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Head-to-head comparison of intensive lifestyle intervention (U-TURN) versus conventional multifactorial care in patients with type 2 diabetes: protocol and rationale for an assessor-blinded, parallel group and randomised trial

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ABSTRACT

Introduction: Current pharmacological therapies in patients with type 2 diabetes (T2D) are challenged by lack of sustainability and borderline firm evidence of real long-term health benefits. Accordingly, lifestyle intervention remains the corner stone in the management of T2D. However, there is a lack of knowledge regarding the optimal intervention programmes in T2D ensuring both compliance as well as long-term health outcomes. Our objective is to assess the effects of an intensive lifestyle intervention (the U-TURN intervention) on glycaemic control in patients with T2D. Our hypothesis is that intensive lifestyle changes are equally effective as standard diabetes care, including pharmacological treatment in maintaining glycaemic control (ie, glycated haemoglobin (HbA1c)) in patients with T2D. Furthermore, we expect that intensive lifestyle changes will decrease the need for antidiabetic medications.

Methods and analysis: The study is an assessor-blinded, parallel group and a 1-year randomised trial. The primary outcome is change in glycaemic control (HbA1c), with the key secondary outcome being reductions in antidiabetic medication. Participants will be patients with T2D (T2D duration <10 years) without complications who are randomised into an intensive lifestyle intervention (U-TURN) or a standard care intervention in a 2:1 fashion. Both groups will be exposed to the same standardised, blinded, target-driven pharmacological treatment and can thus maintain, increase, reduce or discontinue the pharmacological treatment. The decision is based on the standardised algorithm. The U-TURN intervention consists of increased training and basal physical activity level, and an antidiabetic diet including an intended weight loss. The standard care group as well as the U-TURN group is offered individual diabetes management counselling on top of the pharmacological treatment.

Ethics and dissemination: This study has been approved by the Scientific Ethical Committee at the Capital Region of Denmark (H-1–2014–114). Positive, negative or inconclusive findings will be disseminated in peer-reviewed journals, at national and international conferences.

Trial registration number: NCT02417012.

INTRODUCTION

The clinical care of type 2 diabetes (T2D) requires multifactorial intervention, including the pharmacological regulation of hyperglycaemia, hypertension and hyperlipidaemia to minimise T2D complications. The rationale for an increased risk of adverse medication effects, a decreased quality of life and economical costs. Thus, the strategies of lifestyle interventions are equally efficient in maintaining glycaemic control as the pharmacological treatment is well warranted.

Lifestyle changes, such as healthy diet and increased physical activity, are established cornerstones in diabetes management. In addition to the beneficial effects on glycaemic control, improved physical activity and training are also likely to improve mental and physical well-being as well as reducing stress and distress in adults. Only one study has studied the effects of lifestyle-driven weight loss in patients with T2D (the Look AHEAD trial). These reports indicate that short and long-term reductions in antidiabetic, lipid lowering and antihypertensive
medication can be achieved with a weight-loss intervention in patients with T2D. However, the weight loss was partially obtained by pharmacological treatment. Moreover, the pharmacological treatment was not standardised between the intervention and the control group, and the hypoglycaemic treatment was initially performed by the study physicians in the intervention group only. Accordingly, it is difficult to interpret to what extent healthy lifestyle changes (diet, aerobic and strength conditioning as well as decreased physical inactivity) per se can be used as a treatment for T2D as a substitute for the pharmacological treatment without compromising glycaemic control and metabolic health. Thus, studies of the effect of the U-TURN lifestyle intervention on the need for clinical T2D care alongside the effect on glycaemic control are needed in order to implement an intensive lifestyle treatment in clinical care.

A decade after the cessation of the predefined trial period in the UK Prospective Diabetes Study, significant reductions in the risk of myocardial infarction and improved mortality were observed in newly diagnosed patients with T2D allocated to early and aggressive glucose lowering treatment. It was then proposed that intensifying glycaemic control in patients with T2D with short T2D duration could be beneficial in reducing macrovascular and microvascular complications, whereas it had no effect or no adverse effects on patients with severe long-standing T2D. Thus, it could be speculated that an intensive lifestyle intervention could prove to be more efficient in patients with T2D with shorter disease duration.

Study objective and hypothesis
The objective of this study is to assess the clinical efficacy of the U-TURN lifestyle intervention in a sample of patients with short duration of T2D. We hypothesise that the U-TURN intervention would be comparable with the conventional multifactorial care in maintaining glycaemic control, while reducing the need for antidiabetic medications.

METHODS AND ANALYSIS

Trial design and study setting
The study is a parallel-arm, single-blinded, randomised clinical equivalence trial where the primary end point is glycated haemoglobin (HbA1c) monitored across 12 months. The participants are randomised in a 2:1 fashion to the lifestyle intervention (U-TURN) or standard care. The intervention is performed in a free-living environment with partial supervision of training and diet; all data collection will be performed at Copenhagen University Hospital at Rigshospitalet (primary trial sponsor, Blegdamsvej 9, 2100, Copenhagen, Denmark), and Glostrup (Denmark). Participants are recruited from the Capital Region of Denmark and the Region of Zealand, Denmark. The study has been registered at http://www.clinicaltrials.org (NCT02417012) on 14 April 2015. Amendments to the protocol are to be approved by the U-TURN steering committee and the Scientific Ethical Committee at the Capital Region of Denmark. Amendments are reported to http://www.clinicaltrials.org

Participants
Eligibility
Flow of participants is described in figure 1. Initially inclusion and exclusion criteria are identified through a phone interview (preinclusion). If eligible after the phone interview, the participant will be included after providing informed oral and written consent before any additional study procedures are initiated. Next step includes a blood sample (postinclusion) and a thorough medical screening (postinclusion). The latter two screenings are included in the procedure to identify latent exclusion criteria. Inclusion and exclusion criteria are described in box 1.

Interventions
The intervention and standard care are summarised in figure 2. The U-TURN lifestyle intervention is a 1-year intervention consisting of two main components (1 and 2) with four online supplementary intervention components (3–6):

1. Increased levels of structured and supervised training
2. Antidiabetic diet
3. Increased levels of basal physical activity
4. Increased sleep duration
5. Self-monitoring of behaviours related to components 1–4 as well as perceived stress level, mood and motivation
6. Diabetes management education and networking.

The U-TURN intervention is delivered in three phases (see figure 2). The standard care intervention group receives the standard treatment according to the Danish clinical diabetes guidelines. The pharmacological treatment is delivered by the study’s endocrinologist using the same titration and regulatory algorithms for both the groups. Both interventions and their rationale are described in detail below.

The U-TURN intervention

Intervention component 1: Increased levels of structured training

Current guidelines recommend that patients with T2D perform at least 150 min of moderate to vigorous intensity aerobic training for at least 3 days per week, with not more than two consecutive days between each training bout. In addition, resistance training is recommended three times per week at moderate to vigorous intensity. However, as evidence suggests, a greater reduction in HbA1c levels occurs with more than 150 min of structured training per week compared with 150 min of structured training per week or less. The training volume in the intervention will aim for 240–420 min training/
week. During phase 1 and 2 (figure 2), the participants will complete four aerobic training sessions per week of 45–60 min duration. Additionally, two combined training sessions consisting of aerobic and resistance training are included. The aerobic part of the training session will have a time duration of 30–35 min and the resistance part will be of 30 min duration. In phase 3, the participants will complete two aerobic training sessions and three combined training sessions. The duration span of the aerobic training will be maintained in phase 3. Once a week the training session will take place outside whereas all the other training sessions will take place in a fitness centre. The training will be structured, with supervision across the entire project period as evidence supports the beneficial effect of structured training on improving glycaemic control in patients with T2D. The supervision is reduced across the intervention period (figure 2) and is supported with online supervision of the participants’ self-reported and objective training data.

All training is performed in groups of 4–8 participants. The groups are composed based on the geographical location of the participants’ home address. Each group will be assigned at least two certified

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**Figure 1** Flow of participants through the U-TURN study.
coaches (minimum one physiotherapist) with one trainer being present at a supervised training session. Each week, a training programme is delivered from the intervention coordination centre to the coaches. An example of a weekly training programme sent to the coaches is provided in Table 1. This will contain the overall distribution of aerobic and resistance training as well as the detailed programming. In the programme, the aerobic training is described, including the duration and intensity. Furthermore, the resistance training is described with muscle groups, sets and repetitions so that participants from across all groups follow the same training programme. The training modality within the aerobic training (eg, power walking and cycling) and resistance training (eg, machines and bodyweight) is the only factor that may vary between groups. The modality is decided by the trainers in order to prevent and minimise the frequency and severity of injuries. No running is permitted during phase 1. An example of a training programme is presented in Table 1. Supervision is performed directly by the trainers during the training sessions, using heart rate measurements from the Polar V800 (Finland) and the online tool ‘Polar Flow for coach’ (Polar, Denmark). In phase 2 and 3, the heart rate of the participant and compliance to the unsupervised training session will be monitored online via the Polar V800 (Polar Inc, Denmark) and ‘Polar Flow for coach’.

### Intervention component 2: Antidiabetic diet, including an intended weight loss

The American Diabetes Association and the Canadian Diabetes Association support a macronutrient distribution within the range of 45–60% carbohydrate, 15–20% protein and 20–35% fat (<7% saturated fat). The U-TURN dietary intervention will be in line with these macronutrient distribution spans and will additionally focus on macronutrient quality: in particular, a diet with low glycaemic index (GI)/load (GL) as low GI or GL diets are related to a reduced HbA1c level, compared with high GI or GL diets, without inducing hypoglycaemia. As T2D is associated with comorbidities like cardiovascular disease and saturated fat intake is related to cardiovascular disease risk, the U-TURN intervention aims at reducing saturated fat intake to <7% as proposed by ADA. As successful management of T2D is highly related to diets rich in whole grains, fruits, vegetables and nuts and legumes and low on refined grains, red or processed meat and sugar sweetened beverages, focus on these items will be central part of the meal plans.

A clinical dietician will prepare individual meal plans and the implementation is continuously discussed during group sessions (same groups as the training groups) and during individual counselling (Figure 2). The meal plans will cover six daily meals (three main meals and three snack meals). Recipes will be changed continuously throughout the intervention. The principles of the meal plans by the dietician are described in Table 2.

![Figure 2](http://bmjopen.bmj.com/)

**Figure 2:** The participants’ actual body weight is used for calculation of the energy requirement if the body mass index (BMI) is <25 kg/m². If BMI >25 kg/m², the body weight in the equation is adjusted to equal a BMI=25 kg/m². The weight loss phase is discontinued immediately for all participants if the BMI becomes lower than 25 kg/m². At the individual counselling session primo phase 2 (Figure 2), the clinical dietician will decide in collaboration with the participant whether to initiate another weight loss period. If BMI is >30 kg/m² or waistline is >94 cm for men and >80 cm for women, the clinical dietician will recommend another weight loss period; otherwise, a maintenance period will be initiated. In the maintenance phase, the actual weight is applied in order to obtain energy balance. For all days, including structured training, 200 kcal/day will be added to the energy intake. In case of hypoglycaemic events, energy intake will be reassessed. In parallel to the training intervention, the clinical dietician will offer...
cooking classes and workshops on how to develop a meal plan and implement the plan. Participants are allowed to contact the clinical dietician by email once/week in case of any issues regarding implementation of or concerns about the meal plan.

To reduce the risk of hypoglycaemia, the participants are instructed to eat a snack meal just before (100–200 kcal) and after (200 kcal) a training session, and a main meal 2–3 h before a training session. In case of subjective signs of light hypoglycaemia (hunger, sweating, increased heart rate, feeling uncomfortable, dizziness and confusion), the participants are instructed to eat either one piece of fruit, drink a glass of juice in combination with a piece of rye bread or crisp bread.

**Intervention component 3: Increased levels of basal physical activity**

Physical inactivity and prolonged sedentary time comprises clinically important health risk factors. In a recent review it was concluded that minor increases in light intensity physical activity could improve glycaemic control in healthy persons and patients with T2D. As some evidence supports the beneficial effect of walking on glycaemic control in persons with T2D persons and as walking might prevent a deterioration in glycaemic control, U-TURN has adapted walking as a mode of physical activity to replace sedentary behaviour.

The aim is to reach an individual level of minimum 10 000 steps per day by gradually increasing the number of daily steps within the first month of intervention. Participants are encouraged to choose walking when possible and to incorporate light intensity physical activity breaks during prolonged sitting. Information to the participants about daily steps, the level of basal physical activity and sitting is provided using the Polar V800.

**Intervention component 4: Increased sleep duration**

It has been suggested that sleep duration is associated with improved glycaemic control in healthy persons and sleep deprivation reduces insulin sensitivity. Thus, in order to increase sleep duration, regular bedtimes and regular waking times are recommended throughout the week aiming at 7–8 h of sleep every night, with an additional requirement of 15–20 min in bed in order to fall asleep. All individuals will be recommended to shut down all electronic devices and dim the light at least 30 min before bedtime.

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**Figure 2** Description of the interventions and co-interventions. Participants are randomised either to intervention (U-TURN) (upper panel) or Standard care (Middle panel). Both groups receive pharmacological treatment and standard diabetes education (Lower panel—in grey). The intervention consists of three phases (1–3). The overall content in phases 1 through 3 is depicted in the green, light blue and light red boxes, respectively. HRR, heart rate reserve; Avg, average per training session.
| Week day | Aerobic training | Resistance training | Notes to the trainers |
|----------|------------------|---------------------|-----------------------|
| **Monday** | Duration: 40 min  
1. 5 min warm up at 60–65% of HR max  
2. 20 min at 70–78% of HR max  
3. 15 min at 76–83% of HR max | Duration: 30 min  
1. 5 primary target exercises: anterior chain (thigh), posterior chain (thigh), chest, back and shoulders  
2. Each target exercise is performed in three sets of 10–12 repetition max (RM)  
3. You can use machines, free weights, barbells, body weight, etc  
4. Active breaks containing core exercise are performed between each set. This means that the pause between each set is replaced with a core training exercise  
5. The five core exercises should include three dynamic abdominal exercises and two lower back exercises | Make sure to inform the participant which muscle groups are activated and with time expand their “box” of different exercises. This will help participants to increase variation in their exercise programmes and thus increase their motivation. Furthermore, it will also help to minimize the risk of injuries. |
| **Tuesday** | Duration: 60 min  
1. 5 min warm up at 60–65% of HR max  
2. 5 min at 70–75% of HR max  
3. 20 min at 74–79% of HR max  
4. 10 min at 80–88% of HR max  
5. 5 min at 70–75% of HR max  
6. 15 min consisting of 2 HR min at 76–80% of HR max, 2 min at 83–90% of HR max and 1 min active recovery. Repeat three times | | |
| **Wednesday** | Duration: 60 min  
1. 5 min warm up at 60–65% of HR max  
2. 10 min at 68–73% of HR max  
3. 15 min at 75–80% of HR max  
4. 10 min at 77–84% of HR max  
5. 10 min consisting of 30 s max effort and 30 s active recovery | Duration: 30 min  
1. 5 primary target exercises: Anterior chain (thigh), posterior chain (thigh), chest, back and shoulders  
2. Each target exercise is performed in three sets of 10–12 RM  
3. You can use machines, free weights, barbells, body weight, etc  
4. Active breaks containing core exercise are performed between each set. This means that the pause between each set is replaced with a core training exercise | |
| **Thursday** | Duration: 30 min  
1. 5 min warm up at 60–65% of HR max  
2. 25 min at 73–83% of HR max | Duration: 30 min  
1. 5 primary target exercises: Anterior chain (thigh), posterior chain (thigh), chest, back and shoulders  
2. Each target exercise is performed in three sets of 10–12 RM  
3. You can use machines, free weights, barbells, body weight, etc  
4. Active breaks containing core exercise are performed between each set. This means that the pause between each set is replaced with a core training exercise | |
Intervention component 5: Self-monitoring of behaviours related to components 1–4

Self-regulation theory posits that self-monitoring is a prerequisite to self-evaluation of progress made towards one’s goal and self-reinforcement for the progress made.31 Thus, the process of changing habits may require well-developed self-regulatory skills. Self-monitoring is central to this process and includes paying deliberate attention to one’s own actions as well as conditions under which these occur. In a review of 22 studies focusing on self-monitoring of diet, training or physical activity, Burke et al.32 found that more frequent self-monitoring was significantly and consistently associated with larger weight loss.

The U-TURN lifestyle intervention will entail a self-monitoring component, which is based on subjective evaluation on a daily basis. A simple questionnaire-containing eight inquiries regarding the intervention components and personal development is emailed to the participants every day. The participants rate the statements on a scale from 1 to 10 (1 is non-compliant and 10 is highly compliant). They rate the components of training (ability to follow training programme), daily physical activity (ability to comply with the physical activity goal—10 000 steps/day), diet (ability to follow the diet plan), sleep (sleep duration and sleep quality) and personal issues such as stress (perceived level), mood and general motivation. In case the participants’ score 1 (very low) in any of the items, the participants are asked to indicate the primary reasons for scoring so low. In case of a low score (figure 3) or when a participant repeatedly (three times/week) does not fill out the questionnaire, an email will be sent or a phone call will be made to the participant from the intervention coordination centre once a week. However, if the participant has contacted the coordination centre and informed them about circumstances which do not allow them to fill in the questionnaire (eg, vacation, work, etc) the follow-up procedure is not initiated. The dietician or training coach will follow-up with the participant directly based on the guidelines made by the intervention coordination centre. The self-monitoring serves two primary purposes: (1) to create a heightened awareness in each participant about their daily lifestyle choices in order to increase compliance with intervention guidelines, and (2) to prevent loss to follow-up. In addition, it offers the opportunity to adjust and individualise programmes within the overall framework for higher compliance and to prevent loss to follow-up.

Intervention component 6: Diabetes management education and networking

The literature supports individual33 and group-based diabetes counselling34 for improving glycaemic control, although the effect of group-based counselling may be more efficient.35 Thus, a group-based structure has been adapted. In addition, online encouragement and a participant platform for general experience sharing could potentially create a strong community feeling and enhance the face-to-face interaction.36

The U-TURN intervention participants are included in groups (see above). It also includes educational and informative elements, where the entire intervention group will participate in three 2 h lectures. The focus will be on disease pathology and diabetes management, diet and diet plans, training, sleep and motivational science. The participants will be assigned to a closed web-based group on http://www.facebook.com. It is essential to be aware of privacy settings; thus, U-TURN participants and health personnel will be requested to keep their engagement in the closed group and not ‘friend’ each other on

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Table 1 Continued

| Week day | Aerobic training | Resistance training | Notes to the trainers |
|----------|-----------------|-------------------|----------------------|
| Friday   | Duration: 60 min  
1. 5 min warm up at 60–65% of HR max  
2. 15 min at 73–83% of HR max  
40 min consisting of 5 min at 76–82% of HR max, 3 min towards max and 2 min active recovery. Repeat four times | Duration: 15 min  
1. Core training. Free of choice | 5. The five core exercises should include three dynamic abdominal exercises and two lower back exercises |
| Saturday | Rest day | Duration: 60 min  
1. 45 min walking  
2. 15 min walking/jogging uphill or on stairs | |
| Sunday   | Duration: 60 min  
1. 45 min walking  
2. 15 min walking/jogging uphill or on stairs | |

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Ried-Larsen M, et al. BMJ Open 2015;5:e009764. doi:10.1136/bmjopen-2015-009764
Participants allocated to the standard care group will receive the standard Danish T2D treatment. The medical regulation will be performed by two experienced endocrinologists. Owing to the blinding to group allocation, a study nurse is responsible for the main contact with the participants. Following the medical screening, but prior to the baseline measurements, all eligible participants will have their antidiabetic, lipid lowering and antihypertensive pharmacological treatment standardised using the predefined medications (figure 4A–C and table 4). The standardisation is employed in order to decrease the risk of reductions in medications not related directly to the intervention, but rather due to ongoing medical treatment at inclusion to the programme. The titration period is ≥6 weeks. Table 4 outlines treatment goals and intensification of treatment in U-TURN; while details of medication adjustments are outlined below and in figure 4A–C.

Procedures for regulation of medication (baseline to 12 months)
The study nurse presents the anonymised data to the blinded endocrinologists every third month and a decision on regulation of antidiabetic (baseline and every third month), cholesterol (baseline and every sixth month) and/or antihypertensive pharmacological treatment (baseline and every third month) is made (figure 4A–C). No information on group allocation is provided to the endocrinologists. The decisions will be based on HbA1c, cholesterol and home blood pressure measurements (18 home-based measurements over 3 days before each test round (Contour Next, Bayer, Copenhagen, Denmark)) using the algorithms described below. If insulin treatment is initiated, the antidiabetic pharmacological treatment is adjusted based on home glucose monitoring every 2–4 weeks (see below). Also, in case of glucagon-like peptide-1 analogue (GLP-1 analogue) or insulin treatment, an information meeting is arranged with the study nurse to educate the participant in glucose monitoring and insulin injection technique. If the treatment target is

| Table 2 Principles of the U-TURN meal plan |
|-------------------------------------------|
| Principle                                    | Additional comment |
| Homemade food                                | Recipes are included |
| Limit processed food items                  | 200 g should be ‘fat’ fish, for example, salmon or mackerel |
| Include seasonal greens and fruits (minimum 600 g/day) | |
| Maximum two pieces of fruits per day         | |
| Limit the amount of sodium                  | |
| Include fish (350 g/week)                   | |
| Fibre rich food items (3 g/MJ)              | |
| Hot meals should include fish once per week, one vegan meal per week Minced meat maximum twice per week Organic food items | |
| Hot meals should contain minimum 200 g vegetables per meal, maximum one-fourth of the plate should be meat, maximum one-fourth of the plate should be high glycaemic index/load food items Ad libitum intake of water and tea is allowed Maximum two cups of coffee/day No sugar sweetened beverages (including soda pops, juice or artificial sweetened beverages) Alcohol is discouraged throughout the intervention period | Not a demand—but participants are encouraged to use organic food items |
| Juice is allowed in case of subjective signs of hypoglycaemia in relation to training (see below) | |

Facebook. The intervention coordinators, diet and training councillors will post positive encouragements or encourage sharing success, fear, hope, etc. The diet and training councillors will monitor the discussions and add relevant inputs to the discussions.

Preventing discontinuation: The U-TURN Toolbox
Several rescue mechanisms are implemented to prevent loss to follow-up. The first line of action in case of problems with adherence relates to self-monitoring (see above). An extended toolbox contains procedures to modify the intervention components 1 (training) and 2 (diet). These procedures are described in table 3.

Conventional multicomponent care
Participants allocated to the standard care group will receive the standard Danish T2D treatment.33 Briefly, in Denmark patients with T2D are stratified based on the capability and severity of their condition determined by the patient’s general practitioner (GP) to receive varying levels of rehabilitation and clinical care.37 The GP will cover the coordination with rehabilitation programmes, foot and eye specialists. However, the pharmacological treatment will be administered solely by the U-TURN endocrinologists (see description below). Furthermore, the participants receiving standard care will be interviewed 2 weeks postrandomisation about concerns regarding the allocation and will be provided with valid arguments to complete the project. The participants can contact a diabetes nurse by phone and/or email throughout the entire intervention period to discuss their concerns regarding the treatment.

Cointerventions (U-TURN and standard care)
Medical treatment in U-TURN
The medical regulation will be performed by two experienced endocrinologists. Owing to the blinding to group allocation, a study nurse is responsible for the main contact with the participants. Following the medical screening, but prior to the baseline measurements, all eligible participants will have their antidiabetic, lipid lowering and antihypertensive pharmacological treatment standardised using the predefined medications (figure 4A–C and table 4). The standardisation is employed in order to decrease the risk of reductions in medications not related directly to the intervention, but rather due to ongoing medical treatment at inclusion to the programme. The titration period is ≥6 weeks. Table 4 outlines treatment goals and intensification of treatment in U-TURN; while details of medication adjustments are outlined below and in figure 4A–C.

Procedures for regulation of medication (baseline to 12 months)
The study nurse presents the anonymised data to the blinded endocrinologists every third month and a decision on regulation of antidiabetic (baseline and every third month), cholesterol (baseline and every sixth month) and/or antihypertensive pharmacological treatment (baseline and every third month) is made (figure 4A–C). No information on group allocation is provided to the endocrinologists. The decisions will be based on HbA1c, cholesterol and home blood pressure measurements (18 home-based measurements over 3 days before each test round (Contour Next, Bayer, Copenhagen, Denmark)) using the algorithms described below. If insulin treatment is initiated, the antidiabetic pharmacological treatment is adjusted based on home glucose monitoring every 2–4 weeks (see below). Also, in case of glucagon-like peptide-1 analogue (GLP-1 analogue) or insulin treatment, an information meeting is arranged with the study nurse to educate the participant in glucose monitoring and insulin injection technique. If the treatment target is
reached, the dose of the compound is halved at the following control time point (3 months later). In case of unchanged values or an additional drop in level, the compound is discontinued.

**Regulation of medication (12–24 months)**

All pharmacological treatment for the participants will be performed by their usual GP.

**Medical regulation algorithms**

**Algorithms for regulation of antidiabetic medication**

It is well recognised that tight glycaemic control in fragile diabetes patients (advanced age, known history of severe hypoglycaemia, overt cardiovascular disease) might be associated with higher incidence of cardiovascular mortality. Patients eligible for inclusion in the study are without known history of severe hypoglycaemia.

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**Figure 3** One electronic inquiry with eight (1–8 in figure) sub-inquiries is administrated to the participants’ intervention (U-TURN) and rated on a daily basis. The participants rate the inquiry from 1 (worst) to 10 (best). If rating is one, the participants are asked for the primary reason. Based on the frequency of the reasons action is taken. The answers are reviewed by the intervention coordination centre on a weekly basis. Based on ratings and frequency of reasons actions (red boxes) are taken.
GLP-1 analogues exhibit several other positive effects, compared with sulfonylurea, the treatment. Despite an eventual less beneficial effect on glycaemic control in comparison with metformin alone, whereas the training intensity will be maintained unsatisfactory glycaemic control on metformin alone, a GLP-1-analogue (Victoza) is added as second-line treatment due to its long-standing evidence base for efficacy compared to the addition of prandial insulin with a subsequent weight loss and less hypoglycaemia. Eventually, if HbA1c is still above the target while fasting blood glucose is below 7 mmol/L, metformin, fast acting Novorapid, is initiated.

### Algorithm for regulation of lipid lowering and antihypertensive medication

Clinical studies in T2D have shown beneficial effects on cardiovascular outcomes from lowering of blood pressure to at least below 140 mm Hg systolic and 85 mm Hg diastolic. Therapeutic goal for blood pressure during the U-TURN study is 130/80 mm Hg. An angiotensin receptor blocker (Losartan) is chosen as first-line antihypertensive treatment due to its additional protective effect on kidney function in patients with T2D. If the patients exhibit unsatisfactory blood pressure control from Losartan alone, a thiazide (Gentiptyl) is added as second-line treatment based on its well validated and long-standing evidence as an efficient diuretic with an effect on cardiovascular outcomes. As a third and eventually fourth antihypertensive drug in the treatment algorithm, a calcium channel blocker (amlodipine) and a mineralocorticoid antagonist (spironolactone) is chosen.

The increased prevalence of lipid abnormalities in patients with T2D and the associated risk of cardiovascular disease make lipid lowering treatment necessary. Therapeutic goals for lipids in the U-TURN study are low-density lipid cholesterol below 2.5 mmol/L and triglycerides below 5 mmol/L. Statin medication is preferred therapy as it helps to decrease all-cause and vascular mortality among diabetic patients. In the U-TURN study, two different kinds of statins are used, with atorvastatin replacing simvastatin in case of insufficient effect.

### Safety criteria and adverse events

Participants will be informed about side effects as well as subjective signs of hypoglycaemia (hunger, sweating, increased heart rate, feeling uncomfortable, dizziness and confusion) and hyperglycaemia (thirst, polyuria, polydipsia, pruritis, increased weight). Hypoglycaemia, advanced atherosclerosis, severe comorbidity or advanced age which allows for tighter glycaemic control in U-TURN. The goal is an HbA1c of 48 mmol/mol (6.5%) giving due consideration to the risk of hypoglycaemia in the individual patient.

Biguanid (tablet Metformin) is initiated as first-line treatment due to its long-standing evidence base for efficacy and safety; it is inexpensive, and may reduce the risk of cardiovascular events. If the patient exhibits unsatisfactory glycaemic control on metformin alone, GLP-1-analogue (Victoza) is added as second-line treatment. Despite an eventual less beneficial effect on glycaemic control in comparison with sulfonylurea, the GLP-1 analogues exhibit several other positive effects, including less frequent hypoglycaemia and weight loss.

In case the patients experience unacceptable side effects on GLP-1 analogue treatment (eg, nausea), the treatment can be changed to metformin in combination with a dipeptidyl peptidase inhibitor (DPP-4 inhibitor, Januvia). However, with time supplementation with subcutaneous injections of insulin or insulin analogues is often necessary in order to compensate for insulin deficiency. As third-line treatment, one daily injection of insulin glargine biosimilar (Abasaglar initiating dose 0.2 U/kg/day) is, therefore, eventually added and titrated to an acceptable fasting blood glucose level. Further, analyses between these insulin doses have documented significant reductions in hypoglycaemia mainly at night with the insulin analogues, while no statistically significant differences for severe hypoglycaemia rates were shown in any of the trials. Most studies have shown that the combination of GLP-1 receptor agonists with basal insulin has an equal or slightly superior efficacy compared to the addition of prandial insulin with a subsequent weight loss and less hypoglycaemia. Eventually, if HbA1c is still above the target while fasting blood glucose is below 7 mmol/L, then meal insulin, fast acting Novorapid, is initiated.

| Table 3 | Extended tool box for prevention of loss to follow-up |
|---------|------------------------------------------------------|
| **Intervention component 1 (training)** | **If the participant contacts the therapist in person or by email and express concerns about participation in the training intervention** |
| **Action 1** | The participant is offered a motivational interview with the coordination centre to get an overview over the possible challenges, that is, lack of time or worries. An adjusted plan is made and the trainers will follow-up at the supervised training. If the lacking compliance relates to injuries, pain or resistance to training modality, the training modality may be altered, whereas the training intensity will be maintained if action 1 is insufficient, the participant is invited to a personal motivational interview with a motivational expert not involved with the daily training |
| **Action 2** | If action 1 and 2 are insufficient, two training sessions per week are eliminated from the programme for 4 weeks. The training session will be gradually reintroduced |
| **Intervention component 2 (diet)** | **If the participant contacts the dietician or at group counselling** |
| **Action 1** | Participants are interviewed regarding compliance to the meal plan and provided with specific guidelines to practical changes in the plan by the clinical dieticians. For example, to increase adherence to food items increasing satiety or exchange some food items to match preferences |
| **Action 2** | If action 1 is insufficient and the participant still experience lack of satiety, then the energy intake is increased in steps of 100 kcal/day until the level of satiety is acceptable by the participant. The process is performed via email with the dietician |
fatigue and confusion), and urged to contact the study nurse in case of any adverse symptoms. Severe hypoglycaemic events (see below) will be registered by the study nurse. The safety criteria employed include adverse events, health-related outcomes (for instance, episodes of angina or signs of atrial fibrillation) and subject-reported hypoglycaemic episodes (plasma glucose <4 mmol/L). Non-severe hypoglycaemic events are defined as those that can be self-treated; severe hypoglycaemic events are defined as plasma glucose <3 mmol/L or episodes requiring third-party assistance or medical intervention. In case of adverse effects, medication is changed according to the titration described. In case of hypoglycaemic episodes, antidiabetic medication is eventually adjusted. Severe hypoglycaemic periods are reported to the study nurse. The hypoglycaemic events

Figure 4 (A) Illustration of antidiabetic treatment algorithm: Biguanid (tablet Metformin) is initiated at 500 mg once daily up to 1000 mg twice daily. If treatment goal is not reached, then a GLP-1 analogue (injection Victoza) is added at 1.2 mg increasing to 1.8 mg daily. In case of unacceptable adverse effects, a dipeptidyl peptidase inhibitor-4 inhibitor (tablet Januvia) is used at 100 mg daily instead of the GLP-1 analogue. If treatment goal is not reached, then basal insulin (injection Abasaglar) is added (0.2 units/kg once daily). If treatment goal is not reached then meal insulin is added (injection Novorapid titrated based on self-assessed pre-prandial blood glucose measurements in close cooperation with the study nurse). Detailed insulin adjustment is included in the online supplementary material. (B) Illustration of antihypertensive treatment algorithm: An angiotensin II receptor antagonist (tablet Losartan) is initiated at 50 mg daily up to 100 mg daily. If treatment goal is not reached, then a thiazide (tablet Centyl cum KCL) is added at 2.5 mg increasing to 5 mg daily. If treatment goal is not reached, then a calcium antagonist (tablet Amlodipine) is added at 5 mg increasing to 10 mg daily. In case of unacceptable adverse effects, a mineralocorticoid (tablet Spironolactone) is used at 25 mg increasing to 100 mg daily. (C) Illustration of lipid lowering treatment algorithm: A statin (tablet Simvastatin) is initiated at 40 mg daily. If treatment goal is not reached, treatment is replaced by another statin (tablet Atorvastatin) at 10 mg increasing to 40 mg daily.
are then presented to the endocrinologist and registered in a database. At any time, all necessary information, including information about intervention, medical history and adverse events, on the individual participant is available, but this is blinded to the endocrinologist in order to maintain group concealment. If considered necessary, the blinding will be revealed on a patient-to-patient basis and the participants will be contacted directly by the endocrinologist. This will be decided on a patient-to-patient basis and will be based on information provided by the study nurse. The participant’s GP will be informed about the procedure and encouraged to contact the U-TURN project nurse in case of questions.

Injuries related to the intervention (acute and over-use) will be registered if reported. In case of reports of severe adverse events during the study period, the steering committee will be informed as will the Scientific Ethical Committee of the Capital Region of Denmark.

Diabetes education
All participants are invited to individual educational meetings and diabetes controls (30 min) with a trained diabetes nurse every third month (a total of four meetings). Home blood pressure and home glucose measurements are reviewed. At these meetings, challenges regarding the diabetes treatment, including compliance issues, are addressed. Moreover, general education regarding the importance of a healthy lifestyle is provided. Furthermore, all participants are invited to an introductory 2 h diabetes management course.

Treatment of obstructive sleep apnoea
As the prevalence of obstructive sleep apnoea is increased in patients with T2D and maybe casually linked to T2D, a screening of all participants is performed. Participants diagnosed with sleep apnoea (Apnoea-Hypopnoea Index >15 h) following baseline sleep testing with cardiorespiratory monitoring (see below) are offered sleep apnoea treatment (continuous positive airway pressure (CPAP)). To minimise the effect of CPAP treatment on retesting results, the participants with obstructive sleep apnoea are told to discontinue the CPAP treatment 1–2 days before retesting.

Retention
All participants will receive €300 (2250 Danish kroner) to cover lost earnings, transport and discomfort in relation to the testing procedures. To minimise loss to follow-up in the standard care group, participants are interviewed 2 weeks post allocation by the study nurse regarding their concerns about allocations to standard care. Furthermore, they are offered an interview about their progress after 1 year. All participants are allowed to contact the U-TURN study nurse by phone in case of study-related questions (eg, pharmacological treatment, sports injuries, etc).

Study end points and assessments
Table 5 describes the time points at which the outcomes are assessed during the intervention and follow-up period.

Primary outcome
The primary outcome measure is change in glycaemic control (HbA1c) from baseline to 12-month follow-up.

Secondary outcomes
The key secondary end point includes reductions in anti-diabetic medications from baseline to 12-month follow-up. For exploratory purposes, the change in anti-diabetic medication is also quantified according to dose from baseline to 12-month follow-up. Every available dose will be graded according to the titration algorithms described above and summed into a total dose score at 12-month follow-up. Every step in the titration (figure 4A) is awarded one point. A point is either added to the score when medication is added on (ie, increasing one step up on figure 4A) or subtracted upon discontinuation of medication (ie, if declining one step on figure 4A). A total of eight points can be accumulated if the participant is receiving full dose (ie, ends at step 8 figure 4A at 12-month follow-up) or 0 if no antihyperglycaemic medication is prescribed. Changes from the baseline score are reported. All secondary outcomes are assessed until 12-month follow-up except for changes in body composition, and personal and demographic variables which are also monitored until 24-month follow-up.

Table 4 Treatment goals for medical regulation

| Medication | Treatment goals | Intensification of treatment during standardisation | Intensification of treatment at follow-ups |
|------------|----------------|---------------------------------------------------|------------------------------------------|
| Antidiabetics | HbA1c ≤48 mol/mol | HbA1c >64 mmol/L or 5 mmol/mol increment | HbA1c >58 mmol/L or 5 mmol/mol increment |
| Antihypertensive | BP ≤130/80 mm Hg | BP >150/95 mm Hg | BP >140/85 mm Hg |
| Antilipids | LDL ≤2.0 mmol/L | LDL >2.0 mmol/L | LDL >2.0 mmol/L |
| | TG ≤5.0 mmol/L | TG >5.0 mmol/L | TG >5.0 mmol/L |

The table shows treatment goals for the U-TURN intervention and intensification of treatment. If the treatment target is reached, the dose of the compound is halved at the following control time point (3 months later). In case of unchanged values or an additional drop, the compound is then discontinued.

BP, blood pressure; HbA1c, glycated haemoglobin; LDL, low-density lipoproteins; TG, triglycerides.
| Measurement                      | Description                                                                 | Baseline | 3 months | 6 months | 9 months | 12 months | 24 months |
|----------------------------------|-----------------------------------------------------------------------------|----------|----------|----------|----------|-----------|-----------|
| Blood sampling                   | Following an overnight fast (8 h) a blood sample is drawn. The plasma markers of metabolism (HbA1c*,†‡, TC*, LDL*, HDL*, TG†, fasting insulin† and glucose†) will be analysed |          | V        | V        | V        | V         | –         |
| Medications                      | Dose of antidiabetic*, lipid* and blood pressure lowering* medications       |          | V        | V        | V        | V         | –         |
| Adverse events                   | Major hypoglycaemic episodes*, cardiovascular events*, acute* and overuse* injuries related to the intervention component 1 |          | V        | –        | –        | V         | –         |
| Glucose tolerance                | Following an overnight fast (8 h) and after a 48 h medication and training pause, an antecubital intravenous line is placed and a standard 75 g oral glucose tolerance test will be performed. Blood will be drawn at the following time points: 0 (baseline), 15, 30, 60, 90 and 120 min. The plasma will be analysed for insulin†, C-peptide† and glucose†. Area under the curve will be analysed†‡ |          | V        | –        | –        | –         | V         |
| Physical fitness                 | Fitness† will be assessed by employing a progressive bicycle ergometer test protocol. Oxygen consumption will be assessed using continuous indirect calorimetric measurements (Cosmed, Italy) |          | V        | –        | –        | –         | V         |
| Cognitive testing                | Specific cognitive function areas (short-term memory, attention, executive functions, etc) will be tested using the CANTAB test package from Cambridge Cognition (Cambridge Cognition, UK) and BDNF-α is assessed from blood serum |          | V        | –        | –        | –         | V         |
| Depression                       | Characteristics, attitudes and symptoms of depression will be assessed using the Beck Depression Inventory51 |          | V        | –        | –        | –         | V         |
| Well-being, functional ability and motivation | Functional health and well-being will be assessed using the SF-36.52 Positive and negative effect on separate subscales will be |          | V        | V        | –        | V         | –         |
| Measurement                                | Description                                                                 | Baseline | 3 months | 6 months | 9 months | 12 months | 24 months |
|-------------------------------------------|-----------------------------------------------------------------------------|----------|----------|----------|----------|-----------|-----------|
| Dietary intake                            | A food frequency questionnaire is completed*53                             | V        | –        | –        | –        | V         | –         |
| Body composition, anthropometry and blood pressure | Height**§, weight**§, waist and hip circumference will be measured by standard procedures. Dual X-ray absorptiometry (iDXA; Lunar, Madison, WI) and COREScan will be used to assess whole body composition†. Home-based diastolic* and systolic* blood pressure is assessed using an upper arm blood pressure monitor (Model BPM1C, Kinetik, UK) | V        | V        | V        | V        | V         | V         |
| Personal information                      | Age**§, sex**§, diabetes duration**§ and educational level**§ will be obtained using self-report               | V        | –        | –        | –        | –         | –         |
| Physical activity                         | Information on the physical activity level† will be obtained using questionnaire RPAQ                              | V        | –        | –        | –        | V         | –         |
| Personality type                          | Two questionnaires regarding personality, the NEO-Five Factor Inventory and a SES are incorporated¶54 55                | V        | –        | –        | –        | V         | –         |
| Sleep quality, disturbances, fatigue and sleepiness | Cardiorespiratory monitoring is used to measure the prevalence of obstructive sleep apnoea†. Participants complete sleep diaries for 2 weeks after each cardiorespiratory monitoring in order to record and describe potential changes in their sleep. The ESS† and the MFI† are employed to measure daytime sleepiness. Sleep quality is measured using the PSQI†           | V        | –        | V        | –        | V         | V         |
| Arterial function¶                        | An ultrasound system equipped with vascular software for two-dimensional imaging, colour and spectral                  | V        | –        | –        | –        | V         | –         |

*53 Ried-Larsen M, et al. *BMJ Open* 2015;5:e009764. doi:10.1136/bmjopen-2015-009764

**§ Ried-Larsen M, et al. *BMJ Open* 2015;5:e009764. doi:10.1136/bmjopen-2015-009764

† Ried-Larsen M, et al. *BMJ Open* 2015;5:e009764. doi:10.1136/bmjopen-2015-009764

¶ Ried-Larsen M, et al. *BMJ Open* 2015;5:e009764. doi:10.1136/bmjopen-2015-009764
Sample size considerations

The sample size in this study was based on what was considered feasible, within the local context, enabling up to 120 participants to be enrolled in the trial period (29 April 2015–17 August 2017). The sample size is truncated at 120 participants or the N reached at the end of recruitment period—whichever is reached first. To increase the sensitivity to the U-TURN intervention it was decided to randomise the participants in a 2:1 fashion.

The study was not formally powered as an equivalence trial, but from the content experts it was decided that a reasonable equivalence margin would be ±0.4%-points for HbA1c for the between-group comparison. As presented in figure 5, assuming that the HbA1c is down to 6.5% in both groups, with an SD of 0.9%, we estimated that enrolling 120 participants in the intention-to-treat population (ITT; 80:40), testing a 2-tailed superiority hypothesis (based on 95% CIs) would be reasonably precise to estimate within a reasonable equivalence margin; −0.34% to 0.34%). Further, according to the principle of sensitivity, our estimates support that even if we include only 90 (60:30) participants our confidence limits will be acceptable.

Randomisation, sequence generation and allocation concealment

Participants will be assigned randomly (2:1) in permuted blocks of three and six, according to computer-generated random numbers, to undergo either U-TURN or standard care after the baseline measurements. Participants will be stratified according to sex (male vs female). The sequence is generated centrally by a researcher not involved in the testing or allocation, and delivered to the data manager. The allocation information to the participants is given by a study nurse not involved in the testing, randomization or evaluation procedures. The study nurse receives the participant allocation directly from the data manager and the group allocation is delivered to the participants. The data will not be accessible until completion of the 12-month follow-up data collection.

Table 5

| Measurement | Description | Baseline | 3 months | 6 months | 9 months | 12 months | 24 months |
|-------------|-------------|----------|----------|----------|----------|-----------|-----------|
| Doppler is used to assess flow mediated dilation and shear stress in the femoral artery§ |

*Article 1: The effects of a head-to-head comparison of intensive life style intervention (U-TURN) versus standard glucose lowering medications in patients with type 2 diabetes mellitus: an assessor-blinded, parallel group, randomised trial.
†Article 2: The effects of the U-TURN lifestyle intervention on sleep quality and sleep apnoea in patients with type 2 diabetes.
‡Article 3: Predictors for improvements in glycemic control after a comprehensive lifestyle intervention; A sub-study to the randomised trial study U-TURN.
§Article 4: Effect of lifestyle intervention on endothelial function in patients with type 2 diabetes assessed by flow mediated dilatation; A sub-study to the randomised trial U-TURN.
¶Measured in a subset of the sample (N=40).
–, not assessed; BREQ-2, Behavioral Regulation In Exercise Questionnaire 2; ESS, Epworth Sleepiness Scale; GMS, Global Mood Scale; HbA1c, glycated haemoglobin; HDL, high-density lipoprotein; LDL, low-density lipoprotein; MFI, Multidimensional Fatigue Inventory; PSQI, Pittsburgh Sleep Quality Index; RPAQ, Recent physical activity questionnaire; SES, Sensation Seeking Scale; TC, total cholesterol; TG, triglyceride; V, assessed.

Figure 5

Forrest plot depicting three scenarios of total sample size from a total of 90 participants (scenario 1), 105 (scenario 2) and 120 participants (scenario 3) with the respective with of 95% CIs. HbA1c, glycated haemoglobin N, number; MD, mean difference.
Blinding
Outcome assessors as well as the endocrinologists are blinded to the allocation at baseline and follow-up. The participants will be informed that they are not allowed to discuss their allocation during the follow-up measurements. Owing to the nature of the trial, participants, diabetes nurse and intervention coordinators cannot be blinded to the allocation.

Statistical methods
All data being collected longitudinally (including the primary outcome) will be analysed according to the ITT principle using repeated-measures analysis of covariance applied in mixed linear models. Patterns of missing data will be investigated. A priori, the less restrictive missing at random (MAR) assumption is considered more reasonable than the missing data being missing completely at random; MAR assumes that drop-out may depend on observed outcomes or covariates, but does not depend on unobserved data. The prespecified efficacy analyses are based on the full-analysis set, which include all participants who are randomised and have had an outcome assessment at baseline. Under the MAR assumption, likelihood-based approaches, such as mixed effects models, will produce valid inferences. Thus, based on mixed linear models, our primary model is based on analysis of covariance for continuous end points. The model will include group and sex as fixed effects, with the baseline value of the relevant variable as a covariate. Categorical data for dichotomous end points will be analysed with the use of logistic regression with the same fixed effects and covariates as the respective analysis of covariance.

Assuming that the data on potential drop-outs are MAR; both linear mixed models and multiple imputation procedures would be applicable to handle missing data.

While the null hypothesis in a superiority trial is that treatment effects are identical (with $H_A$ implying that these are not), the null hypothesis for the primary outcome in this trial is defined with reference to an acceptable clinical difference in treatment effects (ie, equivalence margin). As mentioned above, we define the threshold for not ‘too different’ or not ‘unacceptably worse’ as a potential difference (95% CI) in HbA1c of±0.4% points. Sensitivity analyses will be performed to assess robustness of the primary analyses, including repeated measures with baseline observation carried forward, and multiple-imputation techniques (with the latter using model based approaches to ‘replace’ missing data).

Exploratory analyses of the treatment effects will be performed on the secondary outcomes. The analysis plan will be developed prior to data analysis, and will be performed in parallel independently by two blinded researchers. Group allocation is not disclosed before consensus about the interpretation of the data is reached. In essence, the information about treatment and the N is concealed until consensus is reached. Discrepancies in the analysis outcome between the researchers will be resolved using a blinded third party statistician.

ETHICS AND DISSEMINATION
The U-TURN study is expected to provide evidence that lifestyle change (without use of weight loss enhancing pharmacological additives as a part of the treatment) is equally effective in treating T2D as the recommended multicomponent care. Furthermore, the study will provide valuable insights into motivational mediators and moderators for adherence to intensive lifestyle treatment. Although intensive, the intervention contains components which can be directly included in the rehabilitation of the patients with T2D and thus, it might have a broad appeal. If the intervention is effective in maintaining glycaemic control and reduces the need for antidiabetic pharmacological care, it will provide an alternative to the current standard care and thus decrease some of the side effects induced by the medications used in standard clinical care.

The study is unique as the methodology is stringent and systematic, thus providing scientifically viable data. A major strength of the study is the blinding of assessors and participants at the baseline assessment, and the assessor and endocrinologist blinding at all follow-up assessments thereby limiting the risk of bias. It is not possible to blind the participants during follow-up in this type of trial, which could introduce a bias. The inclusion of predefined medications used in the pharmacological care (ie, not all combinations and medications are considered) limits the interpretation to patients receiving these combinations. As the choice of medications in the study is in line with the recommended first line-treatments, the observations made in this study will still apply to a large number of patients. Furthermore, the inclusion of self-reported physical activity and adherence pose a potential information bias, limiting the interpretation of the effects of the single intervention components on HbA1c.

The U-TURN study is initiated with a 6-week prerandomisation open label run-in period for all participants, in which the medical treatment is adjusted according to the treatment algorithm used in the study. This is done in order to adjust for differences in presstudy medication due to either non-compliance or insufficient prescriptions (too little or too much medication) and to have a uniform status of medication at baseline. A titration period of 6 weeks is chosen to ensure steady state of new prescribed medication before allocation. In order to dissect the combined effect of medication and intervention in contrast to medication alone, we aim for homogenous pharmacological treatment and reliable estimates of adherence. This is done by a clear pharmacoeutical treatment algorithm and administering of an interview every third month focusing on adherence to medication. The study is designed as a treat-to-target
study aiming at specific goals for glycaemic control, blood pressure and lipids. This implies intensification or decrease in medical treatment according to clinical data (HbA1c, glucose measurements, blood pressure and lipids). We are aware that regulation of medication prior to allocation and during the study period may have a ‘carry-over effect’ affecting the following measurements. Hence, a 3-month period between follow-ups is chosen as this should be sufficient time to obtain sustained effect of medical regulation. Also tapering of medication is done gradually to ensure that any improvement in clinical outcome is sustained even after withdrawal of medication. The study will be conducted according to the principles of the Helsinki Declaration II. Prior to inclusion, all participants received written and oral information about the study and provided written as well as oral informed consent. The participants can discontinue participation in the full study or part of the study at all times with no obligation to provide a reason. This will not have any consequences for their future treatment. If discontinued, the participants will receive the standard T2D care according to the Danish clinical guidelines; however, this will be delivered by their own GP. Medical regulation will be performed by experienced endocrinologists throughout the study in accordance with the predefined algorithms and strict safety criteria, and registration of adverse outcomes is employed. Thus, the study is expected to result in limited risks, adverse effects and discomfort to the participants. The results are to be reported according to the CONSORT guidelines. Negative, positive and inconclusive results will be disseminated in international peer-reviewed scientific journals at national and international conferences. Access to data can be granted on approval for a formal request to the steering committee.

All participants will be provided with their results. Postrandomisation, all participants are ascribed a unique participant identifying information is stored alongside study data. A transfer key (ID to personal information) is available, but is encrypted and stored separately from the trial data. After trial cession all data are anonymised and stored on the server of Copenhagen University Hospital. Additional biological materials are stored in a research bio-bank for up to 20 years.

As far as we are aware, the U-TURN study is the first study to investigate if lifestyle intervention is effective in maintaining glycaemic control while aiming at discontinuing pharmacological care in patients with T2D. The results from this trial are of great importance for clinical care of T2D and can provide important knowledge on how to implement and conduct treatments using the lifestyle changes.

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