Four-year HbA1c and LDL-cholesterol trajectories among individuals with mental disorders and newly developed type 2 diabetes

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Funding information
Novo Nordisk Foundation, Grant/Award Numbers: NNF17SA0031406, NNF20SA0062874; Novo Nordisk Foundation Challenge Programme, Grant/Award Number: NNF14OC011633; Lundbeck Foundation, Grant/Award Number: R358-2020-2342; Danish Cancer Society, Grant/Award Number: R283-A16461; Independent Research Fund Denmark, Grant/Award Number: 7016-00048B

Abstract
The impact of different types of mental disorders on long-term glycemic and lipid trajectories following newly diagnosed type 2 diabetes (T2D) remains unknown. We used real-world clinical data in a population-based cohort to fill this knowledge gap. We found that individuals with new T2D and preexisting personality, anxiety, unipolar depression, or psychotic disorder had higher mean HbA1c levels over 4 years following the onset of T2D, whereas no differences were found regarding LDL-C levels. This knowledge should be considered in the management of T2D in these vulnerable groups.

KEYWORDS
cholesterol, diabetes mellitus, glycated hemoglobin a, glycemic control, LDL, mental disorders

Good glycemic and lipid control is critical to prevent development of disabling micro and macrovascular complications in type 2 diabetes (T2D) (ADA: Standards of Medical Care in Diabetes-2021, 2021). Results of a recent study suggested that individuals with new T2D and preexisting depression are as likely to successfully attain early Hemoglobin A1C (HbA1c) and low-density lipoprotein cholesterol (LDL-C) targets as are T2D individuals without depression (Rohde et al., 2021). However, the impact of depression and other mental disorders on long-term glycemic and lipid trajectories remain unknown. We used data from unique population-based Danish registers to fill this knowledge gap (Arendt et al., 2020; Schmidt et al., 2019).

We identified two nonmutually exclusive populations with incident T2D among residents of the Central and Northern Denmark Regions in the period from 2000 to 2016:

1. Incident HbA1c-defined T2D: Individuals with their first measurement of elevated HbA1c (≥6.5% [48 mmol/mol]) at age ≥30 years.
2. Incident glucose lowering drug (GLD)-treated T2D: Individuals that redeemed their first prescription for a GLD at age ≥30 years.

For both cohorts, we excluded individuals with a prior redeemed prescription for a GLD or prior hospital-based diagnosis of diabetes [Anatomical Therapeutic Chemical (ATC) code A10 and International Classification of Diseases (ICD-10) codes E10-E15, E160-E162, O24, T383A, M142, G590, G632, H280, H334, H450, H360, or N083]. Within these two cohorts, we identified all individuals who had been diagnosed (not mutually exclusive) with psychotic disorder (ICD-10 codes: F02-F29), bipolar disorder (ICD-10 codes: F30-F31), unipolar depression disorder (ICD-10 codes: F32-F33), anxiety disorder...
FIGURE 1  Trajectories of mean HbA1c and mean LDL-C levels based on everyday clinical practice measurements following incident HbA1c-defined (a, b) or incident GLD-treated (c, d) T2D, stratified according to history of hospital-diagnosed mental disorders.

HbA1c and LDL-C: HbA1c and LDL-C values from measurements ordered by any healthcare provider at any patient visit in the primary or secondary health care sector were included. The mean HbA1c and LDL-C levels were calculated for intervals spanning 4 months, using all available HbA1c and LDL-C values within these intervals.

p values: We used a t-test to compare HbA1c and LDL-C levels measured at 12, 24, 36, and 48 months of follow-up in individuals with a mental disorder versus no mental disorder.

Number of individuals with incident HbA1c-defined T2D: Personality disorder: 392, psychotic disorder: 753, anxiety disorder: 966, unipolar depression disorder: 1260, bipolar disorder: 284, no hospital-diagnosed mental disorder: 85,835.

Proportion initiating GLDs among individuals with incident HbA1c-defined T2D diabetes (median time to GLD, quartiles): 77.3% with personality disorder (67 days, 12–505), 72.6% with psychotic disorder (124 days, 14–522), 71.3% with anxiety disorder (75 days, 13–487), 66.1% with unipolar depression disorder (99 days, 15–550), and 65.5% with bipolar disorder (99 days, 15–550), and 65.1% with psychotic disorder, and 72.0% with no mental disorder.

Number of individuals with incident GLD-defined T2D diabetes: Personality disorder: 559, psychotic disorder: 926, anxiety disorder: 1240, unipolar depression disorder: 1520, bipolar disorder: 381, no hospital diagnosed mental disorder: 101,472.
HbA1c-defined T2D, the mean HbA1c levels at baseline were between 7.5% (58 mmol/mol) and 8.0% (64 mmol/mol), and then decreased by approximately 1% point over the course of approximately eight months, to mean levels between 6.6% (49 mmol/mol) and 7.0% (53 mmol/mol) (Figure 1a). Subsequently, a slight gradual increase in HbA1c levels were observed for all subgroups, but the mean HbA1c levels generally remained under the recommended 7% (53 mmol/mol) target (ADA: Standards of Medical Care in Diabetes-2021, 2021), except for the cohort members with personality, anxiety, or psychotic disorder. Specifically, individuals with personality, anxiety, unipolar depression, or psychotic disorder had higher (0.10%–0.36%) mean HbA1c levels compared to individuals without mental disorder, both at baseline and throughout follow-up (see Figure 1a for p values). The mean LDL-C levels at baseline were between 3.05 mmol/L and 3.34 mmol/L across groups (Figure 1b). During follow-up, the mean LDL-C gradually decreased in all subgroups to levels below the recommended 2.6 mmol/L (ADA: Standards of Medical Care in Diabetes-2021, 2021) at 48 months. Although the mean LDL-C levels at baseline were between 3.05 mmol/L and 3.34 mmol/L across groups (Figure 1b). During follow-up, the mean LDL-C gradually decreased in all subgroups to levels below the recommended 2.6 mmol/L (ADA: Standards of Medical Care in Diabetes-2021, 2021) at 48 months. Although the mean LDL-C levels tended to be higher for individuals with versus without mental disorders, no statistically significant differences were found (Figure 1b).

Among those with incident GLD-defined T2D, baseline HbA1c levels were higher than for the HbA1c-defined diabetes cohort (as expected since decision of GLD initiation is a proxy for severity). However, the HbA1c and LDL-C trajectories over time were similar to those described above for the HbA1c-defined cohort (Figure 1c, d).

In this study of incident T2D, we report that, compared to individuals without mental disorders, those with personality, anxiety, unipolar depression, or psychotic disorder had higher mean HbA1c levels over 4 years following the onset of T2D. No differences were found regarding LDL-C levels. These findings suggest that the management of T2D in individuals with personality, anxiety, unipolar depression, or psychotic disorder is suboptimal, possibly due to challenges with self-care or lacking initiation of compliance with GLD treatment and control regimens (Ward & Druss, 2015).

A central limitation to this study is that prior to 2012, HbA1c measurement was restricted mainly to individuals with anticipated problems regarding glycemic control and rarely used diagnostically. This may have caused selective inclusion of high HbA1c levels before 2012. Furthermore, only ≈50% of the patients had an HbA1c measurement taken in each 4-month interval. This proportion was however independent of mental disorder status (data not shown), which speaks against bias in the comparisons.

In conclusion, compared to those without mental disorder, individuals with T2D and personality, anxiety, unipolar depression, or psychotic disorder had higher HbA1c levels over time. This knowledge should be considered in the management of T2D in these vulnerable groups.

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How to cite this article: Kristensen, F. P., Rohde, C., Østergaard, S. D., & Thomsen, R. W. (2021). Four-year HbA1c and LDL-cholesterol trajectories among individuals with mental disorders and newly developed type 2 diabetes. Brain and Behavior, 11, e2372. https://doi.org/10.1002/brb3.2372