





Clinical Study Protocol

Study Title: An open-label, two-centre, randomised, 2-period cross-over study to assess the efficacy, safety and utility of fully closed-loop insulin delivery in comparison with standard care, in adults with type 2 diabetes requiring maintenance dialysis.

Short Title: Closed-loop in adults with T2D requiring dialysis (AP-Renal)

Protocol Version: 1.1 12 June 2019

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This protocol has been written in accordance with current ISO 14155:2011 standard

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PROTOCOL SIGNATURE PAGE

The signature below documents the approval of the protocol entitled "An open-label,
two-centre, randomised, 2-period cross-over study to assess the efficacy, safety
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adults with type 2 diabetes requiring maintenance dialysis." <code>Version</code> $_$. $_$ dated $_$ $_$ / $_$
$_{\rm -}/_{\rm -}$ and provides the necessary assurances that this study will be conducted
according to all stipulations of the protocol, the principles of GCP and the appropriate
reporting requirements.
Signature Date

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SITE SIGNATURE PAGE

I have read the attached protocol entitled " An open-label, two-centre, randomised, 2-period cross-over study to assess the efficacy, safety and utility of fully closed-loop insulin delivery in comparison with standard care, in adults with type 2 diabetes requiring maintenance dialysis." Version ___ dated _ _/_ _/_ _, and agree to abide by all provisions set forth therein.

I agree to comply with the conditions and principles of Good Clinical Practice as outlined in the European Clinical Trials Directives 2001/20/EC and the GCP Directive 2005/28/EC.

I agree to ensure that the confidential information contained in this document will not be used for any other purpose other than the evaluation or conduct of the clinical investigation without the prior written consent of the Sponsor.

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List of abbreviations and relevant definitions

ADA American Diabetes Association

ADE Adverse Device Effect

ASADE Anticipated Serious Adverse Device Effect

AE Adverse Event

AP Artificial Pancreas

AR Adverse Reaction

AUC Area Under the Curve

BMI Body Mass Index

CCTU Cambridge Clinical Trials Unit

CE Conformité Européenne (CE-mark)

CGM Continuous Glucose Monitoring

Cl Chief Investigator or Confidence Interval

CL Closed Loop

CRF Case Report Form

CSII Continuous Subcutaneous Insulin Infusion

DCCT Diabetes Control and Complications Trial

DKA Diabetic Ketoacidosis

DSMB Data Safety and Monitoring Board

eCRF Electronic Case Report Form

EudraCT European Clinical Trial Database

FDA US Food and Drug Administration

GCP Good Clinical Practice

HbA1c Glycated haemoglobin A1c

HD Haemodialysis

HFS Hypoglycaemia Fear Survey

IDE US Investigational Device Exemption

IRB Institutional Review Board

i.v. Intravenous

MDI Multiple Daily Injection therapy

MHRA Medicine and Healthcare products Regulatory Agency

MPC Model-Predictive-Control

NHS National Health Service

NICE National Institute for Health and Care Excellence

PI Principal Investigator

PD Peritoneal Dialysis

QALY Quality-Adjusted Life Years

R & D Research and Development

RCT Randomised Controlled Trial

REC Research Ethics Committee

RRT Renal Replacement Therapy

s.c. Subcutaneous

SADE Serious Adverse Device Effect

SAE Serious Adverse Event

SAP Sensor Augmented Pump Therapy

SD Standard Deviation

T1D Type 1 Diabetes Mellitus

T2D Type 2 Diabetes Mellitus

TSC Trial Steering Committee

USADE Unanticipated Serious Adverse Device Effect

WHO World Health Organisation

1 Study synopsis

Title of clinical trial	An open-label, randomised, 2-period cross-over study to assess the efficacy, safety and utility of fully-automated closed-loop insulin delivery in comparison with usual care, in adults with type 2 diabetes requiring maintenance dialysis
Short title	Closed-loop in adults with type 2 diabetes requiring dialysis (AP-Renal)
Sponsors name	Cambridge University Hospitals NHS Foundation Trust and University of Cambridge, Cambridge, UK
Medical condition or disease under investigation	Type 2 diabetes
Purpose of clinical trial	To determine the efficacy, safety and utility of fully automated closed-loop insulin delivery in the home setting in adults with type 2 diabetes requiring maintenance dialysis.
Study objectives	The study objective is to compare fully automated closed-loop insulin delivery with usual care in adults with type 2 diabetes requiring maintenance dialysis. 1. EFFICACY: The objective is to assess the ability of fully-automated closed-loop insulin delivery in maintaining CGM glucose levels within the target range from 5.6 to 10.0 mmol/l as compared to usual care in adults with type 2 diabetes requiring maintenance dialysis. 2. SAFETY: The objective is to evaluate the safety of fully automated closed-loop insulin delivery in terms of episodes and severity of hypoglycaemia, and nature and severity of other adverse events. 3. UTILITY: The objective is to determine the acceptability and duration of use of the closed-loop system.
Study design	An open-label, two-centre, randomised, two-period crossover study comparing fully automated closed-loop insulin delivery with usual care in adults with type 2 diabetes requiring dialysis. Two intervention periods in the home setting will last 20 days each with a 2-4 week washout period. The order of the two interventions will be random.

Cturdy and naints	The primary endpoint is the time spent in the target	
Study endpoints	glucose range from 5.6 to 10.0 mmol/l based on	
	CGM glucose levels during the 20 day home stay.	
	Other key endpoints:	
	Time spent with sensor glucose above	
	target (10.0 mmol/l)	
	Time spent with sensor glucose <3.9 mmol/l	
	Average of sensor glucose levels	
	Secondary endpoints include:	
	Time spent with sensor glucose below	
	target (5.6 mmol/l)	
	 Time spent with sensor glucose <3.0 mmol/l) 	
	Time spent with sensor glucose levels in significant by paraly coming (glucose).	
	significant hyperglycaemia (glucose levels > 20 mmol/l)	
	Standard deviation and coefficient of	
	variation of sensor glucose levels	
	AUC of glucose below 3.5 mmol/l (63	
	mg/dl) Total daily insulin requirements	
	Average inter-dialytic weight gain	
Safety evaluation	Assessment of frequency and severity of	
-	hypoglycaemic episodes and nature and severity of	
	other adverse events.	
Utility evaluation	Assessment of the acceptability and duration of use	
	of the closed-loop system.	
Participating clinical centres	UK	
	Addenbrooke's Hospital, Cambridge	
	University Hospitals NHS Foundation Trust,	
	Cambridge,	
	Switzerland	
	Bern University Hospital, Bern	
Sample size	32 adults completing the study. Recruitment will target up to 40 adults to allow for drop-outs.	
Summary of eligibility criteria	Key inclusion criteria:	
	1. Age 18 years or over	
	Diagnosis of type 2 diabetes using standard diagnostic practice	
	diagnostis prastiss	

	3. Requirement for maintenance dialysis		
	4. Current treatment with subcutaneous insulin		
	5. Screening HbA1c ≤ 11% (97mmol/mol) on		
	analysis from local laboratory 6. Subject is willing to perform regular finger-		
	prick blood glucose monitoring		
	7. Willingness to wear study devices		
	8. Literate in English (UK) or German		
	(Switzerland)		
	,		
	Kay avaluaian aritaria		
	Key exclusion criteria:		
	 Physical or psychological condition likely to interfere with the normal conduct of the 		
	study and interpretation of the study results		
	as judged by the investigator2. Known or suspected allergy to insulin		
	Lack of reliable telephone facility for contact		
	4. Pregnancy, planned pregnancy, or breast		
	feeding		
	Severe visual impairment		
	Severe hearing impairment		
	Medically documented allergy towards the		
	adhesive (glue) of plasters		
	8. Serious skin diseases located at places of		
	the body, which potentially are possible to		
	be used for localisation of the glucose		
	sensor		
	Illicit drugs abuse		
	Prescription drugs abuse		
	11. Alcohol abuse		
Maximum duration of study for a participant	12 weeks		
Recruitment	Participants will be recruited through the adult		
	diabetes outpatient clinics or haemodialysis units at		
	participating centres.		
Consent	Written informed consent will be obtained from		
	participants according to Research Ethics		
	Committee (REC) requirements.		
Screening and baseline	Eliaible porticipante will un derre a baseline		
assessment	Eligible participants will undergo a baseline evaluation including medical (diabetes) history and		
	current therapy.		
Randomisation	Eligible participants will be randomized in a 1:1		
	Eligible participants will be randomised in a 1:1 ratio using randomisation software to the use of		
	fully automated closed-loop insulin delivery or to		
i	,		

	usual care for 20 days, with a 2-4 week washout period between the two interventions.
1. Closed loop arm	Following randomisation, participants in the closed-loop group will receive training to cover key aspects of insulin pump use and CGM. Competency on the use of study devices will be evaluated
	Once competent in the use of the study pump and CGM, participants will receive training required for safe and effective use of the closed-loop system. During a 2-4 hour session participants will operate the system under the supervision of the clinical team. Competency on the use of closed-loop system will be evaluated. Thereafter, participants are expected to use closed-loop for 20 days without supervision or remote monitoring.
	 All participants will be provided with 24 hour telephone helpline and will also be given written instructions about when to contact clinical team.
2. Standard therapy (control arm)	Participants in the control group will continue with standard insulin therapy with blinded CGM for 20 days.
Study contacts	Follow up contacts will be conducted within 24 hours of starting each treatment arm and then at weekly intervals thereafter.
End of study assessments	Validated questionnaires evaluating the impact of the technology on diabetes management and quality of life will be completed. Participants will resume usual care.
Procedures for safety monitoring during trial	Standard operating procedures for monitoring and reporting of all adverse events (AE) will be in place, including serious adverse events (SAE), serious adverse device effects (SADE).
	A data safety and monitoring board (DSMB) will be informed of all serious adverse events and any unanticipated serious adverse device effects that occur during the study and will review compiled adverse event data at periodic intervals.
Criteria for withdrawal of patients on safety grounds	A participant may terminate participation in the study at any time without necessarily giving a reason and without any personal disadvantage. An investigator can stop the participation of a subject after consideration of the benefit/risk ratio. Possible reasons are:
	Serious adverse events

- 2. Significant protocol violation or non-compliance
- 3. Failure to satisfy competency assessment
- 4. Decision by the investigator, or the Sponsor, that termination is in the participant's best medical interest
- 5. Pregnancy, planned pregnancy, or breast feeding
- 6. Allergic reaction to insulin

Efforts will be made to retain participants in follow up for the final primary outcome assessment even if the intervention is discontinued, unless the investigator believes that it will be harmful for the participant to continue in the trial.

2 Summary

The main objective of this study is to determine the efficacy, safety and utility of fully automated closed-loop glucose control in the home setting over a 20 day period in adults with type 2 diabetes (T2D) requiring maintenance dialysis. This study builds on previous and on-going studies of closed-loop systems that have been performed in Cambridge in adults with type 1 diabetes in the home setting, and in adults with type 2 diabetes in the inpatient setting.

This is an open-label, two-centre, randomised, cross-over study, involving two home study periods during which glucose levels will be controlled either by a fully automated closed-loop system or by participants' usual insulin therapy in random order. Each treatment arm is 20 days long with a 2-4 week washout period between treatments. A total of up to 40 adults with T2D requiring maintenance dialysis will be recruited through outpatient clinics or the dialysis unit, to allow for 32 completed participants available for assessment.

Participants will receive appropriate training by the research team on the safe use of the study devices (insulin pump and continuous glucose monitoring (CGM) and closed-loop insulin delivery system). Participants in the control arm will continue with standard therapy and will wear a blinded CGM system.

The primary outcome is time spent with glucose levels in the target range between 5.6 and 10.0 mmol/L as recorded by CGM. Secondary outcomes are the time spent with glucose levels above and below target, as recorded by CGM, and other CGM-based metrics in addition to insulin requirements. Safety evaluation comprises the tabulation of severe hypoglycaemic episodes.

3 Background

3.1 Introduction

Diabetic nephropathy is the principal cause of end-stage renal disease (ESRD) accounting for 25% of incident cases in the UK (1). The number of adults with type 2 diabetes (T2D) is increasing and therefore the incidence of ESRD associated with diabetes is predicted to rise, with even more patients with T2D requiring renal replacement therapy with haemo- or peritoneal dialysis (HD or PD).

People with diabetes requiring dialysis are a vulnerable group, at high risk of adverse outcomes, with cardiovascular events the leading cause of mortality in this population. The overall survival in patients with diabetes on maintenance HD is approximately half that of their non-diabetic peers (3.7 vs. 7 years) (1).

The diabetes management of patients requiring dialysis is complex for both patients and health care professionals, but guidance on glycaemic targets and management algorithms is lacking. There is a clear need for improved delivery of care and novel approaches to the management of diabetes for people requiring dialysis.

3.2 Management of type 2 diabetes in patients requiring dialysis

Many oral anti-hyperglycaemia therapies are contraindicated in people with diabetes and ESRD, and insulin use is common in this population. There are important changes to both glucose and insulin metabolism that occur with dialysis. Once glomerular filtration rate (GFR) is sufficiently low, insulin clearance becomes markedly reduced, leading to higher levels of circulating insulin (3). The loss of clearance of insulin and reduction in gluconeogenesis in the kidneys leads to falling insulin requirements and subsequently, to a higher risk of hypoglycaemia if insulin is not adjusted (4). Uraemia-induced anorexia and weight loss may occur, reducing insulin requirements. Dialysis significantly improves insulin sensitivity by removing uraemic toxins (5) and people with diabetes using insulin are often dialysed against a dialysate containing supra-physiological glucose concentrations to mitigate against intra-dialytic hypoglycaemia.

Day to day variability in glucose levels and insulin requirements is high and therefore achieving glucose control can be very challenging. At present, basal bolus insulin regimens are the most flexible and often best suited to the glycaemic variability seen in patients with diabetes requiring maintenance haemodialysis; however demand a significant degree of self-management.

Regular glucose monitoring is essential for the assessment of glycaemic control in patients with diabetes on maintenance dialysis receiving insulin therapy. Hypoglycaemia can be a frequent occurrence, and symptoms of hypoglycaemia are often less pronounced in individuals requiring haemodialysis (hypoglycaemia unawareness). At present, individuals are required to undertake regular self-monitoring with finger-stick blood glucose measurements.

Continuous glucose monitoring (CGM) can provide an accurate assessment of glucose excursions in people with type 2 diabetes requiring haemodialysis (6, 7). CGM provides information on short-term glucose fluctuations associated with dialysis. Studies have shown that using CGM data to guide diabetes treatment significantly improves HbA1c and diabetes control (8).

3.3 Glycaemic control in people with type 2 diabetes requiring dialysis

Fear of hypoglycaemia, in addition to monitoring difficulties, complexity regarding the use of available treatments and therapeutic inertia means that glucose levels are often significantly above the usual glucose targets for individuals with type 2 diabetes requiring dialysis. In the short-term, patients on maintenance haemodialysis with diabetes are more likely to be able to maintain lower intra-dialytic weight gain if glucose control is optimised.

The long-term clinical benefits from maintaining effective glycaemic control in diabetes before ESRD is established and renal replacement therapy is required are well known (12, 13). The benefits of improving chronic hyperglycaemia at the stage of haemodialysis are less clear. Observational studies have shown that better glycaemic control predicts better survival among patients with diabetes on maintenance HD (14, 15). However, the threshold defining good glycaemic control was HbA1c <7.5% (58 mmol/mol), which is higher than conventional target HbA1c values. A one-year follow-up study of 23,000 subjects with diabetes suggested that low HbA1c levels may not confer survival benefit in ESRD (16). However, the association between poor glycaemic control and greater survival may be explained by confounding from factors such as malnutrition and anaemia (17). Overall, higher HbA1c was associated with increased mortality risk and lower HbA1c levels not related to malnutrition or anaemia appeared to be associated with improved survival in patients on maintenance HD.

In a prospective study in 444 patients on renal replacement therapy (HD or PD), glycated albumin (GA) and HbA1c levels were measured longitudinally (18). For each 5% increase in GA, the risk of death increased by 14%, whilst HbA1c did not predict survival. Restricting the analysis to the patients on haemodialysis, GA significantly predicted risk of death after adjustment for age, gender, race, and BMI, whereas HbA1c did not. A study in 9,201 haemodialysis patients with type 1 or type 2 diabetes showed that mortality was lowest at HbA1c 53–63 mmol/mol (7.0–7.9%) and increased progressively for either lower or higher HbA1c levels (19).

Attempts to intensify glycaemic control have the potential to increase mortality associated with severe hypoglycaemia (20). The increased risk of hypoglycaemia associated with conventional

insulin therapy used to achieve tight glycaemic control limits treatment intensification. With current management strategies, it is often necessary for target HbA1c levels among patients with diabetes requiring haemodialysis to be less stringent than levels recommended for other people with diabetes.

3.4 Hypoglycaemia in people with type 2 diabetes requiring dialysis

People with diabetes and chronic kidney disease (CKD) have twice the frequency of hypoglycaemia episodes than people with diabetes who do not have CKD (10.7 vs 5.3 episodes per 100 patient-months, respectively) (21). People with diabetes requiring haemodialysis are at even higher risk of hypoglycaemia. Severe hypoglycaemia, particularly nocturnal episodes, is also more common in individuals with ESRD due to reduced hypoglycaemia awareness, and is associated with arrhythmias.

Among patients admitted to hospital with hypoglycaemia, those with ESRD had a higher mortality rate, longer length-of-stay (LOS) and higher hospitalization costs compared to those without ESRD. In multivariate analysis, ESRD was significantly associated with increased odds for mortality (OR 2.92, 95% CI 1.98, 4.29, p<0.01), longer LOS (p<0.001) and higher hospitalization costs (p<0.001) (22).

Fear of hypoglycaemia for both patients and healthcare professionals is common and can impact quality of life and lead to suboptimal glucose control. Avoiding hypoglycaemia remains a priority in this vulnerable population.

3.5 Glycaemic targets in people with diabetes requiring dialysis

Glycated haemoglobin (HbA1c) measurement is the main biomarker for assessing glycaemic control in patients with diabetes and renal impairment, however, the accuracy of HbA1c values is poor due to the impact of anaemia and iron deficiency, elevated blood urea nitrogen levels, and uraemia. The relationship between HbA1c and average glycaemia has not been confirmed in patients receiving haemodialysis. HbA1c levels underestimate average blood glucose in patients on maintenance haemodialysis, so the quality of glycaemic control is overestimated, especially in patients with good to moderate glycaemic control.

Glycated albumin (GA) may offer a better opportunity to assess glycaemic control over a shorter time period (15–20 days) and with greater accuracy in patients with diabetes on maintenance haemodialysis. While HbA1c is affected by haemoglobin concentrations and erythropoietin dosage, these factors and serum albumin concentration do not significantly impact GA. In best-fit multivariate models, haemodialysis status significantly impacted HbA1c levels, without significant effect on GA (9-11). GA more accurately reflects recent glycaemic control.

National clinical guidelines do not distinguish between glycaemic targets for those with or without diabetic nephropathy (23, 24). Consensus groups have extrapolated from general recommendations, such as with Kidney Disease Outcomes Quality Initiative (KDOQI) in 2012, which suggested a target HbA1c level of 7% (53 mmol/mol) in those with CKD (25).

The Joint British Diabetes Societies (JBDS) and Association of British Clinical Diabetologists suggest a target HbA1c of 7.5-8.4% (58–68 mmol/mol) in patients with diabetes who require haemodialysis, given the hypoglycaemic and cardiovascular safety considerations (26, 27).

3.6 Closed-Loop Insulin Delivery

The emergence of new technologies including CGM (28), sensor augmented pump therapy (SAP) (29), and threshold pump suspend (30, 31) provides new opportunities to improve outcomes in diabetes. The most promising approach is closed-loop insulin therapy (32) which combines real-time CGM with insulin pump therapy to achieve glucose-responsive subcutaneous insulin delivery. The vital component of such a system, also known as an artificial pancreas (AP), is a computer-based algorithm. The role of the control algorithm is to compute the amount of insulin to be delivered by the pump using the real-time sensor glucose levels.

The closed-loop approach has been successfully evaluated in children and adults with type 1 diabetes in controlled laboratory studies (33-35) and in home settings (36-41). The results demonstrated improved glucose control and reduced risk of hypoglycaemia events. Psychosocial assessments supported acceptability and positive impact of this novel therapeutic approach. A fully closed-loop approach to the management of type 2 diabetes has been evaluated in the inpatient setting (42, 43), with results suggesting that this technology is a tangible option to improve glucose control in this population.

3.7 Closed-Loop Research in Cambridge

The University of Cambridge and collaborators have a considerable track record investigating closed-loop glucose control in young children, older children, adolescents, adults, and pregnant women with type 1 diabetes (36, 44-47). Since 2012, the University of Cambridge with collaborators have enrolled over 180 subjects in RCTs of free-living closed loop home conditions lasting 1 week to 2 years focusing on young people.

3.7.1 Preclinical testing of Cambridge closed-loop algorithm

The research conducted at the University of Cambridge focused on developing a closed loop system for overnight glucose (initial approach) and day-and-night control (more recent applications; see below) in subjects with T1D. Studies that have been performed employed model predictive control (MPC) – this algorithm estimates user-specific parameters from CGM measurements taken every 1 to 15 minutes and makes predictions of glucose excursions, which are then used to direct insulin infusion between meals and overnight whilst standard bolus calculator is used to deliver prandial insulin (48).

The MPC algorithm has been studied extensively using *in silico* testing utilising a simulator developed by members of the study team (49). The simulations suggested a reduced risk of nocturnal hypoglycaemia and hyperglycaemia with the use of the MPC algorithm (50).

3.7.2 Studies of closed-loop in children and adolescents with type 1 diabetes in the clinical research facility

To date around sixty children and adolescents with type 1 diabetes have been studied at the clinical research facility. Closed-loop insulin delivery was maintained on more than 100 nights. No episodes of significant hypoglycaemia (plasma glucose concentration less than 2.8 mmol/l) have been observed thus far during closed-loop blood glucose control. Results from these studies were published in The Lancet (33) and showed that overnight closed loop therapy increased the time spent euglycaemic by 37% and reduced the risk of overnight hypoglycaemia eight-fold, as compared to conventional pump treatment. Different real-life scenarios predisposing to nocturnal hypoglycaemia, such as afternoon exercise, were explored and closed-loop therapy reduced the risk of overnight hypoglycaemia as compared to conventional insulin pump therapy in a randomised, cross-over design.

3.7.3 Studies of closed-loop in adults with type 1 diabetes in the clinical research facility

We have completed two randomised overnight closed-loop studies in 24 adults with T1D, testing a similar closed-loop system comprising CGM and pump devices and the MPC algorithm. The first study (n=12) assessed the feasibility and efficacy of overnight closed-loop insulin delivery following a moderate-sized (60g carbohydrate) evening meal compared with conventional pump therapy. We demonstrated that overnight closed-loop insulin delivery, compared with usual continuous subcutaneous insulin infusion (CSII), significantly increased time in target plasma glucose range (3.9-8 mmol/l) by 24% and reduced glycaemic variability as measured by standard deviation of plasma glucose. The improvements in glucose control seen on closed-loop were even greater after midnight, when time in target increased by 41%. In the second study we tested the efficacy of overnight closed-loop following a common situation such as consuming a large (100g carbohydrate) evening meal and drinking alcohol (0.75g ethanol/kg body weight of 13%abv white wine). We showed that overnight closed-loop insulin delivery, compared with conventional CSII, similarly increased time in target plasma glucose between 3.9 and 8.0 mmol/l by 24% and reduced time spent above target by 11%, even following such challenges. Importantly these improvements during closed-loop were achieved with no increased requirement in the average rate of insulin infusion overnight. These results have been published in the British Medical Journal (51).

3.7.4 Overnight closed-loop study in children and adolescents with type 1 diabetes in home setting

Following successful demonstration of safety and efficacy of closed-loop insulin delivery in the research facility, overnight closed-loop studies under free living conditions were commenced in July 2012. The first study compared the efficacy and safety of closed-loop with sensor augmented pump therapy in 16 adolescents over a three week duration (36). Closed-loop was activated over at least 4 hours on 269 nights (80%); sensor data were collected over at least 4 hours on 282 control nights (84%). Closed-loop increased the time when glucose was in target range by a median 15% (interquartile range –9 to +43), P<0.001. Mean overnight glucose was reduced by a mean 0.8±3.2 mmol/l, P<0.001. Time when glucose was below 3.9 mmol/l was low in both groups but nights with glucose below 3.5mmol/l for at least 20min were less frequent during closed-loop (10% vs. 17%, P=0.01). Despite lower total daily insulin doses by a median 2.3 (interquartile range -4.7 to +9.3) units, P=0.009, overall 24h glucose was reduced by a mean 0.5 (standard deviation 2.3 mmol/l (P=0.006) during closed-loop.

In a second multicentre, crossover, randomised, controlled study, we compared 12 week use of an overnight closed-loop insulin delivery system with sensor augmented pump therapy in children and adolescents aged 6 to 18 years (37). The proportion of time with the night-time glucose level in the target range (3.9 to 8.0 mmol/l) was higher during the closed-loop phase than during the control phase (by 24.7 percentage points; 95% CI, 20.6 to 28.7; P<0.001), and the mean night-time glucose level was lower (difference, -1.6 mmol/l; 95% CI,-2.2 to -1.1; P<0.001). The area under the curve for the period in which the day-and-night glucose levels were less than 3.5 mmol/l was lower by 42% (95% CI, 4 to 65; P = 0.03). Two severe hypoglycaemic episodes occurred during the closed-loop phase when the closed-loop system was not in use.

3.7.5 Overnight closed-loop studies in adults with type 1 diabetes in home setting

A four week overnight closed-loop study under free living conditions in 24 adults with type 1 diabetes on insulin pump therapy in a multicentre crossover study design was completed in 2014 (41). Closed-loop was utilised over median 8.3 (interquartile range 6.0, 9.6) hours on 555 nights (86%). The proportion of time when overnight glucose was in the overnight target range between 3.9 and 8.0 mmol/l from midnight to 07:00 was significantly higher during closed-loop compared to sensor augmented pump therapy (52.6%±10.6 vs. 39.1%±12.8, mean±SD; p<0.001). Mean overnight glucose (8.2±0.9 vs. 9.0±1.3 mmol/l, p=0.005) and time spent above target (44.3%±11.9 vs. 57.1%±15.6, p=0.001) were significantly lower during closed-loop. Time spent below target was low and comparable between interventions [1.8%(0.6, 3.6) vs. 2.1%(0.7, 3.9), p=0.28].

3.7.6 Day-and-night closed-loop studies in adolescents with type 1 diabetes in home setting

We completed a randomised, crossover design study in adolescents aged 10 to 18 years who underwent two 7-day home periods of sensor-augmented insulin pump therapy or closed-loop insulin delivery without supervision or remote monitoring (39). The proportion of time when the sensor glucose level was in the target range (3.9–10 mmol/L) was increased during closed-loop insulin delivery compared with sensor-augmented pump therapy (72% vs. 53%, P < 0.001; primary end point), the mean glucose concentration was lowered (8.7 vs. 10.1 mmol/L, P = 0.028), and the time spent above the target level was reduced (P = 0.005) without changing the total daily insulin amount (P = 0.55). The time spent in the hypoglycaemic range was low and comparable between interventions. A three week single centre study in children and adolescents has also been completed (N = 12).

3.7.7 Day and night closed-loop studies in adults with type 1 diabetes in home setting

In 2014, we completed a first study testing a day and night home system over a seven day period in 17 adults. This randomised clinical trial adopted a multicentre, multi-national, crossover design. During the home phase, the percentage time when glucose was in target range (3.9 to 10.0 mmol/l) was significantly higher during closed loop compared to sensor augmented pump therapy (75 [61, 79] vs. 62 [53, 70]%, median [IQR], p=0.005). Mean glucose (8.1 vs. 8.8 mmol/l, p=0.027) and time spent above target (p=0.013) were lower during closed-loop while time spent below target was comparable (p=0.339). Increased time in target was observed during both day-time (p=0.017) and night-time (p=0.013).

We completed a multicentre, multinational, crossover, randomised, controlled study under free living home conditions comparing 24/7 closed-loop insulin delivery with sensor augmented pump therapy (control intervention) in 33 adults with type 1 diabetes (37). The proportion of time that the glucose level was in the target range (3.9 to 10.0 mmol/l) was 11.0 percentage points (95% confidence interval [CI], 8.1 to 13.8) greater with the use of the closed-loop system day and night than with control therapy (P<0.001). The mean glucose level was lower during the closed-loop phase than during the control phase (difference, -0.6 mmol/l; 95% CI, -0.9 to -0.3; P<0.001), as were the area under the curve for the period when the glucose level was less than 3.5 mmol/l (39% lower; 95% CI, 24 to 51; P<0.001) and the mean glycated haemoglobin level (difference, -0.3%; 95% CI, -0.5 to -0.1; P = 0.002).

3.7.8 Closed-loop studies in adults with type 2 diabetes

The Cambridge closed-loop system has been shown to be safe and feasible in insulin-naive patients with type 2 diabetes in a controlled research facility setting (52).

We have previously assessed fully automated closed-loop insulin delivery in non-critical care patients with type 2 diabetes hospitalised in the general wards (53). Forty participants were randomised to either automated fully closed-loop insulin delivery or usual insulin therapy for a 72h study period. Results showed that closed-loop significantly increased time spent within target glucose range (5.6-10.0mmol/l), without any increase in the risk of hypoglycaemia. Closed-loop insulin delivery without meal-time boluses is effective and safe in insulin-treated adults with type 2 diabetes.

In a larger multi-national study, 136 adults with type 2 diabetes who required insulin therapy were randomised to receive either closed-loop insulin delivery or conventional subcutaneous insulin therapy for up to 15 days or until hospital discharge (43). The mean percentage of time that the sensor glucose measurement was in the target range was 65.8% in the closed-loop group and 41.5% in the control group, a difference of 24.3 percentage points (P<0.001); values above the target range were found in 23.6% and 49.5% of the patients, respectively, a difference of 25.9 percentage points (P<0.001). The mean glucose level was 8.5 mmol/l in the closed-loop group and 10.4 mmol/l in the control group (P<0.001). There was no significant between-group difference in the duration of hypoglycaemia or in the amount of insulin that was delivered. Among inpatients with type 2 diabetes receiving noncritical care, the use of an automated, closed-loop insulin-delivery system resulted in significantly better glycaemic control than conventional subcutaneous insulin therapy, without a higher risk of hypoglycaemia.

3.8 Risk and benefits

A potential key benefit of closed-loop insulin delivery is a reduction in hypoglycaemia which has been shown to be associated with cardiovascular events, increased hospital admissions and increased mortality. The long-term impact of improved glucose control in this population may be reduced rates of diabetes complications and improved quality of life.

Any potential risks presented by this investigation have been minimized and adequate testing, safeguards, and safety monitoring will be incorporated into the investigation to further minimize and mitigate these risks. A detailed Risk Management File adopting risk management processes complying with EN ISO 14971:2012 Medical Devices – Application of Risk Management to Medical Devices, will be submitted as part of the regulatory submission to the MHRA.

3.9 CamAPS HX fully-automated closed loop system to be used in the present study

In the present study, we will use the CamAPS HX closed-loop system comprising:

- Dana insulin pump (Diabecare, Sooil, Seoul, South Korea)
- Dexcom G6 real-time CGM sensor (Dexcom, Northridge, CA, USA)
- An Android smartphone hosting CamAPS HX Application with the Cambridge model predictive control algorithm and communicating wirelessly with the insulin pump
- Cloud upload system to monitor CGM/insulin data.

An overview of this proposed automated closed loop system is given in Figure 1.



Figure 1: CamAPS HX comprises Samsung Galaxy phone (or similar) running Cambridge control algorithm, Dana insulin pump (Sooil), G6 real-time CGM sensor (Dexcom).

3.10 Rationale for the present study

The study builds on recent technological advances of closed-loop insulin delivery (artificial pancreas). Studies from our group have assessed the safety and efficacy of closed-loop insulin delivery in T1D in a controlled research setting and at home. Closed-loop use in hospitalised patients has been evaluated in the intensive care setting and on the general wards, and has demonstrated efficacy and safety in achieving target glucose range when compared to standard treatment (42, 54-57). Despite studies in people with diabetes requiring maintenance dialysis reporting an adverse relationship between dysglycaemia and clinical outcome, glycaemic management of these patients remains suboptimal.

Closed-loop insulin delivery may be of benefit to such patients in whom the optimal dosing regimen is difficult to establish, and hence may be better facilitated by an algorithm-initiated and driven insulin therapy. Most importantly, closed-loop may provide a safer method of insulin delivery with the added benefit of continuous monitoring of glucose levels, thus minimising the likelihood of hyper- and hypoglycaemic events and their known associated worse outcomes.

The purpose of this study is to test the impact of closed loop insulin delivery in patients with type 2 diabetes requiring maintenance dialysis on time in target glucose range and frequency of hypoglycaemia. The present study will also test the feasibility and acceptance of this therapy so that it could be considered as a standard treatment modality in the future.

4 Objectives

4.1 Efficacy

To assess efficacy of day-and-night fully automated closed-loop insulin delivery in maintaining glucose levels within the target range from 5.6 to 10.0 mmol/l based on subcutaneous continuous glucose monitoring (CGM), during the 20 day home stay as compared to usual care.

4.2 Safety

To evaluate the safety of fully-automated closed-loop glucose control in terms of episodes and severity of hypoglycaemia and nature and severity of other adverse events.

4.3 Utility

To determine the acceptability, duration and frequency of use of the closed-loop system.

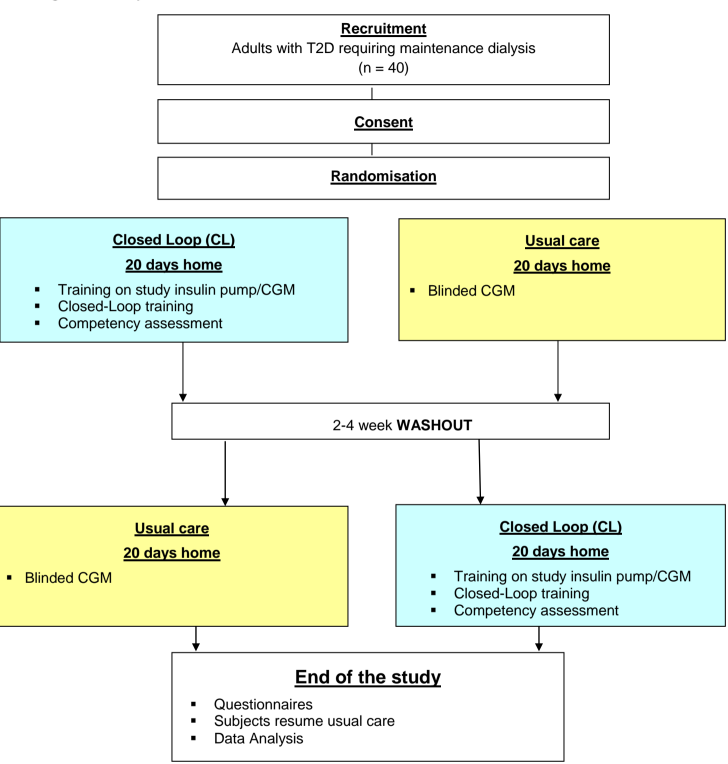
5 Study design

An open-label, two-centre, randomised, two-period crossover study assessing the efficacy, safety and utility of fully automated closed-loop insulin delivery compared with usual care in adults with type 2 diabetes requiring maintenance dialysis. Two intervention periods in the home setting will last 20 days each with 2-4 weeks washout period. The order of the two interventions will be random.

Up to a total of 40 adults with T2D requiring maintenance dialysis will be recruited, to allow for 32 completed subjects available for assessment.

The study flow chart is outlined in Figure 2.

Figure 2: Study flow chart



6 Study participants

6.1 Study population

Adults with type 2 diabetes requiring maintenance dialysis will be recruited

6.1.1 Inclusion criteria

- 1. The subject is age 18 years or over
- 2. Diagnosis of type 2 diabetes using standard diagnostic practice
- 3. The subject requires maintenance dialysis
- 4. The subject requires current treatment with subcutaneous insulin
- 5. Screening HbA1c ≤ 11% (97mmol/mol) on analysis from local laboratory
- 6. Subject is willing to perform regular finger-prick blood glucose monitoring
- 7. The subject is literate in English (UK) or German (Switzerland)
- 8. The subject is willing to wear study devices 24/7 during intervention arm and follow study specific instructions

6.1.2 Exclusion criteria

- Physical or psychological condition likely to interfere with the normal conduct of the study and interpretation of the study results as judged by the investigator
- 2. Known or suspected allergy to insulin
- 3. Lack of reliable telephone facility for contact
- 4. Pregnancy, planned pregnancy, or breast feeding
- 5. Severe visual impairment
- 6. Severe hearing impairment
- 7. Medically documented allergy towards the adhesive (glue) of plasters or unable to tolerate tape adhesive in the area of sensor placement
- 8. Serious skin diseases located at places of the body, which potentially are possible to be used for localisation of the glucose sensor
- 9. Illicit drugs abuse
- 10. Prescription drugs abuse
- 11. Alcohol abuse

6.2 Recruitment and informed consent

The study will aim for 32 completed subjects. Recruitment will target up to 40 subjects to allow for drop-outs. Participants will be recruited from the adult diabetes outpatient clinics or from the dialysis units at the following centres.

- Addenbrooke's Hospital, Cambridge University Hospitals NHS Foundation Trust, Cambridge, UK
- 2. Bern University Hospital, Bern, Switzerland

Potential participants will be identified by their treating clinicians and a contact with the research team will be established if agreed. Study information leaflets and/or similar recruitment material will be handed out or sent to participants by the research team including an invitation to join the study. Written informed consent will be obtained from all participants before any study related activities.

7 Methods under investigation

7.1 Name and description of the method of investigation

The investigational treatment is a closed-loop system, see section 3.9 or follow up prototypes of the automated closed-loop insulin delivery system manufactured by the Cambridge University Hospitals NHS Foundation Trust. Component versions will be identified during regulatory submission to the national regulatory bodies.

7.2 Intended purpose

The intended purpose of the investigational treatment is automated day and night fully closed-loop insulin delivery. The investigated medical device is used to manage glucose levels in adults with type 2 diabetes, using a fully closed-loop approach.

7.3 Method of administration

The closed-loop system consists of components directly attached to the patient, which are the CGM sensor/transmitter and the insulin pump. The component not directly attached to the patient is the handheld smartphone containing closed-loop algorithm and communicating wirelessly with the insulin pump.

7.4 Required training

Prior to commencement of the study, the research team nurses/clinicians will be trained to use the closed-loop system and its components. Prior to the use of study devices, participants will be trained to use the study CGM device, the study pump and the closed-loop system. Competency assessments of the participants' capability to use study devices and the closed-loop system will be made.

7.5 Precautions

During treatment with insulin there is a risk of hypoglycaemia and hyperglycaemia. In-hospital testing and hazard analysis have documented reduced risk of hypoglycaemia and hyperglycaemia during closed loop compared to conventional treatment.

7.6 Accountability of the method under investigation

The local Investigator will provide training for the study participants and will make every effort, through regular contact, to ascertain that the closed loop system is used for the study purposes only. Devices will be identified using batch/lot/serial numbers and the location of investigational devices and their dates of use by subjects will be documented throughout the study.

8 Study schedule

8.1 Overview

The study will be co-ordinated from the Institute of Metabolic Science, Addenbrooke's Hospital, Cambridge. Participants in the UK will be recruited through the Dialysis unit in Addenbrooke's Hospital, Cambridge, and other satellite haemodialysis units in the Cambridgeshire region. Participants in Switzerland will be recruited through the Dialysis unit in University Hospital Bern.

The study will consist of up to 5 visits and 6 contacts over the two study periods (closed-loop vs. usual care). The study periods will last 20 days each. The order of the two interventions will be random. There will be a 2-4 week washout period between the two study periods.

Prior to the closed-loop intervention, there will be a training visit, which will be conducted at the CRF/dialysis unit, followed by a 20 day study period at the participant's home. The training visit will last approximately 4 hours and will return home when the participant is competent and confident in using the closed-loop system. Maximum time in study is 12 weeks.

Table 1 outlines study activities when CL intervention precedes standard care.

Table 2 outlines study activities when standard care precedes CL intervention.

Table 1: Schedule of study visits / phone contacts when closed-loop intervention precedes usual care

	Visit/ contact	Description	Start relative to previous / next Visit / Activity	Duration		
	Visit 1	Recruitment visit: Consent, baseline assessments including questionnaires	-	1-2 hours		
	RANDOMISATION					
Visit 2	CGM, Insulin pump and closed loop training Competency assessment and initiation of closed loop	Within 1 to 3 weeks of Visit 1 Training visits can be repeated if competency not achieved	3-4 hours			
ntion s)	Contact 1*	Review use of study devices	24h after Visit 2 (±3 days)	<0.5 hour		
CL Intervention (20 days)	Contact 2*	Review use of study devices	7 days after Visit 2 (±3 days)	<0.5 hour		
Contact	Contact 3*	Review use of study devices	14 days after Visit 2 (±3 days)	<0.5 hour		
	Visit 3	End of closed-loop treatment period (20 days) Return devices. Revert back to usual diabetes therapy. Questionnaires	After 20 days of Visit 2	1-2 hours		
		Washout period	Immediately after Visit 3	2-4 weeks		
	Visit 4	Blinded CGM insertion Review of diabetes management	Within 2-4 weeks of Visit	1-2 hours		
e (c	Contact 4*	Review diabetes management	24h after Visit 4 (±3 days)	<0.5 hour		
Usui (20	Contact 5*	Review diabetes management	7 days after Visit 4 (±3 days)	<0.5 hour		
	Contact 6*	Review diabetes management.	14 days after Visit 4 (±3 days)	<0.5 hour		
	Visit 5	End of usual care period (20 days) Questionnaires	After 20 days of Visit 4	1-2 hours		
	* could be done at home or phone/email					

Table 2: Schedule of study visits / phone contacts when usual care precedes closed loop intervention

	Visit/ contact	Description	Start relative to previous / next Visit / Activity	Duration		
	Visit 1	Recruitment visit: Consent, baseline assessments including questionnaires	-	1-2 hours		
	RANDOMISATION					
	Visit 2	Blinded CGM insertion Review of diabetes management	Within 1-3 weeks of Visit	1-2 hours		
re ()	Contact 1*	Review diabetes management	24h after Visit 2 (±3 days)	<0.5 hour		
Usual Care (20 days)	Contact 2*	Review diabetes management	7 days after Visit 2 (±3 days)	<0.5 hour		
ă S	Contact 3*	Review diabetes management	14 days after Visit 2 (±3 days)	<0.5 hour		
	Visit 3	End of usual care period (20 days) Questionnaires	After 20 days of Visit 2	1-2 hours		
		Washout period	Immediately after Visit 3	2-4 weeks		
Vis		CGM, Insulin pump and closed	Within 2 to 4 weeks of			
	Visit 4	loop training Competency assessment and initiation of closed loop	Visit 3 Training visits can be repeated if competency not achieved	3-4 hours		
tion	Visit 4 Contact 4*	Competency assessment and	Training visits can be repeated if competency	3-4 hours		
ntervention 20 days)		Competency assessment and initiation of closed loop	Training visits can be repeated if competency not achieved			
CL Intervention (20 days)	Contact 4*	Competency assessment and initiation of closed loop Review use of study devices	Training visits can be repeated if competency not achieved 24h after Visit 4 (±3 days) 7 days after Visit 4 (±3	<0.5 hour		
CL Intervention (20 days)	Contact 4* Contact 5*	Competency assessment and initiation of closed loop Review use of study devices Review use of study devices	Training visits can be repeated if competency not achieved 24h after Visit 4 (±3 days) 7 days after Visit 4 (±3 days) 14 days after Visit 4 (±3	<0.5 hour <0.5 hour		

8.2 Baseline visit (Visit 1)

Once participants have agreed to participate in the study, they will be invited for the baseline visit, when the following activities will be performed by the research team:

- written informed consent
- · checking inclusion and exclusion criteria
- medical (diabetes) history
- record of current insulin therapy
- body weight measurement
- questionnaires will be distributed to assess quality of life and diabetes management.

Switzerland only

Woman of reproductive age will be required to take a pregnancy test and will be advised to use contraception during study participation.

8.3 Randomisation

On completion of Visit 1, eligible participants will be randomised in a 1:1 ratio using randomisation software to the initial use of fully automated closed-loop glucose control or to usual care for 20 days, with a 2-4 week washout period before crossing over to the second intervention arm.

8.4 Post-randomisation training (Visits 2 and 4)

8.4.1 Closed loop intervention

Participants starting the closed-loop arm will receive training to cover key aspects of insulin pump use and CGM, prior to training on closed-loop insulin delivery. Particular attention will be paid to:

- Insulin cartridge and infusion set changes and correct priming procedure
- Sensor insertion and calibration
- Blood glucose targets and alarm settings
- Hypo- and hyperglycaemia management
- Connection and disconnection of the closed-loop system

Written easy to use guidelines for the operation of insulin pump, CGM and closed-loop will be provided. This session will be conducted by a professional pump educator and/or member of the study team. Device manual guides will be provided.

Competency in the use of study pump, CGM and closed-loop system will be assessed by the study team. Only subjects who demonstrate competency in use of the system will be allowed to continue to the home study phase.

Subjects will be advised to use closed-loop 24/7 for the next 20 days. Written step by step guidance will also be provided, including how to deal with low and high glucose at home. Subjects will be provided with 24 hour telephone helpline and information on when to contact study team.

The subject is allowed to drive while adhering to usual precautions and country specific rules and regulations.

8.4.2 Standard therapy (control intervention)

Participants in the control arm will continue to follow their current diabetes management plan for the 20 day study period. During the control arm, participants and/or the clinical team are free to adjust insulin therapy as per usual clinical practice, but no active treatment optimisation will be undertaken by the study team. For self-monitoring of blood glucose (SMBG) throughout the study, subjects will continue using their own glucose meter provided the meter type meets ISO standards (Standard 15197: 2013).

Participants will be shown how to insert the study CGM and they will be asked to wear a blinded continuous glucose monitoring (CGM) system during the 20 day home stay. If the sensor fails or sensor function is interrupted prematurely (detached sensor), another sensor will be inserted. The sensor(s) will be sent back/collected by the research team once the sensor life has expired and/or the sensor has detached.

8.5 Contacts after initiation of study arm

Participants will be contacted 24 hours and 1 and 2 weeks after initiation of the respective study arm. These contacts can be via telephone/email. The purpose of this contact would be to troubleshoot any problems, and to record any adverse events, device deficiencies, and changes in insulin therapy.

8.6 End of first study arm visit (Visit 3)

On completion of the first study arm, participants will be invited to attend the research facility/clinical area 20 days after study arm initiation. Participants will return all study devices and then resume usual care. Participants will be asked to complete questionnaires.

8.7 Washout

A minimum washout period of 2 weeks must be ensured between treatment periods. Duration of the wash out period has been chosen to minimise any carry over effect between the two interventions. The subject will continue with their usual diabetes care during the washout period. Following the washout period subjects will cross over to alternative intervention.

8.8 End of study visit (Visit 5)

On completion of the second study arm, participants will be invited to attend the research facility/clinical area 20 days after study arm initiation. Participants will return all study devices and then resume usual care. Participants will be asked to complete questionnaires.

8.9 Participant withdrawal criteria

The following pre-randomisation withdrawal criterion will apply:

1. Subject is unable to demonstrate safe application of insulin therapy as judged by the investigator

The following pre- and post-randomisation withdrawal criteria will apply:

- Subject is unable to demonstrate safe use of insulin injections or study insulin pump and/or CGM and/or closed loop during post randomisation training period as judged by the investigator
- 3. Subjects may terminate participation in the study at any time without necessarily giving a reason and without any personal disadvantage
- 4. Significant protocol violation or non-compliance
- 5. Recurrent severe hypoglycaemia events not related to the use of the closed loop system
- 6. Recurrent severe hyperglycaemia event/DKA unrelated to infusion site failure and related to the use of the closed loop system
- 7. Decision by the investigator or the Sponsor that termination is in the subject's best medical interest
- 8. Allergic reaction to insulin

9. Allergic reaction to adhesive surface of infusion set or glucose sensor

Subjects who are withdrawn for reasons stated in (4) to (9) will be invited to complete questionnaires at the end of the planned study intervention. Subjects who discontinue study intervention prior to the final visit will receive an exit survey.

8.10 Study stopping criteria

The study may be stopped if three consecutive participants withdraw from the study, the study may be stopped for safety reasons or on the advice of the Data Safety Monitoring Board.

8.11 Support telephone line

In addition to standard clinical advice, there will be a 24-hour telephone helpline to the research team for subjects in case of any technical device or problems related to diabetes management such as hypo- or hyperglycaemia.

8.12 Subject reimbursement

The study will provide the CGM device, insulin pump, CL components and related consumables. A study payment will be made to reflect local practice. The amount paid will be specified in the participant information sheet/informed consent form and REC application form. Reasonable travel expenses will be reimbursed. After completing the study, subjects will not keep the study devices. They will revert to their usual diabetes management.

9 Endpoints

9.1 Efficacy endpoints

9.1.1 Primary efficacy endpoint

The primary endpoint is the time spent in the target glucose range from 5.6 to 10.0 mmol/l during the 20 day study periods based on continuous glucose monitoring (CGM).

9.1.2 Other Key Endpoints

- Time spent above target glucose (10.0 mmol/l)
- Average of glucose levels
- Time spent with glucose levels <3.9 mmol/l

9.1.3 Secondary efficacy endpoints

Secondary endpoints include:

- Time spent below target glucose (5.6 mmol/l)
- Standard deviation and coefficient of variation of glucose levels
- The time with glucose levels <3.0 mmol/l
- The time with glucose levels in significant hyperglycaemia (glucose levels > 20 mmol/l)
- Total, basal and bolus insulin dose
- AUC of glucose below 3.5mmol/l

Endpoints regarding glucose levels will be based on sensor glucose data.

9.1.4 Exploratory endpoints

Average inter-dialytic weight gain

9.2 Safety evaluation

Safety evaluation will comprise the number of episodes of severe hypoglycaemia as well as the number of subjects experiencing severe hypoglycaemia, severe hyperglycaemia (>20 mmol/l) and number, nature and severity of any other adverse events.

All subjects including those who withdraw will be included in the safety evaluation.

9.3 Utility evaluation

Utility evaluation is the frequency and duration of use of the closed-loop system. Expectations, attitudes and responses to the closed-loop system will be assessed using questionnaires.

10 Assessment and reporting of adverse events

10.1 Definitions

10.1.1 Reportable Adverse Events

A reportable Adverse Event is any untoward medical occurrence that meets criteria for a serious adverse event or any unanticipated medical occurrence in a study subject that is study or device-related. Device deficiencies that could have led to a serious adverse device effect will also be reported (ISO 14155: 8.2.5 and 9.8).

10.1.2 Adverse Events

An adverse event (AE) is any untoward medical occurrence, unintended disease or injury, or untoward clinical signs (including abnormal laboratory findings) in a subject who has received an investigational device, whether or not related to the investigational medical device (ISO 14155: 3.2). This definition includes events related to the device under investigation or the comparator or to the study procedures. For users or other persons, this definition is restricted to events related to the investigational device. The following anticipated adverse events will not be recorded:

- Non clinically significant skin reactions as judged by investigator
- Pre-existing medical conditions
- New illnesses or conditions not requiring concomitant medication or medical intervention/procedures
- Non severe hypoglycaemia
- Hyperglycaemia without significant ketonaemia

10.1.3 Adverse Device Effect

An Adverse Device Effect (ADE) is an adverse event related to the use of an investigational medical device (ISO 14155: 3.1). This includes adverse events resulting from insufficient or inadequate instructions for use, deployment, implantation, installation, or operation, or any malfunction of the investigational medical device. This definition also includes any event resulting from use error or from intentional misuse of the device under investigation.

10.1.4 Serious Adverse Event

A serious adverse event (SAE) is an adverse event that:

- led to a death
- led to a serious deterioration in the health of the subject, that either resulted in:
 - o a life threatening illness or injury
 - o a permanent impairment of a body structure or function
 - o in-patient hospitalisation or prolonged hospitalisation
 - medical or surgical intervention to prevent life-threatening illness or injury or permanent impairment to a body structure or a body function
- led to foetal distress, foetal death or a congenital abnormality or birth defect

A planned hospitalisation for pre-existing condition, or a procedure required by the study protocol, without a serious deterioration in health, is not considered to be a serious adverse event.

More than one of the above criteria can be applicable to one event. Life-threatening in the definition of a serious adverse event or serious adverse reaction refers to an event in which the subject was at risk of death at the time of the event; it does not refer to an event which hypothetically might have caused death if it were more severe. Medical judgement should be exercised in deciding whether an adverse event or reaction is serious in other situations.

Important adverse events or reactions that are not immediately life-threatening or do not result in death or hospitalisation but may jeopardise the subject or may require intervention to prevent one of the other outcomes listed in the definition above, should also be considered serious.

The following serious adverse events are anticipated:

- Severe hypoglycaemia
- DKA

10.1.5 Serious Adverse Device Effect

A Serious Adverse Device Effect (SADE) is an adverse device effect that has resulted in any of the consequences characteristic of a serious adverse event.

10.1.6 Unanticipated Serious Adverse Device Effect

An Unanticipated Serious Adverse Device Effect (USADE) is a serious adverse device effect which by its nature, incidence, severity or outcome has not been identified in the current version of the protocol or the risk analysis report (ISO 14155: 3.42).

An Anticipated Serious Adverse Device Effect (ASADE) is a serious adverse device effect which by its nature, incidence, severity or outcome has been identified in the protocol.

10.1.7 Device Deficiencies

A device deficiency is an inadequacy of a medical device with respect to its identity, quality, durability, reliability, safety or performance such as malfunction, misuse or user error and inadequate labelling (ISO 14155: 3.15). A device deficiency may lead to an Adverse Device Effect or Serious Adverse Device Effect. The following anticipated device deficiencies and device-related issues will not be recorded:

- Infusion set occlusion/leakage not leading to ketonaemia
- Sensor failure due to significant over/under-reading (difference>3mM) or detachment
- Premature interruption of sensor-life

- Battery lifespan deficiency due to inadequate charging or extensive wireless communication
- CAD error messages not needing system replacement
- Intermittent device communication failure not leading to system replacement

10.1.8 Adverse event intensity

Intensity	Definition
Mild	Patient is aware of signs and symptoms but they are easily tolerated
Moderate	Signs / symptoms cause sufficient discomfort to interfere with usual activities
Severe	Patient is incapable to work or perform usual activities

NB. The term "severe" is often used to describe the intensity (severity) of a specific event. This is not the same as 'serious', which is based on patient/event outcome or action criteria (see definition 10.1.4). For example, itching for several days may be rated as severe, but may not be clinically serious.

10.1.9 Adverse event causality

Intensity	Definition
Not assessable	A report suggesting an adverse event, which cannot be judged because information is insufficient or contradictory, and which cannot be supplemented or verified.
Unlikely	A clinical event, including laboratory test abnormality, with a temporal relationship, which makes a causal relationship improbable, and in which other drugs/treatments, chemicals or underlying disease(s) provide plausible explanations.

Possible	A clinical event, including laboratory test abnormality, with a reasonable time sequence to administration of the treatment/use of investigational treatment/device, but which also could be explained by concomitant diseases or other drugs/treatments or chemicals.
Probable	A clinical event, including laboratory test abnormality, with a reasonable time sequence to administration of the treatment/use of medical method/device, unlikely to be attributable to concomitant disease(s) or other drugs/treatments or chemicals, and which follows a clinically reasonable response on withdrawal (dechallenge). Rechallenge information is not required to fulfil this definition.
Definite/certain	A clinical event, including laboratory test abnormality, occurring in a plausible time relationship to study treatment/use of medical method/device and which cannot be explained by concomitant disease(s), other drugs/treatments or chemicals. The response to withdrawal of the treatment (dechallenge) should be clinically plausible. The event must be unambiguous, either pharmacologically or as phenomenon, using satisfactory rechallenge procedures if necessary.

(Reference: WHO-UMC Causality Categories)

10.2 Recording and reporting of adverse events, serious adverse events and device deficiencies

10.2.1 Monitoring period of adverse events

The period during which adverse events will be reported is defined as the period from the beginning of the study (obtaining informed consent) until 72 hours after the end of the study participation. Adverse events that continue after the subject's discontinuation or completion of the study will be followed until their medical outcome is determined or until no further change in the condition is expected. The follow up of AEs may therefore extend after the end of the clinical investigation; however no new AEs will be reported after the trial reporting period.

10.2.2 Recording and reporting of adverse events

Throughout the course of the study, all efforts will be made to remain alert to possible adverse events or untoward findings. The first concern will be the safety of the subject, and appropriate medical intervention will be taken. The investigator will elicit reports of adverse events from the subject at each visit and complete adverse event forms. All AEs, including those the subject reports spontaneously, those the investigators observe, and those the subject reports in response to

questions will be recorded on paper or electronic AE forms at each site within seven days of discovering the event.

The study investigator will assess the relationship of any adverse event to be device-related or unrelated by determining if there is a reasonable possibility that the adverse event may have been caused by the study device or study procedures. The individual investigator at each site will be responsible for managing all adverse events according to local protocols, and decide if reporting is required.

10.2.3 Severe hypoglycaemia

Severe hypoglycaemia will be defined as an event requiring assistance of another person to actively administer carbohydrate, glucagon, or other resuscitative actions. These episodes may be associated with sufficient neuroglycopaenia to induce seizure or coma. If plasma glucose measurements are not available during such an event, neurological recovery attributable to the restoration of plasma glucose to normal is considered sufficient evidence that the event was induced by a low plasma glucose concentration.

Severe hypoglycaemia will be regarded as a foreseeable adverse event and an adverse event form will be completed. Severe hypoglycaemia is not necessarily a serious adverse event and hence may not require immediate reporting to the Sponsor. Non-severe hypoglycaemia will not be reported or considered an adverse event.

10.2.4 Reporting of serious adverse events and serious adverse device effects

When reporting adverse events, all pertinent data protection legislation must be adhered to.

The serious adverse event report should contain the following information*:

- 1. Study identifier (EudraCT number if applicable)
- 2. Participant's unique study number
- 3. Date of birth
- 4. Event description
- 5. Start date of event
- 6. Laboratory tests used and medical interventions used to treat the SAE
- 7. Planned actions relating to the event, including whether the study device was discontinued
- 8. Statement on the patient's current state of health

- 9. Criterion for seriousness (i.e. death, life threatening, hospitalisation, disability/incapacity or other)
- 10. Evaluation of causality (including grade of relatedness) with the following (more than one may apply):
 - a. the investigational treatment/medical device
 - b. the clinical study/a study specific procedure
 - c. other: e. g. concomitant treatment, underlying disease
- 11. Date of procedure
- 12. Reporter's name, date and signature

*In the case of incomplete information at the time of initial reporting, all appropriate information should be provided as soon as this becomes available.

UK only

The relationship of the SAE to the investigational treatment / medical device should be assessed by the investigator at site, as should the anticipated or unanticipated nature of any SAEs and SADEs.

All SAEs whether or not deemed investigational method/device related and whether anticipated or unanticipated must be reported to the Sponsor by email or fax within 24 hours (one working day) of the Investigator learning of its occurrence.

SAEs should be reported to:

Stephen Kelleher

Cambridge University Hospitals

NHS Foundation Trust

Box 277, Addenbrooke's Hospital

Hills Road, Cambridge, CB2 0QQ, UK

Phone: +44 (0) 1223 217418

Fax: +44 (0) 1223 348494

E-mail: enquiries@addenbrookes.nhs.uk

A written report must follow within five working days and is to include a full description of the event and sequelae, in the format detailed on the Serious Adverse Event reporting form. If applicable, the Sponsor will notify the competent authority of all Serious Adverse Events in line with pertinent legal requirements.

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The Investigator will notify the Research Ethics Committee (REC) in UK of all Serious Adverse

Events in line with pertinent legal requirements. The Investigator will inform the Sponsor about all

reports sent to the reporting organisation including follow-up information and answers by the

reporting organisation. The local investigator is responsible for informing other site principal

investigators and the CI of all SAEs.

The regulatory authority (MHRA) will be notified of all SAEs as soon as possible within ten days of

the event occurring during the study. The main REC will be notified of all unexpected and related

SAEs within 15 days of the occurrence of the event.

Switzerland only

The following events are reported to the Representative of the Sponsor and Principal Investigator

within 24 hours upon becoming aware of the event:

All SAEs and SADEs

Device deficiencies that might have potentially led to an SAE

Health hazards that require measures

Dr Lia Bally

Department of Diabetes, Endocrinology, Clinical Nutrition and Metabolism

Inselspital, Bern University Hospital

University of Bern, Bern

Switzerland

Tel: 0041 31 632 36 77

Email: lia.bally@insel.ch

SAEs will evaluated by the Representative of the Sponsor and Principal Investigator with regard to

causality and seriousness. Device deficiencies are assessed regarding their potential to lead to an

SAE.

10.2.5 Recording and reporting of device deficiencies

All device deficiencies will be documented throughout the study. The investigator will be responsible

for managing all device deficiencies and determine and document in writing whether they could have

led to a serious adverse device effect.

All device deficiencies that might have led to a serious adverse device effect(s) if: suitable action

had not been taken; intervention had not been made, or if circumstances had been less fortunate,

must be reported to the Sponsor as for SAEs/SADEs.

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10.2.6 Healthcare arrangements and compensation for adverse events

Healthcare arrangements for subjects who suffer an adverse event as a result of participating in the study may include advice from clinical members of the study team or the patient's treating diabetes team, or use of emergency health services.

If an adverse event occurs, there are no special compensation arrangements unless this was due to the negligence of one of the clinical investigators or due to harm resulting from study protocol design. In this case subjects may have grounds for legal action for compensation. The normal national complaints mechanism will be available. In addition, any harm arising due to study design (both negligent and non-negligent) will be covered under Sponsor's insurance policy as applicable.

10.2.6.1 Country specific requirements

- 1. UK The Investigator will notify the ethics committee of all Serious Adverse Events in line with pertinent legal requirements. The Investigator will inform the Sponsor about all reports sent to the ethics committee including follow-up information and answers by the ethics committee. The MHRA and REC will be notified of all SUSARs occurring during the study according to the following timelines: fatal and life-threatening within 7 days of notification and non-life threatening within 15 days.
- 2. Switzerland The Representative of the Sponsor and Principal Investigator reports SAEs and SADEs in Switzerland within 7 days to the local Ethics Committee which are
 - related or possibly related to the medical device under investigation
 - related or possibly related to study procedures

The Representative of the Sponsor and Principal Investigator reports the above both locally and internationally, within 7 days to Swissmedic. In addition, all device deficiencies that could have led to a SADE in Switzerland and study centres abroad will be notified to Swissmedic and the local Ethics Committee within 7 days (ClinO Rt: 42)

Health hazards that require measures are reported to the Ethics Committee and Swissmedic within 2 days (ClinO Art. 37).

An annual safety report including adverse events and device deficiencies including local and international events will be provided to Ethics Committee and Swissmedic every year by the Representative of the Sponsor (ClinO Art. 43).

The Representative of the Sponsor and Principal Investigator notifies Swissmedic and Ethics Committee of the completion of the study within 90 days (as of last patient, last visit). A discontinuation or an interruption of the trial, and the reasons for this are notified within 15 days. A final report, with contents in accordance with EN ISO 14155, will be submitted within one year of completion to Swissmedic and Local Ethics Committee (ClinO Art 38).

10.3 Anticipated adverse events, risks and benefits

10.3.1 Risks and anticipated adverse events

Known risks represent hazardous situations which may result in anticipated adverse events. In the following text, where appropriate, the term "risk" and "anticipated adverse events" are used interchangeably without affecting meaning.

10.3.2 Hypoglycaemia and hyperglycaemia

Subjects with type 2 diabetes requiring insulin therapy have a pre-existing risk for hypoglycaemia and hyperglycaemia. Potential risks are:

- Risk of mild to moderate hypoglycaemia and associated symptoms such as sweating, trembling, difficulty thinking and dizziness. There is also a rare risk of severe hypoglycaemia when conscious level is altered, needing help from a third party to correct the hypoglycaemia. These risks are pre-existent in any patient with type 2 diabetes requiring insulin and the study objective is to develop systems to minimise these risks
- Risk of possible mild to moderate hyperglycaemia similar to the risk that a subject with type
 2 diabetes requiring insulin experiences on a daily basis

10.3.3 Finger-stick blood glucose measurements

Finger-stick tests may produce pain and/or bruising at the site.

10.3.4 Insulin injection therapy

Potential risks associated with multiple daily injection therapy include:

- Slight discomfort at the time of insulin injection (common)
- Slight bruising at the site of injection (common)
- Bleeding at injection site (rare)
- Infection at the site of injection (rare)
- Insulin pen malfunction and mechanical problems (rare)

- Allergy to insulin (very rare)
- Lipodystrophy / lipoatrophy (very rare)

10.3.5 Insulin pump therapy

Potential risks associated with insulin pump therapy include:

- Slight discomfort at the time of insertion of the insulin delivery cannula (common)
- Slight bruising at the site of insertion (common)
- Bleeding at insertion site (rare)
- Infection at the site of insertion (rare)
- Allergy to the insulin delivery cannula or adhesive (rare)
- Infusion set and cannula occlusions (rare)
- Insulin pump malfunction and mechanical problems (rare)
- Allergy to insulin (very rare)
- Lipodystrophy / lipoatrophy (very rare)

10.3.6 Continuous glucose monitoring

Potential risks associated with CGM:

- Slight discomfort at the time of insertion of CGM (common)
- Slight bruising at the site of insertion (unlikely)
- Bleeding at insertion site (rare)
- Infection at the site of insertion (rare)
- Allergic reaction to the CGM sensor material (rare)

If a skin reaction is classified as severe (the observation is noticeable and bothersome to subject and may indicate infection or risk of infection or potentially life-threatening allergic reaction), an adverse event form will be completed.

10.3.7 Dialysis related events

Potential risks associated with haemodialysis include:

- Intra-dialytic hypotension
- Muscle cramps
- Cannulation difficulties
- Clotting of the dialysis circuit

• Infiltration of the haemodialysis access

10.3.8 Questionnaires

As part of the study, subjects will complete questionnaires which include questions about their private attitudes, feelings and behaviour related to diabetes. It is possible that some people may find these questionnaires to be mildly upsetting. Similar questionnaires have been used in previous research and these reactions are uncommon.

10.3.9 Risk Analysis and residual risk associated with the investigational device

After in-depth analysis and consideration of all the potential hazards in relation to use of the CamAPS HX system in the home environment, it is concluded that the CamAPS HX system is safe, if used as intended.

Risk Assessment of the CamAPS HX system has been carried out in accordance with ISO 14971:2012. A preliminary Hazard Determination has been carried out including consideration of the questions in Annex C of ISO 14971:2012.

One hazard 'Hazard S16: "Unknown CGM accuracy in Infants/Children, Pregnant Women and those on dialysis treatment or with critical illness' is the only hazard identified that could not be reduced to an acceptable risk level, post mitigation. Our in-detail risk/benefit assessment concluded that the benefits of the system outweigh the risk with respect to this specific hazard.

As per our risk management process, further risk analysis shall be undertaken post production and release as to ensure any issues raised are acted upon to ensure the CamAPS HX system continues to improve and develop.

10.4 Benefits

It is expected that day and night closed loop system may have an important role in the management of diabetes in this patient group. The closed loop system may impact on the frequency of hypoglycaemia with suspected fewer low glucose levels with closed loop insulin delivery compared with usual care. In addition to this, higher blood glucose levels above target should be reduced with use of the closed loop algorithm. During the closed-loop period, participants will not need to self-

administer insulin, which may facilitate diabetes management. Therefore, participation in this study is likely to be beneficial for study participants.

As this will be a proof-of-concept study, it is possible that subjects will not directly benefit from being a part of this study. However, it is also possible that the blood glucose information from the CGM devices along with the information about insulin dosing during day and night closed loop will be useful for subjects' diabetes self-management.

10.5 Data Safety Monitoring Board (DSMB)

An independent Data Safety Monitoring Board (DSMB) will be informed of all serious adverse events and any unanticipated adverse device effects that occur during the study and will review compiled adverse event data at periodic intervals.

11 Methods and assessments

11.1 Procedures

11.1.1 Weight

Weight will be recorded at the study initiation visit at baseline and before and after each dialysis session. Weight will be measured in kilograms using a calibrated electronic scale.

11.1.2 Blood Glucose Meter Data

The blood glucose meter will be downloaded periodically during the duration of the study.

11.2 Assessment for Safety

Subjects will receive education about treating hypo- and hyperglycaemia during closed loop insulin delivery. If the low glucose alert from the CGM becomes activated in closed-loop, a capillary glucose sample will also be measured. They will be advised to treat any finger stick capillary glucose level below 4 mmol/l with quick acting carbohydrate. Written guidelines in keeping with subjects' usual treatment guidelines will be provided for dealing with hypo- and hyperglycaemia.

11.3 Assessment for Efficacy

Continuous glucose monitoring during the study for 20 days in each arm will be used in each participant to assess for efficacy.

Two continuous glucose monitoring systems (CGM) will be used throughout this study: a blinded CGM with retrospective sensor glucose data read out, and a real-time system providing a contemporaneous display of sensor readings.

Blinded CGM will be used throughout the control arm. Sensors will be returned to the research team thereafter. Secondary glucose endpoints as outlined in 10.1 will be based on glucose data derived from data captured during this 20 day period.

Real-time CGM will be applied during closed loop intervention. The control algorithm will use the real-time CGM's continuous stream of glucose data to control insulin titration.

11.4 Questionnaires

Quantitative data on health-related quality of life will be assessed using validated questionnaires. Participants will complete a series of questionnaires including an evaluation of their experience and views of the diabetes treatment received. It is estimated that participants will take 10-15 minutes to complete the questionnaires. All results will be evaluated at the end of the study.

Questionnaires will include:

- Problem Areas in Diabetes Questionnaire
- Hypoglycaemia Fear Survey (HFS)
- Closed-loop experience questionnaire

12 Study materials and products

12.1 Insulin

During the control intervention and wash out period, subcutaneous insulin therapy will be administered using CE-marked insulin pen devices as per usual clinical practice.

During closed-loop intervention, faster acting insulin aspart (Fiasp, Novo Nordisk, Copenhagen, Denmark) will be administered via an insulin pump as described below (see 12.3).

12.2 Multiple daily insulin injections during control intervention and wash out period

During the control intervention when multiple daily injection therapy will be applied, insulin will be administered using CE-marked insulin pen devices as per usual clinical practice.

12.3 Insulin pump

During day and night automated closed loop glucose control the Dana insulin pump (SOOIL) or similar CE-marked insulin pump will be used.

12.4 Continuous subcutaneous glucose monitor

The Dexcom G6 real-time sensor with sensor applicator (Dexcom, Northridge, CA, USA) will be the study CGM. The sensor will be calibrated according to manufacturer's instructions.

12.5 Blood Glucose Meter

Study participants will use their own approved glucose meter for self-monitoring of capillary blood glucose (SMBG) during the study. The capillary glucose meter readings may be used to calibrate the sensor according to manufacturer's instructions.

12.6 Computer-based algorithm

The Cambridge closed loop controller has been used safely and effectively in the closed loop studies in both children and adults with T1D (study REC Ref. 06/Q0108/350, REC Ref. 07/H0306/116, REC Ref. 08/H0304/75, REC Ref. 08/H0308/297, REC Ref. 09/H0306/44, REC Ref. 10/H0304/87, REC Ref. 12/EE/0155, REC Ref. 12/EE/0034, REC Ref. 12/EE/0424, REC Ref. 13/EE/0120, REC Ref. 13/WM/0498, REC Ref. 13/EE/0251, REC Ref. 13/EE/0321, REC Ref. 13/EE/0018, REC Red 15/EE/0324, REC ref 16/EE/0286, REC ref 16/EE/0380 and REC Ref 17/LO/0576).

13 Data Analysis

13.1 Primary Endpoint Analysis

The primary analysis will evaluate the between group difference in time spent in the target glucose range from 5.6 to 10.0 mmol/l based on CGM glucose levels during the 20 day intervention periods.

Mean ± SD or summary statistics appropriate to the distribution will be reported for the primary endpoint over the 20 day period by treatment intervention. The treatment interventions will be

compared using a linear mixed model. A 95% confidence interval will be reported for the difference between the interventions based on the linear mixed model.

Residual values will be examined for an approximate normal distribution. If values are highly skewed, then a ranked normal score transformation will be used instead. However, previous experience suggests that the primary endpoint will follow an approximately normal distribution.

A 5% significance level will be used to declare statistical significance for the primary comparison. A two-sided p-value will be reported.

The primary analysis will be a single statistical comparison of a single outcome measure. No formal corrections for multiple comparisons will be performed for the key or secondary analyses in this study.

The primary analysis will be performed on an intention-to-treat basis using the treatment group assigned by randomization.

A per-protocol analysis restricted to participants with a minimum of 60% CGM data during control period and 60% use of closed-loop during closed-loop period will be conducted for the primary endpoint.

Primary endpoint hypotheses

- Null Hypothesis: There is no difference in the true mean time spent in the target range
 (5.6 to 10.0 mmol/L) over the 20 day period between the two treatment groups.
- Alternative Hypothesis: There is a nonzero difference in the true mean time spent in the target range over the 20 day period between the two treatment groups.

13.2 Other Key Endpoints

For the following key endpoints will be assessed:

- Time spent above target glucose (10.0 mmol/l)
- Average of glucose levels
- Time spent with glucose levels <3.9 mmol/l

13.3 Secondary Endpoints

The following endpoints will be considered exploratory:

CGM derived indices:

- Percentage time spent at glucose <5.6 mmol/l to quantify time spent below target
- Percentage time spent at glucose <3.0 mmol/l
- Percentage time spent at glucose > 20.0 mmol/l
- Standard deviation of glucose to quantify the glucose variability
- Number of hypoglycaemia events (glucose < 3.5mmol/l)

Insulin and Other Endpoints:

- Total amount of insulin delivered
- Average inter-dialytic weight gain

For all secondary endpoints, summary statistics appropriate to the distribution will be tabulated by treatment group. Analysis of key endpoint, and all secondary CGM and insulin endpoints will parallel the primary analysis. A ranked normal score transformation will be applied to all highly skewed secondary endpoints.

Summary statistics for the following outcome metrics will also be tabulated separately for daytime (defined as 8am to less than 12am) and night time (defined as 12am to less than 8am), and dialysis days and non-dialysis days over the 20 day period:

- Percent time with glucose levels spent in the target range (5.6 to 10.0 mmol/L)
- Mean of glucose levels
- Standard deviation of glucose levels
- Percent time with glucose levels below 3.9 mmol/L
- Total insulin dose

13.4 Safety Evaluation

For each of the following safety outcomes, mean \pm SD or summary statistics appropriate to the distribution will be tabulated by treatment group:

- Number of subjects with any severe hypoglycaemia events
- Number of episodes of severe hypoglycaemia events per subject and incidence rate per 100 person years
- Number of adverse events per subject
- Number of serious adverse events per subject

For purposes of analysis, a severe hypoglycaemic event will be defined as an event requiring assistance of another person actively to administer carbohydrate, glucagon, or other resuscitative actions. These episodes may be associated with sufficient neuroglycopaenia to induce seizure or 261016_Protocol V1.1_12 June 2019

coma. If plasma glucose measurements are not available during such an event, neurological recovery attributable to the restoration of plasma glucose to normal is considered sufficient evidence that the event was induced by a low plasma glucose concentration.

All of the above safety outcomes will be tabulated for all subjects (including dropouts and withdrawals), regardless of whether CGM data are available and irrespective of whether closed loop was operational. All adverse events will be listed for the entire study duration, including washout period.

For severe hypoglycaemia (if enough events), the event rates will be compared using a repeated measures Poisson regression model.

13.5 Utility Evaluation

The amount of CGM use will be tabulated for each treatment arm, in addition to the amount of closed loop system use in the CL arm. Summary statistics appropriate to the distribution and range will be reported for the percentage of time using the CGM over the 20 day period for each treatment group. The same will be done for the percentage of time using the closed loop system in the CL arm. Tabulations of summary statistics will also be performed for the percentage of time spent using the closed-loop system while using the CGM in the CL arm.

13.6 Questionnaires

Descriptive tabulations of questionnaires will be carried out, and scores will be calculated using provided scaling and scoring tools as appropriate.

13.7 Evaluative periods

Where appropriate, sensor based measures will also be calculated for day and night-time periods and dialysis days and non-dialysis days. The interval from 8.00 to 24:00 defines day-time period, 00:00 to 08:00 am defines the night-time period. The primary and secondary measures will be calculated from day 1 until the end of each 20 day study intervention.

13.8 Interim monitoring and analyses

No formal interim analysis will be performed.

13.9 Sample size and power calculations

This is an exploratory analysis involving 32 subjects with at least 48 hours of data. Previous studies in people with type 2 diabetes may not provide reliable information about the within group variability

in this particular population requiring maintenance renal replacement therapy. No formal power calculations thus apply.

Allowing for 20% loss to follow up means we would need a total of 40 randomised participants.

14 Case Report Forms

The Case Report Form (CRF) is the printed, optical, or electronic document designed to record all the protocol required information to be reported to the Chief Investigator for each study participant.

CRFs will be completed in accordance with GCP and ISO 14551 Guidelines. Corrections to the CRF will be performed by striking through the incorrect entry and by writing the correct value next to the data that has been crossed out; each correction will be initialled and explained (if necessary) by the Investigator or the Investigator's authorised staff.

If any amendments to the protocol or other study documents are made, CRFs will be reviewed to determine if an amendment to these forms is also necessary.

15 Data Management

Confidentiality of subject data shall be observed at all times during the study. Personal details for each subject taking part in the research study and linking them to a unique identification number will be held locally on a study screening log in the Trial Master File at the investigation centre. These details will not be revealed at any other stage during the study, and all results will remain anonymous (in Switzerland, data will be considered coded).

Case report forms (CRFs) will be used for recording anonymised (in Switzerland, data will be considered coded) study data. CRFs will be completed in accordance with GCP and ISO 15197: 2013 Guidelines. The study identification number will be used on CRF. Names and addresses will not be used.

Electronic data will be stored on password-protected computers. All paper records will be kept in locked filing cabinets, in a secure office at the investigation centre. Only members of the research team and collaborating institutions will have password access to the anonymised (in Switzerland, data will be considered coded) electronic data. Only members of the research teams will have

access to the filing cabinet. All data will be stored for at least 15 years in line with the General Data Protection Regulation (GDPR) (EU) 2016/679. In case of withdrawal of participants, data that were obtained before will be further used anonymised (in Switzerland, data will be considered coded).

Direct access to the source data will be provided for monitoring, audits, REC review and regulatory authority inspections during and after the study. The fully anonymised data may be shared with third parties (EU or non-EU based) for the purposes of advancing management and treatment of diabetes.

Appropriate procedures agreed by the Chief Investigator and Clinical Principal Investigators will be put in place for data review, database cleaning and issuing and resolving data queries.

15.1 Further information on data management – Switzerland only

15.1.1 Hardware and software

Data will be managed using a research cluster with three servers. The cluster provides enhanced availability, reliability and scalable performance. The servers are located in locked dedicated server rooms with restricted access.

Two dedicated virtual machines are installed for research data management with REDCap – one productive instance and one test instance. The servers are running on Microsoft Windows Server and using Apache HTTP Server.

Study data will be collected and managed using the REDCap (Research Electronic Data Capture) software. REDCap is a secure, web-based application designed to support data capture for research studies, providing 1) an intuitive interface for validated data entry; 2) audit trails for tracking data manipulation and export procedures; 3) automated export procedures for seamless data downloads to common statistical packages; and 4) procedures for importing data from external sources [https://projectredcap.org/resources/ citations/]. The REDCap tools are hosted at the Department of Anaesthesiology and Pain Medicine, University Hospital Bern.

15.1.2 Data security, access and backup

The servers are behind a firewall and cannot be accessed through the internet. Furthermore, Apache HTTP Server and REDCap were configured to run under Secure Sockets Layer (SSL) hence data is being encrypted and is transmitted securely.

Available disk space is monitored actively. If free disk space is less than 10 percent then administrators get an email and more storage capacity will be added accordingly.

The application has a group and role-based security model. Each user belongs to one or more security groups with specific sets of permissions in relation to folder or projects in the system. Only dedicated site administrators have access to the admin console enabling user management and changing security settings.

The system can only be accessed entering a user name and password. All events are recorded in the user event list of the audit log files.

All servers are regularly backed up on storage servers in a separate server room using a multi-level system.

15.1.3 Analysis and archiving

REDCap provides data analysis by integrated tools for creating reports and charts.

All data can be exported in different formats (Microsoft Excel, CSV, PDF, SAS, Stata, R, SPSS) suitable for transfer to a statistical software package of choice. All data will be archived and secured in the database as long as required by legislation.

16 Ethics

The study will be conducted in accordance with the Declaration of Helsinki Ethical Principles for Medical Research involving Human Subjects (64th WMA General Assembly, Fortaleza, Brazil, October 2013).

16.1 Research Ethics Committee and Institutional Review Board

Prior to commencement of the study, the protocol, any amendments, subject information and informed consent and assent forms, any other written information to be provided to the subject, subject recruitment procedures, current investigator CVs, and any other documents as required by the Research Ethics Committee (REC) or Institutional Review Board will be submitted. Written approval will be obtained from the REC prior to the commencement of the study. Any additional requirements imposed by the REC or regulatory authority shall be followed.

16.2 Informed consent of study subjects

In obtaining and documenting informed consent, the investigator will comply with the applicable regulatory requirements and will adhere to GCP standards and to the ethical principles that have their origin in the Declaration of Helsinki. Prior to the start of the study, the Investigator will obtain

favourable ethical opinion of the written informed consent form and any other written information to be provided to subjects.

Subjects will be given full verbal and written information regarding the objectives and procedures of the study and the possible risks involved. The study team will avoid any coercion or undue improper inducement of the subject to participate and subjects will be given ample time to consider participation in the study.

All subjects will have the right to leave the study at any time, without stating any reason, and without any negative consequences to their subsequent medical treatment. The subject will be informed in a timely manner should any new information become available during the course of the study that may affect their well-being, safety and willingness to participate in the study.

Written consent will be obtained from participants according to REC requirements. The signed informed consent forms will be photocopied, originals filed in the Investigator's Site File, a copy placed in the patient's notes and a copy given to the subjects. All subjects will receive a copy of the informed consent form, and the Project Coordinator's office will hold copies of the consent forms and Ethics Committee approvals and make them available upon request.

17 Amendments to the protocol

Any substantial amendments to the protocol and other documents shall be notified to, and approved by, the Research Ethics Committee or Institutional Review Board, and the regulatory authority, prior to implementation as per nationally agreed guidelines.

18 Deviations from the protocol

Deviations from the protocol should not occur without prior approval of the REC or sponsor except under emergency circumstances, to protect the rights, safety and well-being of subjects. If deviations do occur, they will be documented, stating the reason and the date, the action taken, and the impact for the subject and for the study. The documentation will be kept in the Investigator's Site File.

Deviations affecting the subject's rights, safety and well-being or the scientific integrity of the study will be reported to the REC and sponsor as soon as possible in a timely manner, following nationally-agreed guidelines.

19 Study management

19.1 Data and Safety Monitoring Board (DSMB)

An independent Data Safety Monitoring Board (DSMB) will comprise an independent chairperson and two external experts. The DSMB aims to safeguard the interests of trial participants, assess the safety and efficacy of the interventions during the trial, and monitor the overall conduct of the clinical trial.

The DSMB should receive and review the progress and accruing data of the project clinical trials and provide advice on the conduct of the trial. The DSMB will be informed of all serious adverse events and any unanticipated adverse device effects that occur during the study and will review compiled adverse event data at periodic intervals.

19.2 Trial Steering Committee (TSC)

A trial steering committee (TSC) will supervise the trial, to ensure it is conducted to high standards in accordance with the protocol, the principles of GCP, and with regard to participant safety. This committee will consist of the Chief Investigator and Clinical Investigators.

The TSC will meet (in person or conference call) at regular intervals during active phase, and at the conclusion of the study. The TSC will consider the study and relevant information from other sources, ensuring at all times that ethical considerations are met when recommending the continuation of the trial.

20 Responsibilities

20.1 Chief Investigator

The Chief Investigator (CI) is the person with overall responsibility for the research and all ethical applications will be submitted by the CI. The CI is accountable for the conduct of the study and will ensure that all study personnel are adequately qualified and informed about the protocol, any amendments to the protocol, the study treatments and procedures and their study related duties. The CI should maintain a list of appropriately qualified persons to whom he/she has delegated specified significant study-related duties.

20.2 Principal Clinical Investigator

The Principal Clinical Investigator will be responsible for the day-to-day conduct of the clinical aspects of the study.

21 Reports and Publications

Data will be submitted for publication in internationally peer-reviewed scientific journals; members of the investigator group will all be co-authors. The privacy of each subject and confidentiality of their information shall be preserved in reports and publication of data.

22 Timetable

Inclusion of the first subject in the study is planned to take place in April 2019. The expected completion of the last subject is December 2019 and the planned completion of the Clinical Study Report is March 2020.

23 Retention of Study Documentation

Subject notes must be kept for the maximum time period as permitted by each relevant institution. Other source documents and the Investigator's Site File must be retained for at least 15 years, in line with the General Data Protection Regulation (GDPR) (EU) 2016/679. The Principal Investigator will archive the documentation pertaining to the study after completion or discontinuation of the study.

Switzerland only

The Representative of the Sponsor and Principal Investigator will retain all data pertaining for a minimum of 10 years after completion or discontinuation of the trial (Art 45 ClinO of HRA). Access will be restricted to research team members, Swissmedic and Ethics Committee.

24 Indemnity statements

The clinical investigators are indemnified to cover negligent harm to patients participating in the study by their membership of medical defence organisations. Indemnity for any harm arising from the conduct of research will be provided according to local arrangements in respective centre.

- 1. Cambridge, UK National Health Service indemnity cover will apply for any claims arising from management and conduct of research. Any liability arising from study design will be covered by the clinical trial insurance policy organised by the University of Cambridge.
- 2. Bern, Switzerland Study insurance will be provided by the University Hospital of Bern. A copy of the certificate will be filed in the investigator site file and the trial master file.

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26 Document amendment history

Version number	Date	Amendment information
1.1	12 June 2019	 Inselspital, Bern University Hospital logo added Switzerland funder information added Language requirements in Switzerland updated Recruitment in Switzerland clarified Requirement for women of reproductive age to take a pregnancy test and use contraception during the study in Switzerland Country specific requirements in Switzerland regarding safety reporting and data management