Effect of the GLP-1 receptor agonist semaglutide on metabolic disturbances in clozapine-treated or olanzapine-treated patients with a schizophrenia spectrum disorder: study protocol of a placebo-controlled, randomised clinical trial (SemaPsychiatry)

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ABSTRACT

Introduction Clozapine and olanzapine are some of the most effective antipsychotics, but both are associated with weight gain and relevant metabolic disturbances, including pre-diabetes and diabetes. Non-pharmacological/behavioural interventions have had limited effects counteracting these adverse effects. Semaglutide, a glucagon-like peptide 1 receptor agonist, is approved for the treatment of type 2 diabetes and obesity. We will investigate the long-term effects of add-on treatment with semaglutide once a week versus placebo once a week on the metabolic status in pre-diabetic patients diagnosed with a schizophrenia spectrum disorder who initiated clozapine or olanzapine treatment within the last 60 months.

Methods and analysis This is a 26-week, double-blinded, randomised, placebo-controlled trial. Altogether, 104 patients diagnosed with a schizophrenia spectrum disorder, aged 18–65 years, with pre-diabetes or diabetes will be randomised to injections of 1.0 mg semaglutide once a week or placebo for 26 weeks. The primary endpoint is change from baseline in HbA1c. Secondary endpoints include changes in body weight, hip and waist circumference and plasma levels of insulin, glucagon, glucose, and C-peptide, insulin sensitivity, beta cell function, hepatic function, fibrosis-4 score, lipid profile, incretin hormones, bone markers, bone composition, bone density, proteomic analyses and oxidative stress markers. Together with alcohol, tobacco and drug use, potential effects on the reward value of a sweet-fat stimulus, psychopathology, level of activity and quality of life will also be assessed.

STRENGTHS AND LIMITATIONS OF THIS STUDY

⇒ A 26-week, double-blinded, randomised, parallel-group, placebo-controlled, good clinical practice-monitored, multicentre trial has been chosen in accordance with the trial objectives.
⇒ This study will evaluate the effects of add-on therapy of early glucagon-like peptide 1 (GLP-1) receptor agonist (semaglutide) intervention to dysglycaemic clozapine-treated or olanzapine-treated patients diagnosed with a schizophrenia spectrum disorder.
⇒ The viability of GLP-1 injection treatment will be determined by an assessment of the patients’ adherence to self-administration once a week.
⇒ Lack of rescue medication may cause more patients to be excluded from the placebo arm compared with the active arm during the trial period.
⇒ The 26-week trial period could limit exploring the full potential of the treatment.

INTRODUCTION

Background and rationale Schizophrenia is a mental disorder with a heterogeneous genetic and neurobiological aetiology, affecting approximately 1% of the general population.1 Patients with schizophrenia have a shorter life expectancy of...
10–15 years compared with the general population.\textsuperscript{2–6} This excess mortality is largely explained by a high prevalence of obesity, metabolic disturbances, type 2 diabetes and cardiovascular diseases.\textsuperscript{4–7,11} Researchers have suggested that glucose homeostasis is altered in patients with schizophrenia from illness onset, indicating that this group of patients may be at increased risk of developing impaired glucose homeostasis and type 2 diabetes.\textsuperscript{12–14} This risk could be further aggravated as a consequence of a high prevalence of a sedentary lifestyle, low physical activity, dietary risk, smoking and reduced access to adequate healthcare in patients with schizophrenia.\textsuperscript{13–19} Antipsychotic medications play a central role in the treatment of schizophrenia, but, unfortunately, several antipsychotics are also associated with body weight increase and the development of metabolic disturbances including pre-diabetes and type 2 diabetes.\textsuperscript{20–22} The two antipsychotics olanzapine and clozapine are quite efficacious,\textsuperscript{23} but regrettably, olanzapine and clozapine confer the highest risk of inducing weight gain and metabolic disturbances, including pre-diabetes compared with other psychotropics.\textsuperscript{8,20} The weight gain after clozapine initiation is on average 5 kg within the first 12 weeks of treatment.\textsuperscript{20,26–27} Clozapine is associated with long-term weight gain even though the risk of weight gain appears to be highest in the initial treatment period.\textsuperscript{28,29} The manifestation of type 2 diabetes in this group of patients follows a similar trend, with around 10% of those receiving clozapine developing the condition within the first year of treatment, approximately 35% within the first 10 years, and nearly 45% within the first 21 years of treatment.\textsuperscript{30,31} Body weight loss and improvements in metabolic disturbances can be achieved by switching to more weight-neutral antipsychotic medication.\textsuperscript{32,35} However, both olanzapine and clozapine are potent treatments for schizophrenia, with clozapine as the recommended choice for treatment-resistant schizophrenia.\textsuperscript{34–37} Due to the efficacy of the compound in otherwise partial or fully treatment-resistant patients, switching to another antipsychotic is often not possible without worsening psychotic symptoms.\textsuperscript{38} Despite the adverse effects, long-term antipsychotic treatment is associated with decreased all-cause mortality compared with non-use of antipsychotic drugs,\textsuperscript{11,39–40} and clozapine is associated with the lowest all-cause mortality compared with other antipsychotic drugs.\textsuperscript{11,39–40} This may be the outcome of improved control of psychiatric symptoms due to antipsychotic use, which promotes secondary prevention as a healthier lifestyle and more frequent use of care services for physical illness.\textsuperscript{39,41}

Previously, limited effects have been demonstrated for counteracting antipsychotic-induced body weight gain and metabolic disturbances with non-pharmacological/behavioural interventions\textsuperscript{42–51} and adjunct pharmacological treatments.\textsuperscript{51–60} Several meta-analyses conclude evidence in favour of topiramate and metformin as adjunct pharmacological treatment.\textsuperscript{51–53,61} while individual lifestyle counselling is found to be the most effective non-pharmacological intervention for weight reduction.\textsuperscript{51}

Growing evidence suggests a potential benefit of adjunct pharmacological treatment with glucagon-like peptide 1 (GLP-1) glucagon-like peptide 1 receptor agonists (GLP-1RAs).\textsuperscript{62} GLP-1 is an incretin hormone secreted from the L cells in the intestinal mucosa in response to nutrients.\textsuperscript{63} GLP-1 stimulates insulin secretion and inhibits glucagon secretion, thereby lowering plasma glucose levels.\textsuperscript{63} Both effects are strictly glucose-dependent (more pronounced at higher levels of plasma glucose), and the effects cease as the level of plasma glucose reaches values below 4–5 mmol/L. Thus, the GLP-1RA keeps plasma glucose concentrations at lower levels without increasing the risk of hypoglycaemia.\textsuperscript{63} Additional effects include decreased gastric emptying, increased satiety, and decreased appetite and food intake leading to body weight loss\textsuperscript{65} with reduction of fat mass, android fat, trunk fat and lean body mass.\textsuperscript{64,65} Furthermore, recent data suggest that GLP-1RAs have beneficial effects on bone structure,\textsuperscript{66–68} and large-scale cardiovascular outcome trials have shown beneficial effects of GLP-1RAs on major adverse cardiovascular events in patients with type 2 diabetes at high risk of CVD.\textsuperscript{69–71} Currently, several studies have been conducted to investigate if these beneficial findings could be extended to the psychiatric population receiving antipsychotic medications.\textsuperscript{62,72–75} Our group previously reported that subcutaneous treatment for 16 weeks with the GLP-1RA liraglutide as adjunctive treatment to clozapine or olanzapine in patients with schizophrrenia-spectrum disorder with pre-diabetes improved glycated haemoglobin A1c (HbA1c) with approximately 0.2% (2.3 mmol/mol) glucose tolerance, and several other cardiometabolic disturbances and induced a 5.3 kg body weight loss compared with placebo.\textsuperscript{76} Recent meta-analyses have reported that GLP-1RAs are effective and tolerable for managing antipsychotic-associated body weight gain,\textsuperscript{77} especially for clozapine/olanzapine-treated patients.\textsuperscript{62} However, studies involving a large number of participants and longer treatment duration are needed in order to replicate the results.\textsuperscript{78,79} Future research should focus on GLP-1RAs with an extended half-life as they allow subcutaneous administration once a week as opposed to short-acting GLP-1RAs like liraglutide, which requires daily subcutaneous administration. Furthermore, results have indicated that this newer group of GLP-1RAs is more effective.\textsuperscript{80}

We will perform a clinical study with semaglutide (Ozempic), a GLP-1RA with an extended half-life of approximately 1 week, which permits once weekly subcutaneous administration. Ozempic 1.0 mg once a week was approved for the treatment of type 2 diabetes by the US Food and Drug Administration (FDA) in 2017 and the European Medicines Agency (EMA) in 2018. This trial will include a large number of participants, a long treatment duration and a novel GLP-1RA with a more potent efficacy, and an extended half-life expected to increase clinical outcomes.
Box 1 Endpoints

<table>
<thead>
<tr>
<th>Primary endpoint</th>
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<td>Glycated haemoglobin A1c.</td>
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<tr>
<th>Secondary endpoints</th>
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<tr>
<td>Body weight, hip and waist circumference.</td>
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<td>Vitals signs (blood pressure and pulse).</td>
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<tr>
<td>Peptide hormones (insulin, C-peptide and glucagon).</td>
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<td>Insulin sensitivity and beta cell function (evaluated by homeostatic model assessment).</td>
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<td>Lipid profile (low-density lipoprotein, high-density lipoprotein, triglycerides and cholesterol).</td>
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<tr>
<td>Hepatic function and fibrosis-4 score (thrombocytes, alanine aminotransferase, aspartate aminotransferase, alkaline phosphatase and bilirubin).</td>
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<tr>
<td>Incretin hormones (glucagon-like peptide 1, glucagon-like peptide 2 and glucose-dependent insulinotropic polypeptide).</td>
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<tr>
<td>Bone markers (vitamin D, calcium, phosphate, magnesium, parathyroid hormone, procollagen type 1 N-terminal propeptide, C-terminal telopeptide of type 1 collagen and osteocalcin).</td>
</tr>
<tr>
<td>Body composition and bone density (dual-energy X-ray absorption scan).</td>
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<tr>
<td>Psychopathology (Positive and Negative Syndrome Scale–Six Items, Clinical Global Impression Severity Scale), psychosocial disability (Cognitive Test and the Global Assessment of Psychosocial Disability Scale) and quality of life (Schizophrenia Quality of Life Scale).</td>
</tr>
<tr>
<td>Reward value of sweet–fat stimulus (clicker test).</td>
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<tr>
<td>Activity measurements (wearable activity device).</td>
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<tr>
<td>Alcohol, drug and tobacco use (Alcohol Use Disorders Identification Test, Drug Use Disorders Identification Test and Fagerström Test for Nicotine Dependence).</td>
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<tr>
<td>Proteomic analyses and measurement of oxidative stress.</td>
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</tbody>
</table>

Hypothesis

We hypothesise that add-on therapy with semaglutide once a week in dysglycaemic patients diagnosed with a schizophrenia spectrum disorder who initiated treatment with clozapine or olanzapine within 60 months, will improve overall glycaemic control and mitigate the antipsychotic-induced metabolic disturbances.

Objective and endpoints

This study was designed to evaluate the effects of 1.0 mg semaglutide once a week on the metabolic state in dysglycaemic patients diagnosed with a schizophrenia spectrum disorder, who initiated treatment with clozapine or olanzapine within 60 months prior to inclusion. The primary endpoint is the change from baseline in HbA1c. For secondary endpoints, see box 1.

METHODS AND ANALYSIS

Trial design, setting and timeline

This trial is a 26-week investigator-initiated, double-blinded, randomised, two-arm, parallel group, multicentre, good clinical practice (GCP)-monitored, and placebo-controlled study. All clinical visits will be conducted by investigators at Psychiatric Centre Copenhagen (PCK), Denmark; Psychiatric Centre Nordsjaelland (PCN), Denmark; and Aarhus University Hospital–Psychiatry (AUHP), Aarhus, Denmark, with PCK as the coordinating centre. It is the responsibility of the sponsor–investigator to inform sites of the terms and conditions of the protocol. All investigators will be trained in the study requirements, standardised clinical measurements and requirements for laboratory specimen collection including urine samples, counselling for adherence, and the eliciting of information from study participants in a uniform reproducible manner. Patient recruitment was initiated in September 2021. The number of patients to be included is 104. It is expected that PCK, PCN and AUHP will include 74, 10 and 20 patients, respectively. Initial and final visits will be performed at PCK and AUHP only. The study is expected to be completed in 2024.

Study population

Pre-diabetic and diabetic patients diagnosed with a schizophrenia spectrum disorder according to the criteria of the International Classification of Diseases, 10th Revision, WHO or the Diagnostic and Statistical Manual of Mental Disorders (Fifth Edition, the American Psychiatric Association) who initiated treatment with clozapine or olanzapine within the last 60 months and who can comply with the requirements for participation in the study (box 2) will be included.

The planned number of patients to be screened, that is, documented informed consent, is 125 (expected screen failure rate: up to 20%). The number of patients to be included, randomised and started on semaglutide or placebo is 104. The number of patients expected to complete the trial is 52, with 26 in each arm (figure 1). The patients will be recruited via one of the psychiatric centres in the Capital Region of Copenhagen involved in the study; PCK, Psychiatric Centre Glostrup, Psychiatric Centre Ballerup (PCB), and Psychiatric Centre Amager (PCA), PCN or from AUHP, Central Denmark Region. The expected number of dropouts is difficult to estimate. In a recent study in patients diagnosed with a schizophrenia spectrum disorder and in stable treatment with clozapine or olanzapine receiving daily subcutaneous injections of the GLP-1RA liraglutide or placebo over a 16-week trial period, we had a drop-out rate of approximately 7%.76 We expect the study withdrawal to be higher in the present, longer-lasting study in patients that have more recently, that is, within 60 months, initiated treatment with clozapine or olanzapine. Consequently, we expect a drop-out rate of up to 50% in our study population over the 26-week trial period. If the drop-out rate should turn out to be higher than 50%, more patients will be included to reach the goal of 52 patients completing the study. In that case, a change in sample size will be documented in a substantial protocol amendment.

Recruitment, inclusion and randomisation

To expand the knowledge about the project, we will contact psychiatrists and other employees working at the
Box 2 Eligibility criteria

Inclusion criteria
⇒ Informed oral and written consent.
⇒ Diagnosed with a schizophrenia spectrum disorder according to the criteria of the International Classification of Diseases, 10th Revision, or the Diagnostic and Statistical Manual of Mental Disorders, Fifth Edition.
⇒ Age 18–65 years (both included)
⇒ Body mass index of ≥25 kg/m².
⇒ Diagnosed with pre-diabetes or type 2 diabetes, with the following plasma levels: pre-diabetes: HbA1c 35–47 mmol/mol (5.4%–6.4%) or type 2 diabetes: HbA1c 48–57 mmol/mol (6.5%–7.4%).

Exclusion criteria
⇒ Acute worsening of psychosis based on a clinical evaluation (score of 6 or 7 on the Clinical Global Impression Severity Scale).
⇒ Coercive measures.
⇒ Women of childbearing potential who are pregnant, breastfeeding or have the intention of becoming pregnant.
⇒ Women who are not willing to use an adequate contraceptive during the full length of the study.
⇒ Patients treated with corticosteroids or other hormone therapy (except oestrogens).
⇒ Any active substance abuse or dependence (except for nicotine).
⇒ Impaired hepatic function (plasma liver transaminases >2 times upper normal limit).
⇒ Impaired renal function (serum creatinine >150 µmol/L).
⇒ Impaired pancreatic function (acute or chronic pancreatitis and/or plasma amylase >2 times upper normal limit).
⇒ Cardiac problems defined as decompensated heart failure (New York Heart Association class III/IV), unstable angina pectoris and/or myocardial infarction within the last 12 months.
⇒ Hypertension with systolic blood pressure of >180 mm Hg or diastolic blood pressure of >100 mm Hg.
⇒ Any condition that the investigator feels would interfere with trial participation.
⇒ Receiving any experimental or premarketing drug within the last 3 months.
⇒ Use of diabetes medication or weight-lowering pharmacotherapy within the last 3 months.
⇒ Known type 1 diabetes.
⇒ Suicidal behaviour as judged by the investigator and based on clinical evaluation.
⇒ Plasma HbA1c of >57 mmol/mol (7.4%) (tested twice), in which case the patient will be excluded from the study.
⇒ Any known contraindication towards the treatment with semaglutide.

Information, diagnosis, HbA1c, body mass index (BMI), medication record, name of the primary psychiatrist and psychiatric department. Patients meeting inclusion criteria will be informed about the possibility of participating in this study after permission to be contacted has been obtained from the treating psychiatrist. Before any trial-related procedures are performed, patients will be thoroughly informed about the study and sign an informed consent form (online supplemental material 1). All patients will receive oral and written information about the trial, including the most common adverse events, and the procedures involved in the study. Patients will have the opportunity to ask questions and have up to 48 hours to consider participation. A pretreatment evaluation will be conducted to screen patients according to the inclusion and exclusion criteria (box 2). When a patient is included in the study, the patients will be block-randomised into two groups. The randomisation will be stratified by age (two levels) and sex (two levels). The supplier of semaglutide pens (Novo Nordisk A/S) will provide a subject randomisation list (SRL) for randomisation purposes. The SRL will come in two versions: one blinded version for the appointed blinded investigators responsible for randomisation and one unblinded version for the unblinded staff who handle the trial product. Novo Nordisk A/S will be responsible for labelling and binding the semaglutide and placebo pens. A total dispensing unit number list (TDL) will be provided by Novo Nordisk A/S for the purpose of selecting dispensing units/pens containing the assigned treatment. The unblinded SRL and the TDL are only accessed and handled by appointed unblinded staff and will be uploaded in a separate randomisation module of the Research Electronic Data Capture (REDCap) program with no access for blinded staff. Via REDCap it will be possible for the blinded staff to automatically assign enrolled patients to the respective intervention via the blinded SRL. Unblinded staff will handle the dispensing of project medicine with double data entry to ensure integrity. Unblinded staff will not participate in any other parts of the trial.

Treatment protocol
The total trial duration is 26 weeks for each patient (table 1). The patients will receive blinded treatment in one of the two study arms: injection with 1.0 mg semaglutide or placebo once a week. Every 4 weeks, each patient will attend regular visits throughout the trial or be assessed by phone to evaluate adverse effects, and adherence to trial medication, and participate in testing with standard blood sampling (box 3) and a clinical examination (table 1).

In addition, all patients will be evaluated on the following rating scales and questionnaires: Schizophrenia Quality of Life Scale, Clinical Global Impression Severity Scale (CGI-S), Alcohol Use Disorders Identification Test, Drug Use Disorders Identification Test, Fagerström Test for Nicotine Dependence and the Global Assessment of Psychosocial Disability.
scale. Women not willing to use an adequate contraceptive during the full length of the study will be excluded. Women with childbearing potential will have a pregnancy test (ie, beta-human chorionic gonadotropin) performed before inclusion or if any suspicion of pregnancy arises during the study. If a patient gives informed consent to participate in future ancillary studies, extra material (serum/plasma/urine) will be stored in a biobank for possible future analyses at the inclusion and final visit. In this case, new approval by the regional health research ethics committee is required. Due to the time frame of the project, patients’ study-related travel expenses will be covered. Investigators will ensure that patients are covered by insurance via the hospital during study participation.

**Patient withdrawal**

Patients are free to withdraw from the trial at any time without providing reason(s) for withdrawal and without prejudice to further treatment. The reason for withdrawal may be withdrawal of consent, discontinuation of clozapine/olanzapine, failure to maintain a present level of adherence with the clinical trial medication, that is, missing more than three consecutive injections or more than 12 injections in total, adverse event(s), pregnancy discovered during the trial, significant worsening of mental health (defined as a Clinical Global Impression–Improvement score of 6 or 7) or an HbA1c higher than 57 mmol/mol (7.4%). No rescue medication will be offered in this study. Dropouts will be replaced until 52 patients have completed the treatment period. Data from dropouts will be included in the data processing. Patients withdrawing from the trial should be encouraged to undergo the same final evaluations as patients completing the trial. Completion or trial termination for any reason will be documented in the clinical record form pages. The supplier of pens will provide emergency sealed codes for ‘emergency breaking only’. In case of emergencies, it is possible for the investigators to reveal treatment based on study ID, that is, to collect information about the specific patient without breaking all randomisation codes.

**Intervention and choice of comparator**

Participants will be randomly assigned to semaglutide (semaglutide 1.34 mg/mL, 1.5 mL prefilled pen-injector) or placebo (no active drug) administered in the same way and volume. The trial medication is administered subcutaneously once a week in the entire treatment period; days 1–28: 0.25 mg semaglutide/placebo, days 29–56: 0.5 mg semaglutide/placebo and days 57–182: 1.0 mg semaglutide/placebo. Patients who, due to adverse events, do not tolerate up titration to 1.0 mg semaglutide will remain on 0.5 mg once a week. Patients who, due to adverse events, cannot tolerate dose escalation to 0.5 mg semaglutide/placebo will be excluded from the study. The patients will be instructed in the subcutaneous injection technique, which can be performed at home. If the patient cannot self-inject, a contact person will assist. Adherence and adverse events will be noted during the entire treatment period. For each patient treated, the batch number of the pen must be documented, and the patients will be asked to return the pens after usage. Patients will be asked to complete a weekly injection diary. If the patient accepts, they will receive a reminder of injections days and scheduled study visits. The importance of following study guidelines for adherence to weekly injections will be emphasised to all patients, and there will be a brief discussion of reasons for possibly missed doses and simple strategies for enhancing adherence. At the termination of study, levels of plasma-semaglutide will be assessed for adherence.

**Sample size**

The study is an explanatory study and the required patient population size is based on significance level (α) of 5%, a power (1–β) of 90%, where β (10%) is the risk of accepting a hypothesis that is false. Based on data from a recent study in similar patients,

\[ \text{Sample size} = \frac{2 \times (\text{SD})^2 \times (\alpha)^2 \times (1-\beta)^2}{(\text{effect size})^2} \]

we estimated that the minimum relevant difference (MIREDF) of HbA1c (primary endpoint) after 26 weeks of intervention would be −0.26% with an SD of 0.28%. With the aforementioned power, significance level, MIREDF and SD, the trial requires 26 patients in each of the two arms, that is, a total of 52 patients.
<table>
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<tr>
<th>Table 1</th>
<th>Treatment procedures</th>
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<td></td>
<td>Screening</td>
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<td>Introduction, informed consent, screening</td>
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<td>Height</td>
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<td>Hip and waist circumference</td>
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<td>Body weight</td>
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<td>Blood pressure and heart rate</td>
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<td>Clinical evaluation of suicide risk</td>
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<tr>
<td>Standard blood sampling and safety markers</td>
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<tr>
<td>Fasting blood and urine samples</td>
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<td>The Schizophrenia Quality of Life Scale</td>
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<td>Clinical Global Impression Severity</td>
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<td>Global Assessment of Psychosocial Disability</td>
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<td>Alcohol Use Disorders Identification Test</td>
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<td>Drug Use Disorders Identification Test</td>
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<td>Fagerström Test for Nicotine Dependence</td>
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<td>Clicker test</td>
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<td>Positive and Negative Syndrome Scale</td>
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<tr>
<td>DXA scanning</td>
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<td>Activity device distributed</td>
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<td>Activity device collected</td>
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<tr>
<td>Drug dispensing/returned</td>
<td>X</td>
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<tr>
<td>Adherence/practice study treatment self-administration</td>
<td>X</td>
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<tr>
<td>Adverse event assessment</td>
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<tr>
<td>Follow-up by phone call</td>
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DXA scan will be performed at baseline and at the termination of study participation. Activity measurements will be collected continuously during the first (week 1) and last week of participation (week 26) by using a wearable activity device\(^91\)\(^{-93}\) (table 1). At baseline and the last visit, potential effects on the reward value of a sweet–fat stimulus will be examined using a progressive ratio task ‘clicker test’\(^94\), a Positive and Negative Syndrome Scale–Six Items interview\(^95\) will be performed, and fasting blood and urine samples will be collected (box 3).

DXA, dual-energy X-ray absorption.
Thus, with an expected dropout of 50%, a total of 104 patients will be randomised.

Data management and statistical analysis

All data will be collected in electronic case report forms in REDCap hosted in the Capital Region of Denmark. The database will be locked after the last participants’ end of study. The sponsor–investigator has the final decision to terminate the trial. All statistical analyses except the post hoc sensitivity analyses will be performed with treatment groups still blinded. Before dividing participants into two groups, the statistical analysis plan will be uploaded at clinicaltrials.gov and the dataset will be locked. Statistical analyses will be performed using R (http://www.R-project.org/), with alpha set at 0.05. Statistical tests will be two-sided, and the level of significance will be 5%. All efficacy and safety analyses will be performed using a modified intention-to-treat principle in which all randomised participants who received at least one dose of the trial compound and have at least one assessment after baseline will be included. For the primary endpoint, repeated mixed-model analysis of covariance will be used to analyse change in the level of HbA1c from week 0 to week 26 for the semaglutide once a week and the placebo groups. All changes in secondary endpoints from baseline to the end of the trial will be analysed using repeated mixed-model analyses for continuous outcomes and mixed-model logistic regression for categorical outcomes. For comparison between the two groups, the covariates of age, sex, illness duration, treatment group, baseline CGILS score and BMI will be included in the analyses together with the baseline value of the relevant variable. Effect sizes (Cohen d) for lowering HbA1c levels and reducing body weight will be calculated by dividing the difference of the means in change from baseline to endpoint (treatment–placebo) by the pooled SD.

Public involvement and data availability

No members of the public or patients were involved in the research question, the design, the conduct of the research. Following publication, researchers who provide a methodological sound proposal can request access to the trial dataset.

ETHICS AND DISSEMINATION

We expect that the present study will generate important information about the long-term effects of continuous-acting GLP-1RA as an add-on therapy for dysglycaemic patients treated with clozapine and olanzapine. Patients are included in this study within the first 5 years of initiating clozapine or olanzapine treatment if they have developed pre-diabetes or diabetes. Clozapine-induced metabolic changes can be observed within the first years after initiation of treatment, which is why we are exploring an GLP-1RA as a choice for early treatment of antipsychotic-induced metabolic changes. This study is not considered as posing ethical problems. Semaglutide once a week is approved for the treatment of type 2 diabetes, obesity and overweight in the presence of weight-related comorbidity by EMA and by the FDA. The treatment in this study is associated with minimal discomfort for the participating patients. Procedures include blood sample collection, and a weekly injection of semaglutide in the subcutis of the abdomen, thigh, or upper arm. At the two DXA scans, participants will be exposed to modest radiation of app, 0.01 mSv, corresponding to two to three times the dosage of a dental X-ray. It is painless and no adverse effects are expected.

Investigators will be given access to the cleaned datasets. At the end of the trial, one or more manuscripts will be prepared for publication in international scientific journals. Data will be processed and merged into one or more scientific articles and published in accordance with the Consolidated Standards of Reporting Trials 2010 statement in international, peer-reviewed scientific journals and presented at national and international scientific meetings. We will seek to publish both positive and negative results. While it is the intention that the sponsor–investigator will be the last author, the published international guidelines for authorship (International Committee of Medical Journal Editors, 1997) will be adhered to.

Protocol, study approval and registration

The Standard Protocol Items: Recommendations for Interventional Trials (SPIRIT) guidelines were used as guidance for writing the protocol. The protocol and amendments are approved by the Danish Medicines Agency (EudraCT: 2020-000102-28) and the Regional Scientific Ethics Committee of the Capital Region of Denmark (committee C, #H-20019008). See online supplemental appendix A for further details. All data will be handled according to the General Data Protection Regulation. A detailed plan for curation has been approved by The Capital Region of Denmark. The investigators will ensure that the study will be carried out according to International Council for Harmonisation

Box 3 Blood and urine analyses

<table>
<thead>
<tr>
<th>Standard blood sampling</th>
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<tbody>
<tr>
<td>Glycated haemoglobin A1c, glucose, insulin, C-peptide, thrombocytes, alanine aminotransferase, aspartate aminotransferase, alkaline phosphatase and bilirubin.</td>
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<tr>
<th>Fasting blood sampling</th>
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<tbody>
<tr>
<td>Glucagon-like peptide 1, glucagon-like peptide 2, glucose-dependent insulinotropic polypeptide, glucagon, vitamin D, calcium, phosphate, magnesium, parathyroid hormone, procollagen type 1 N-terminal propeptide, C-terminal telopeptide of type 1 collagen, osteocalcin, low-density liprotein, high-density lipoprotein, triglycerides, cholesterol, amylase, creatinine, sodium, potassium, semaglutide and proteomic analyses.</td>
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<th>Fasting urine sampling</th>
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<td>DNA and RNA oxidative stress biomarkers.</td>
</tr>
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</table>
Good Clinical Practice (ICH-GCP) guidelines and the Helsinki Declaration and with national laws and regulations for clinical research. The study and a data sharing plan has been registered at ClinicalTrials.gov. In the process of ensuring data completeness and accuracy, source data verification will be performed by the GCP unit of Copenhagen University. The participants will be informed in writing about the possibility of audits and/or inspections. The audit and/or inspection might be performed by the hospital institutional review board/ethics committee or regulatory authority.

The status of the study is active recruitment of patients according to the approved protocol version, 9.0, 02.12.22.

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