings, and thus, has been validated as a reliable outcome measure in clinical trials (26) and linked to long-term diabetes complications beyond HbA1c (27). Taking these observations into consideration, it should be noted that although higher TG indicates lower risk of hypoglycemia, it suggests less time spent in the target range and higher risk of hyperglycemia. In the glycemic management...
of T1D, minimizing hypoglycemia should be the first priority, targeting <4% of TBR, and then addressing the TIR (targeting >70% of TIR) or hyperglycemia (targeting <25% of TAR), as recommended by the international consensus (14). Our findings provide insights into clinical practices for clinicians in optimizing personalized medical regimen and decision-making, suggesting that adults with T1D with high TG might benefit more from adding sodium-glucose cotransporter 2 inhibitors as adjuncts to insulin, which has been demonstrated to help increase TIR and GV, without increasing TBR in T1D (28,29). Additionally, given that a higher level of TG may indicate lower insulin sensitivity, for those with high TG level, metformin might be another alternative of add-on therapy to insulin because metformin has shown to improve insulin sensitivity in T1D, regardless of baseline BMI (30,31). However, the results of metformin as an adjunctive therapy to insulin in T1D are contradictory. Metformin was reportedly not associated with improving overall glycemic control indicated by HbA1c or lowering TG with slight increase of severe hypoglycemia in a meta-analysis (32). Thus, although metformin could yield benefit in improving insulin sensitivity in T1D, and higher TG is associated with the lower TBR, add-on therapy of metformin in

Figure 3 Predictive performance of serum TG level for targets of time in ranges using ROC curves. The area under ROC curves of TG predicting (A) TBR <4% in all participants (n=108), (B) TBR <4% in participants with hemoglobin A1c <7% (n=44), (C) TAR <25% in all participants (n=108), and (D) TIR >70% in all participants (n=108) are 0.704 (95% CI: 0.600–0.807, P=0.001), 0.856 (95% CI: 0.688–1.000, P=0.042), 0.693 (95% CI: 0.588–0.798, P=0.001), and 0.576 (95% CI: 0.449–0.704, P=0.296), respectively. 95% CI, 95% confidence intervals; ROC curves, receiver operating characteristic curves; TBR, time below range (glucose <3.9 mmol/L); TAR, time above range (glucose >10.0 mmol/L); TIR, time in range (glucose 3.9–10.0 mmol/L).
T1D should be prescribed with caution even in those with high TG.

Interestingly, we found an interplay between HbA1c and TG in the TBR determined by FGM in adults with T1D. Our data supports the predictive ability of low serum TG level for longer time spent in hypoglycemia determined by FGM, which differed substantially depending on the overall glucose control indicated by HbA1c, suggesting that HbA1c possibly modifies the association between TG and the time spent in hypoglycemia. This finding may have clinical implications, allowing clinicians to make more targeted and informed therapy decisions in glucose management of T1D.

It has been previously demonstrated that lower HbA1c was associated with more frequent hypoglycemia, suggesting that improvements in glycemic control could increase the risk of hypoglycemia in T1D (33). Our observation provides additional information that is necessary for adults with T1D with good overall glycemic control and lower serum TG level to be especially more cautious about the elevated risk of hypoglycemia, and thus, a less intensive glycemic target should be set. Hence, facilitating individualized therapy regimen of diabetes management of T1D in the clinical practice.

Based on the results of ROC analysis, TG performed well in predicting T1D, achieving the recommended targets of TBR and TAR, whereas the result of TIR was disappointing. Moreover, TG performed significantly better in predicting whether the recommended target of TBR was achieved, with an optimal cutoff value of 0.50 mmol/L and a positive predictive value of 94.4%, suggesting that adults with HbA1c <7.0% accompanied by TG <0.50 mmol/L are at high risk of time spent in hypoglycemia, longer than 1 hour per day. Therefore, it may be necessary to set a less intensive glycemic target to minimize hypoglycemia. However, the sample size of the present study was small and the cutoff point has not been validated in another population, which limits the generalizability of the finding; therefore, it should be interpreted with caution. Further investigations are needed to validate the results in the future.

Although dyslipidemia is frequently observed and has been demonstrated as an important risk factor for cardiovascular diseases in patients with T1D (34), lipid profiles within normal ranges are not unusual for individuals with T1D (35). The study population was characterized with normal levels of TG in most cases, suggesting that the TG within normal range is also beneficial for the clinical decision-making in diabetes management of T1D.

The strengths of the present study are as follows: (I) the participants were blinded to the glucose readings derived from FGM, thus avoiding any bias with respect to FGM data from the participants. (II) We are the first to explore the association of lipid profiles with FGM-derived metrics, and FGM use is rapidly increasing in routine clinical practice. However, FGM-derived metrics have been largely undefined and warrant further evaluation.

Overall, several limitations also deserve to be addressed. First, the small sample size limits the reliability of our findings when applied to clinical scenarios and thus, they should be interpreted with caution and further investigations with larger populations are warranted. Second, the absence of confirmatory blood glucose measurements in this study might also limit the reliability of the FGM readings because the glucose level and rates of glucose concentration changes affect the accuracy of FGM (36,37). Finally, other confounding factors such as diet and exercise also exert significant impact on short-term GV; however, we did not collect and adjust these confounders in the current study.

Conclusions

Collectively, our findings suggest that lower serum TG levels independently predict shorter TBR and TIR, and longer TAR determined by FGM in adults with T1D. The association of TG with TBR was modified by HbA1c, which was significant in those with HbA1c <7.0% but obscure in those with HbA1c ≥7.0%. These data have important clinical implications for individualized therapy and decision-making in glucose management of T1D, suggesting that adults with T1D with HbA1c <7.0% and lower serum TG levels should be more cautious about the increased risk of hypoglycemia, and thus it may be necessary to set a less intensive glycemic target. Nevertheless, further investigations are still warranted to validate these findings and the clinical value in the future.

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Footnote

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Ethical Statement: The authors are accountable for all aspects of the work in ensuring that questions related to the accuracy or integrity of any part of the work are appropriately investigated and resolved. The study was conducted in accordance with the Declaration of Helsinki (as revised in 2013). The study was conducted under a protocol approved by the Ethics Committee of Peking Union Medical College Hospital (Approval No. B275). All participants voluntarily signed the written informed consent form.

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