

## **Clinical trial results:**

A randomized, double-blinded, placebo-controlled trial that evaluates the effect of empagliflozin on oxidative stress in patients with type 2 diabetes

EudraCT number	2016-000370-38
Trial protocol	DK
Global end of trial date	22 January 2020
Result version number	v1 (current)
This version publication date	04 February 2021
First version publication date	04 February 2021
Sponsor protocol code	2016-100
Sporisor protocor code	2010-100
ISRCTN number	-
ClinicalTrials.gov id (NCT number)	NCT02890745
WHO universal trial number (UTN)	-
Notes:	
Sponsor organisation name	Department of Clinical Pharmacology, Bispebjerg- Frederiksberg Hospital - Rigshospitalet, Q7642
Sponsor organisation address	Blegdamsvej 9, Copenhagen, Denmark,
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Notes:	
Is trial part of an agreed paediatric investigation plan (PIP)	No
Does article 45 of REGULATION (EC) No 1901/2006 apply to this trial?	No
Does article 46 of REGULATION (EC) No 1901/2006 apply to this trial?	No

Notes:	
Analysis stage	Final
Analysis stage	
Date of interim/final analysis	01 November 2020 Yes
Is this the analysis of the primary completion data?	res
Primary completion date	22 January 2020
Global end of trial reached?	Yes
Global end of trial date	22 January 2020
Was the trial ended prematurely?	Yes
Notes:	
Martin all trades at 100 at 100	
Main objective of the trial:	
	ydroguanosine (8-oxoGuo) and 8-oxo-7,8-dihydro-2'- fter treatment with empagliflozin compared to placebo patients with type 2 diabetes
Protection of trial subjects:	
All data was stored in accordance with the	ne Danish Law of data protection
Background therapy: -	
Evidence for comparator: -	
Actual start date of recruitment	23 November 2016
Long term follow-up planned	No
Independent data monitoring committee (IDMC) involvement?	Yes
Notes:	
Country: Number of subjects enrolled	Denmark: 35
Worldwide total number of subjects	35
EEA total number of subjects	35
Notes:	
	T <sub>a</sub>
In utero	0
Preterm newborn - gestational age < 37 wk	0
Newborns (0-27 days)	0
Infants and toddlers (28 days-23 months)	0
Children (2-11 years)	0
Adolescents (12-17 years)	0
Adults (18-64 years)	18
From 65 to 84 years	17
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85 years and over	0

## Recruitment details:

Recruitment were done in relation to patients outpatient consultation at Gentofte Hospital and Bispebjerg Hospital as well as throug advertisements at public available material.

## Screening details:

A screening visit is performed prior to enrollment in the study to ensure elibility of study criteria with an physical examination and disease and life style history

Period 1 title	Overall trial (overall period)
Is this the baseline period?	Yes
Allocation method	Randomised - controlled
Blinding used	Double blind
Roles blinded	Subject, Investigator, Monitor, Data analyst, Assessor
Are arms mutually exclusive?	Yes
	Intervention
Arm description:	
Active arm. Empagliflozin 25 mg, once d	aily for 14 days
Arm type	Experimental
Investigational medicinal product name	empagliflozin
Investigational medicinal product code	
Other name	
Pharmaceutical forms	Tablet
Routes of administration	Oral use
Dosage and administration details:	
25 mg, tablet, oral use	
	Placebo
Arm description:	
Placebo	
Arm type	Placebo
Investigational medicinal product name	Placebo
Investigational medicinal product code	
Other name	
Pharmaceutical forms	Tablet
Routes of administration	Oral use

Dosage and administration details:

Placebo, tablet, oral use

	Intervention	Placebo	
Started	16	15	
Completed	16	15	

## Notes:

[1] - The number of subjects reported to be in the baseline period are not the same as the worldwide number enrolled in the trial. It is expected that these numbers will be the same.

Justification: A total number of 35 participants were enrolled in the study. However, 4 participants chose to withdrew before getting trial medicine, and were thus not included in the baseline information.

Reporting group title	Overall trial

Reporting group description: -

	Overall trial	Total	
Number of subjects	31	31	
Age categorical			
Units: Subjects			
In utero		0	
Preterm newborn infants (gestational age < 37 wks)		0	
Newborns (0-27 days)		0	
Infants and toddlers (28 days-23 months)		0	
Children (2-11 years)		0	
Adolescents (12-17 years)		0	
Adults (18-64 years)		0	
From 65-84 years		0	
85 years and over		0	
Age continuous			
Mean: 63 SD: 7			
Units: years			
arithmetic mean	63		
standard deviation	± 7	-	
Gender categorical			
Units: Subjects			
Female	0	0	
Male	31	31	
Body mass index			
Units: kg/m^2			
arithmetic mean	30.9		
standard deviation	± 4.6	-	
HbA1c			
Units: mmol/mol			
arithmetic mean	56		
standard deviation	± 7	-	

Reporting group title Intervention			
Reporting group description:			
Active arm. Empagliflozin 25 mg, once daily for 14 days			
Reporting group title	Placebo		
Reporting group description:			
Placebo			

End point title	Changes in urinary excretion of 8-oxo-7,8-dihydroguanosine
End point description:	
End point type	Primary
End point timeframe:	
Changes from baseline to end of study	

	Intervention	Placebo	
Subject group type	Reporting group	Reporting group	
Number of subjects analysed	16	15	
Units: nmol/24h			
arithmetic mean (standard deviation)	-0.5 (± 7.8)	-1.8 (± 8.5)	

	Between group effects 8-oxo-7,8-dihydroguanosine
Comparison groups	Placebo v Intervention
Number of subjects included in analysis	31
Analysis specification	Pre-specified
Analysis type	equivalence
P-value	= 0.66
Method	t-test, 2-sided
Parameter estimate	Mean difference (final values)
Point estimate	1.3
Confidence interval	
level	95 %
sides	2-sided
lower limit	-4.2
upper limit	4.8

	Intervention	Placebo	
Subject group type	Reporting group	Reporting group	
Number of subjects analysed	16	15	
Units: nmol/mL			
arithmetic mean (standard deviation)	-0.13 (± 0.67)	0.08 (± 0.54)	

	Between group effect malondialdehyde
Comparison groups	Intervention v Placebo
Number of subjects included in analysis	31
Analysis specification	Pre-specified
Analysis type	equivalence
P-value	= 0.33
Method	t-test, 2-sided
Parameter estimate	Mean difference (final values)
Point estimate	-0.22
Confidence interval	
level	95 %
sides	2-sided
lower limit	-0.66
upper limit	0.23

Timeframe for reporting adverse events:

During the study period (14 days) and one week afterwards

Adverse event reporting additional description:

Daily text messages and at the end of study. If any occured the week following the study, participants contacted the investiggator

Assessment type Systematic

Dictionary name	MedDRA
Dictionary version	23.1

Reporting group title	Intervention group	
Reporting group description:		
Empagliflozin 25 mg, once daily for 14 days		
Reporting group title	Placebo	

Reporting group description:

Placebo for 14 days

	Intervention group	Placebo	
Total subjects affected by serious adverse events			
subjects affected / exposed	0 / 16 (0.00%)	0 / 15 (0.00%)	
number of deaths (all causes)	0	0	
number of deaths resulting from adverse events	0	0	

Frequency threshold for reporting non-serious adverse events: 1 %

	Intervention group	Placebo	
Total subjects affected by non-serious adverse events			
subjects affected / exposed	10 / 16 (62.50%)	6 / 15 (40.00%)	
Gastrointestinal disorders			
Increased thirst			
subjects affected / exposed	1 / 16 (6.25%)	0 / 15 (0.00%)	
occurrences (all)	1	0	
Nausea or changed stool consistence			
subjects affected / exposed	0 / 16 (0.00%)	2 / 15 (13.33%)	
occurrences (all)	0	2	
Reproductive system and breast disorders			

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Genital fungal infection			
subjects affected / exposed	1 / 16 (6.25%)	0 / 15 (0.00%)	
occurrences (all)	1	0	
Denal and uninamy discurdant			
Renal and urinary disorders			
Polyuria			
subjects affected / exposed	9 / 16 (56.25%)	1 / 15 (6.67%)	
occurrences (all)	9	1	
Psychiatric disorders			
Tired			
subjects affected / exposed	0 / 16 (0.00%)	2 / 15 (13.33%)	
occurrences (all)	0	2	
Musculoskeletal and connective tissue disorders			
Sprained finger			
subjects affected / exposed	1 / 16 (6.25%)	0 / 15 (0.00%)	
occurrences (all)	1	0	
Infections and infestations			
Upper respiratory tract infection			
subjects affected / exposed	0 / 16 (0.00%)	2 / 15 (13.33%)	
occurrences (all)	0	2	

Were there any global substantial amendments to the protocol? No

Were there any global interruptions to the trial? No

Limitations of the trial such as small numbers of subjects analysed or technical problems leading to unreliable data.

The project did not include 34 participants that succesfully completed the study. 35 partipants were enrolled, however, 4 participants withdrew before starting trial medicine. Thus, 31 participants succesfully completed the study.

Notes: