National Institute for Health and Care Excellence

Draft for consultation

Type 2 diabetes in adults: management (medicines update)

[E1.3] Evidence reviews for initial pharmacological management of type 2 diabetes: appendix D studies L to Z

NICE guideline GID-NG10336

Evidence reviews underpinning recommendations 1.8.6-1.8.32, 1.8.34, 1.8.38-1.8.60 and recommendations for research in the NICE guideline

August 2025

Draft for Consultation

This evidence review was developed by NICE



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ISBN:

Note: In the study characteristics tables, if any baseline characteristic is not mentioned in a table, then this is because the value was either not reported by the study or not reported in a way that could be meaningfully extracted by the analyst assigned to review the study and so was not reported in the data extraction.

67. Lambadiari, 2018

Bibliographic Reference

Lambadiari, V.; Pavlidis, G.; Kousathana, F.; Varoudi, M.; Vlastos, D.; Maratou, E.; Georgiou, D.; Andreadou, I.; Parissis, J.; Triantafyllidi, H.; et, al.; Effects of 6-month treatment with the glucagon like peptide-1 analogue liraglutide on arterial stiffness, left ventricular myocardial deformation and oxidative stress in subjects with newly diagnosed type 2 diabetes;

Cardiovasc Diabetol; 2018; vol. 17 (no. 1); 8

67.1. Study details

07.1. 3	tudy details
Secondary publication of another included study- see primary study for details	No additional information.
Other publications associated with this study included in review	No additional information.
Trial name / registration number	NCT03010683.
Study type	Randomised controlled trial (RCT)
Study location	Greece.
Study setting	Outpatient follow-up.
Study dates	No additional information.
Sources of funding	No funding.
Inclusion criteria	Newly diagnosed and treatment-naïve people with type 2 diabetes.
Exclusion criteria	History or clinical evidence of coronary or valvular heart disease; liver or kidney failure; history of alcohol or drug abuse; treatments able to modify glucose metabolism. All women were premenopausal and their investigations were undertaken during the first week of their menstrual cycles. None were taking oral contraceptives.
Recruitment / selection of participants	No additional information.
Intervention(s)	Liraglutide N=30

	Liraglutide 1.8mg once daily (with weekly dose escalation as instructed) as a subcutaneous injection for 6 months.
	Concomitant therapy: No additional information.
Strata 1: People with type 2 diabetes mellitus and heart failure	People without heart failure
Strata 2: People with atherosclerotic cardiovascular diseases	People without other cardiovascular diseases
Strata 3: People with type 2 diabetes mellitus and chronic kidney disease	People without chronic kidney disease
Strata 4: People with type 2 diabetes mellitus and high cardiovascular risk	People at higher risk of developing cardiovascular disease Based on smoking, hypertension, dyslipidaemia, family history, age, BMI and presence of diabetes.
Subgroup 1: People with frailty	Not stated/unclear
Subgroup 2: Onset of type 2 diabetes mellitus	Not stated/unclear
Subgroup 3: People with non-alcoholic fatty liver disease	People without non-alcoholic fatty liver disease
Subgroup 4: People with obesity	Not stated/unclear
Subgroup 5: eGFR category at baseline	Not stated/unclear

Subgroup 6: Albuminuria category at baseline	Not stated/unclear
Sensitivity analysis category: Enrichment trial status	5) All treatment naïve
Population subgroups	No additional information.
Comparator	Metformin N=30 Metformin 1000mg twice daily for 6 months. Concomitant therapy: No additional information.
Number of participants	60
Duration of follow-up	6 months.
Indirectness	No additional information.
Method of analysis	Not stated/unclear
Additional comments	No additional information.

67.2.1. **Liraglutide (N = 30)**

Liraglutide 1.8mg once daily (with weekly dose escalation as instructed) as a subcutaneous injection for 6 months. Concomitant therapy: No additional information.

67.2.2. Metformin (N = 30)

Metformin 1000mg twice daily for 6 months. Concomitant therapy: No additional information.

67.3. Characteristics

67.3.1. Arm-level characteristics

67.3.1. Arm-level characteristi	CS	
Characteristic	Liraglutide (N = 30)	Metformin (N = 30)
% Male	n = 20 ; % = 67	n = 20 ; % = 67
Sample size		0,
Mean age (SD) (years)	51 (10)	50 (12)
Mean (SD)		(,
Ethnicity	n = NR ; % = NR	n = NR ; % = NR
Sample size		,
Comorbidities	n = NA ; % = NA	n = NA ; % = NA
Sample size		,
Hypertension	n = 17; % = 56.7	n = 16; % = 53.3
Sample size		,
Dyslipidaemia	n = 16; % = 53.3	n = 15 ; % = 50
Sample size		·
Family history of coronary artery disease	n = 6; % = 20	n = 6; % = 20
Sample size		
Presence of frailty	n = NR ; % = NR	n = NR ; % = NR
Sample size		
Time since type 2 diabetes diagnosis	NR (NR)	NR (NR)
Mean (SD)		, ,
HbA1c (%)	8.6 (2)	8.4 (1.2)
Mean (SD)		. ,
Blood pressure (mmHg)	NA (NA)	NA (NA)
Mean (SD)		, ,
Systolic blood pressure	142 (15)	142 (19)
Mean (SD)		. ,
Diastolic blood pressure	90 (8)	89 (9)
Mean (SD)		, ,
Heart rate (beats per minute)	74 (12)	71 (12)

Characteristic	Liraglutide (N = 30)	Metformin (N = 30)
Mean (SD)		
Smoking status	n = 11; % = 36.7	n = 11 ; % = 36.7
Sample size	ND 0/ ND	
Alcohol consumption	n = NR ; % = NR	n = NR ; % = NR
Sample size		
Presence of severe mental illness	n = NR ; % = NR	n = NR ; % = NR
Sample size		
People with significant cognitive impairment	n = NR ; % = NR	n = NR ; % = NR
Sample size	- ND - 0/ ND	
People with a learning disability Sample size	n = NR ; % = NR	n = NR ; % = NR
	09 (16)	
Weight (kg)	98 (16)	78 (12)
Mean (SD)	00.0 (5)	
BMI (kg/m2)	32.9 (5)	27.7 (2)
Mean (SD)		
Number of people with obesity	n = NR ; % = NR	n = NR ; % = NR
Sample size	ND (ND)	
Cholesterol and lipid levels	NR (NR)	NR (NR)
Mean (SD)	()	
Albumin creatinine ratio	NR (NR)	NR (NR)
Mean (SD)		, ,
eGFR (mL/min/1.73m2)	85 (8)	83 (11)
Mean (SD)		
Other antidiabetic medication used	n = NR ; % = NR	n = NR ; % = NR
Sample size		
Blood pressure-lowering medication used	n = NA ; % = NA	n = NA ; % = NA
Sample size		
Beta blockers	n = 5; % = 16.7	n = 6; % = 20
Sample size		

Characteristic	Liraglutide (N = 30)	Metformin (N = 30)
Calcium antagonists	n = 10; % = 33.3	n = 8; % = 26.7
Sample size		
ACE inhibitors/ARBs	n = 9; % = 30	n = 10 ; % = 33.3
Sample size		
Diuretics	n = 4; % = 13.3	n = 4; % = 13.3
Sample size		
Statins/lipid-lowering medication used	n = 14; % = 46.7	n = 12 ; % = 40
Sample size		
Other treatment being received	n = NR ; % = NR	n = NR ; % = NR
Sample size		

68. Lee, 2013

Bibliographic Reference

Lee, Ji; Hong, Soon; Jeong, Han Saem; Joo, Hyung; Park, Jae; Chul-Min, Ahn; Yu, Cheol; Lim, Do-Sun; Effects of a PPAR-γ (Peroxisome Proliferator-Activated Receptor-gamma) Activator on Flow-Mediated Brachial Artery Dilation and Circulating Level of microRNA-21 in Hypertensive Type 2 Diabetic Patients; Journal of the Korean Society of Hypertension; 2013; vol. 19; 99

33111	tady actains
Study location	Korea
Study setting	University hospital cardiovascular centres
Study dates	Recruitment: July 2011 - June 2012
Sources of funding	Grant from Korean Society of Hypertension.
Inclusion criteria	 - 45-75 years old - Essential hypertension - Type 2 diabetes - sitting diastolic bp >= 80 mmHg - sitting systolic bp <= 130 mmHg -previously untreated hypertensive diabetes
Exclusion criteria	 use of pioglitazone within 3 months of enrolment SiSBP > 180 mmHg SiDBP > 110 mmHg heart failure (ejection fraction <45% or signs of heart failure) hepatic dysfunction (serum asparate or alanine aminotransferase levels being above twice the upper limit of normal ranges serum creatinine > 2.0 mg/dl pregnant, breastfeeding, "childbearing potential"
Recruitment / selection of participants	Eligible patients (n = 50, 20 women and 30 men) were randomly assigned to receive either pioglitazone 15 mg (25 patients) or control (25 patients) after measuring baFMD.
Intervention(s)	
Cointervention	

Strata 1: People with type 2 diabetes mellitus and heart failure	People without heart failure
Strata 2: People with atherosclerotic cardiovascular diseases	Not stated/unclear
Strata 3: People with type 2 diabetes mellitus and chronic kidney disease	Not stated/unclear
Strata 4: People with type 2 diabetes mellitus and high cardiovascular risk	Not stated/unclear
Subgroup 1: People with frailty	Not stated/unclear
Subgroup 2: Onset of type 2 diabetes mellitus	Not stated/unclear
Subgroup 3: People with non-alcoholic fatty liver disease	Not stated/unclear
Subgroup 4: People with obesity	Not stated/unclear
Subgroup 5: eGFR category at baseline	Not stated/unclear
Subgroup 6: Albuminuria category at baseline	Not stated/unclear

Sensitivity analysis category: Enrichment trial status	5) All treatment naïve
Population subgroups	
Comparator	"Placebo"
Number of participants	50
Duration of follow-up	6 months
Indirectness	Directly applicable
Method of analysis	ITT

68.3. Characteristics

68.3.1. Arm-level characteristics

Characteristic	Pioglitazone (N = 25)	Placebo (N = 25)
Age (yr)	61.4 (12.4)	59.5 (11.8)
Mean (SD)		
% Male (%)	n = 16 ; % = 64	n = 14 ; % = 56
Sample size		

69. Lewin, 2015

Bibliographic Reference

Lewin, A.; DeFronzo, R. A.; Patel, S.; Liu, D.; Kaste, R.; Woerle, H. J.; Broedl, U. C.; Initial combination of empagliflozin and linagliptin in subjects with type 2 diabetes; Diabetes Care; 2015; vol. 38 (no. 3); 394-402

Secondary publication of another included study-see primary study for details Other publications associated with this study included in review NcT01422876 registration number Study type Randomised controlled trial (RCT) Study location Study setting Study dates Sources of funding Inclusion criteria This study enrolled subjects aged ≥18 years with type 2 diabetes with BMI ≤45 kg/m2 and HbA1c >7% to ≤10.5% (>53 to ≤91 mmol/mol) at screening despite a diet and exercise regimen who had not received treatment with oral antidiabetic therapy, GLP-1 analogue, or insulin for ≥12 weeks prior to randomisation. Exclusion Criteria Uncontrolled hyperglycaemia (glucose level >240 mg/dL after an overnight fast during the placebo run-in, confirmed by a second measurement); estimated glomerular filtration rate (eGFR), 60 mL/min/1.73 m2 (using the Modification of Diet in Renal Disease equation); acute coronary syndrome, stroke, or transient ischemic attack within 3 months prior to consent; bariatric surgery in the last 2 years; and treatment with anti-obesity drugs within 3 months prior to consent. Patients were recruited from 197 centres in 122 countries. All patients had a 2-week placebo run-in period prior to randomisation.		
publications associated with this study included in review Trial name / registration number Study type Randomised controlled trial (RCT) Study location US Study setting No information available Study dates 08/2011 to 09/2013 Sources of funding Inclusion criteria Inclusion Criteria Exclusion Criteria Exclusion Criteria Exclusion Criteria Fast during the placebo run-in, confirmed by a second measurement); estimated glomerular filtration rate (eGFR), 60 mL/min/1.73 m2 (using the Modification of Diet in Renal Disease equation); acute coronary syndrome, stroke, or transient ischemic attack within 3 months prior to consent; bariatric surgery in the last 2 years; and treatment with anti-obesity drugs within 3 months prior to consent. Patients were recruited from 197 centres in 122 countries. All patients had a 2-week placebo run-in period prior to randomisation.	publication of another included study- see primary study	No information available.
registration number Study type Randomised controlled trial (RCT) Study location US Study setting No information available Sources of funding Boehringer Ingelheim and Eli Lilly and Company. Inclusion criteria This study enrolled subjects aged ≥18 years with type 2 diabetes with BMI ≤45 kg/m2 and HbA1c >7% to ≤10.5% (>53 to ≤91 mmol/mol) at screening despite a diet and exercise regimen who had not received treatment with oral antidiabetic therapy, GLP-1 analogue, or insulin for ≥12 weeks prior to randomisation. Exclusion criteria Uncontrolled hyperglycaemia (glucose level >240 mg/dL after an overnight fast during the placebo run-in, confirmed by a second measurement); estimated glomerular filtration rate (eGFR), 60 mL/min/1.73 m2 (using the Modification of Diet in Renal Disease equation); acute coronary syndrome, stroke, or transient ischemic attack within 3 months prior to consent; bariatric surgery in the last 2 years; and treatment with anti-obesity drugs within 3 months prior to consent. Recruitment / selection of Patients were recruited from 197 centres in 122 countries. All patients had a 2-week placebo run-in period prior to randomisation.	publications associated with this study included in	No information available.
Study setting No information available Study dates 08/2011 to 09/2013 Sources of funding Inclusion Criteria This study enrolled subjects aged ≥18 years with type 2 diabetes with BMI ≤45 kg/m2 and HbA1c >7% to ≤10.5% (>53 to ≤91 mmol/mol) at screening despite a diet and exercise regimen who had not received treatment with oral antidiabetic therapy, GLP-1 analogue, or insulin for ≥12 weeks prior to randomisation. Exclusion Criteria Uncontrolled hyperglycaemia (glucose level >240 mg/dL after an overnight fast during the placebo run-in, confirmed by a second measurement); estimated glomerular filtration rate (eGFR), 60 mL/min/1.73 m2 (using the Modification of Diet in Renal Disease equation); acute coronary syndrome, stroke, or transient ischemic attack within 3 months prior to consent; bariatric surgery in the last 2 years; and treatment with anti-obesity drugs within 3 months prior to consent. Patients were recruited from 197 centres in 122 countries. All patients had a 2-week placebo run-in period prior to randomisation.	registration	NCT01422876
Study setting No information available Study dates 08/2011 to 09/2013 Sources of funding Boehringer Ingelheim and Eli Lilly and Company. Inclusion criteria This study enrolled subjects aged ≥18 years with type 2 diabetes with BMI ≤45 kg/m2 and HbA1c >7% to ≤10.5% (>53 to ≤91 mmol/mol) at screening despite a diet and exercise regimen who had not received treatment with oral antidiabetic therapy, GLP-1 analogue, or insulin for ≥12 weeks prior to randomisation. Exclusion criteria Uncontrolled hyperglycaemia (glucose level >240 mg/dL after an overnight fast during the placebo run-in, confirmed by a second measurement); estimated glomerular filtration rate (eGFR), 60 mL/min/1.73 m2 (using the Modification of Diet in Renal Disease equation); acute coronary syndrome, stroke, or transient ischemic attack within 3 months prior to consent; bariatric surgery in the last 2 years; and treatment with anti-obesity drugs within 3 months prior to consent. Recruitment / selection of Patients were recruited from 197 centres in 122 countries. All patients had a 2-week placebo run-in period prior to randomisation.	Study type	Randomised controlled trial (RCT)
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Sources of funding Inclusion Criteria This study enrolled subjects aged ≥18 years with type 2 diabetes with BMI ≤45 kg/m2 and HbA1c >7% to ≤10.5% (>53 to ≤91 mmol/mol) at screening despite a diet and exercise regimen who had not received treatment with oral antidiabetic therapy, GLP-1 analogue, or insulin for ≥12 weeks prior to randomisation. Exclusion Criteria Uncontrolled hyperglycaemia (glucose level >240 mg/dL after an overnight fast during the placebo run-in, confirmed by a second measurement); estimated glomerular filtration rate (eGFR), 60 mL/min/1.73 m2 (using the Modification of Diet in Renal Disease equation); acute coronary syndrome, stroke, or transient ischemic attack within 3 months prior to consent; bariatric surgery in the last 2 years; and treatment with anti-obesity drugs within 3 months prior to consent. Recruitment / selection of Patients were recruited from 197 centres in 122 countries. All patients had a 2-week placebo run-in period prior to randomisation.	Study setting	No information available
Inclusion criteria This study enrolled subjects aged ≥18 years with type 2 diabetes with BMI ≤45 kg/m2 and HbA1c >7% to ≤10.5% (>53 to ≤91 mmol/mol) at screening despite a diet and exercise regimen who had not received treatment with oral antidiabetic therapy, GLP-1 analogue, or insulin for ≥12 weeks prior to randomisation. Exclusion Criteria Uncontrolled hyperglycaemia (glucose level >240 mg/dL after an overnight fast during the placebo run-in, confirmed by a second measurement); estimated glomerular filtration rate (eGFR), 60 mL/min/1.73 m2 (using the Modification of Diet in Renal Disease equation); acute coronary syndrome, stroke, or transient ischemic attack within 3 months prior to consent; bariatric surgery in the last 2 years; and treatment with anti-obesity drugs within 3 months prior to consent. Recruitment / selection of Patients were recruited from 197 centres in 122 countries. All patients had a 2-week placebo run-in period prior to randomisation.	Study dates	08/2011 to 09/2013
 ≤45 kg/m² and HbA1c >7% to ≤10.5% (>53 to ≤91 mmol/mol) at screening despite a diet and exercise regimen who had not received treatment with oral antidiabetic therapy, GLP-1 analogue, or insulin for ≥12 weeks prior to randomisation. Exclusion Criteria Uncontrolled hyperglycaemia (glucose level >240 mg/dL after an overnight fast during the placebo run-in, confirmed by a second measurement); estimated glomerular filtration rate (eGFR), 60 mL/min/1.73 m² (using the Modification of Diet in Renal Disease equation); acute coronary syndrome, stroke, or transient ischemic attack within 3 months prior to consent; bariatric surgery in the last 2 years; and treatment with anti-obesity drugs within 3 months prior to consent. Recruitment / Patients were recruited from 197 centres in 122 countries. All patients had a 2-week placebo run-in period prior to randomisation. 		Boehringer Ingelheim and Eli Lilly and Company.
fast during the placebo run-in, confirmed by a second measurement); estimated glomerular filtration rate (eGFR), 60 mL/min/1.73 m2 (using the Modification of Diet in Renal Disease equation); acute coronary syndrome, stroke, or transient ischemic attack within 3 months prior to consent; bariatric surgery in the last 2 years; and treatment with anti-obesity drugs within 3 months prior to consent. Recruitment / selection of Patients were recruited from 197 centres in 122 countries. All patients had a 2-week placebo run-in period prior to randomisation.		≤45 kg/m² and HbA1c >7% to ≤10.5% (>53 to ≤91 mmol/mol) at screening despite a diet and exercise regimen who had not received treatment with oral antidiabetic therapy, GLP-1 analogue, or insulin for ≥12 weeks prior to
selection of a 2-week placebo run-in period prior to randomisation.		estimated glomerular filtration rate (eGFR), 60 mL/min/1.73 m2 (using the Modification of Diet in Renal Disease equation); acute coronary syndrome, stroke, or transient ischemic attack within 3 months prior to consent; bariatric surgery in the last 2 years; and treatment with anti-obesity drugs
participants		·

Intervention(s)	Linagliptin 5 mg once daily, taken orally in the morning.
Cointervention	No information available.
Strata 1: People with type 2 diabetes mellitus and heart failure	Not stated/unclear
Strata 2: People with atherosclerotic cardiovascular diseases	Not stated/unclear
Strata 3: People with type 2 diabetes mellitus and chronic kidney disease	Not stated/unclear
Strata 4: People with type 2 diabetes mellitus and high cardiovascular	Not stated/unclear
Subgroup 1: People with frailty	Not stated/unclear
Subgroup 2: Onset of type 2 diabetes mellitus	Mixed population
Subgroup 3: People with non-alcoholic fatty liver disease	Not stated/unclear
Subgroup 4: People with obesity	Not stated/unclear

Subgroup 5: eGFR category at baseline	eGFR ≥30mL/min/1.73m2
Subgroup 6: Albuminuria category at baseline	Mixed population
Sensitivity analysis category: Enrichment trial status	3) Selection of specific population
Population subgroups	No information available.
Comparator	Empagliflozin 10 mg once daily, taken orally in the morning.
	Empagliflozin 5 mg once daily, taken orally in the morning.
	The study included a further two arms listed below, but data for these interventions has not been extracted because they are outside the scope of this review.
	 Empagliflozin 25 mg/ linagliptin 5 mg, taken orally in the morning. Empagliflozin 10 mg/ linagliptin 5 mg, taken orally in the morning.
Number of participants	N = 677
Duration of follow-up	Treatment for 52 weeks and 4 weeks follow-up.
Indirectness	22 countries were included however they have not stated which countries were included.
Method of analysis	ACA
Additional comments	 Efficacy analyses were performed on the full analysis set which included subjects treated with ≥1 dose of study drug who had a baseline and ≥1 treatment HbA1c value. Safety was assessed in the treated set, which comprised subjects treated ≥1 dose of study drug. Rescue medication was to be initiated if a subject had blood glucose >240 mg/dL after an overnight fast between weeks 1 and 12, blood glucose >200 mg/dL after an overnight fast between weeks 12 and 24, or blood glucose >180 mg/dL or HbA1c >8% (>63.9 mmol/mol) after an overnight fast between weeks 24 and 52. The initiation, choice, and dosage of rescue medication were at the discretion of the investigator, according to local prescribing information, but the use of DPP-4 inhibitors, GLP-1 analogues, and SGLT2 inhibitors was not permitted.

69.2.1. Empagliflozin 25 mg once daily (N = 133) Taken orally in the morning

69.2.2. Empagliflozin 10 mg once daily (N = 132) Taken orally in the morning

69.2.3. Linagliptin 5 mg once daily (N = 133) Taken orally in the morning

69.3. Characteristics

69.3.1. Arm-level characteristics

Characteristic	Empagliflozin 25 mg once daily (N = 133)	Empagliflozin 10 mg once daily (N = 132)	Linagliptin 5 mg once daily (N = 133)
% Male	n = 77; % = 57.9	n = 64 ; % = 48.5	n = 75 ; % = 56.4
No of events			
Mean age (SD) (years)	56 (9.3)	53.9 (10.5)	53.8 (11.5)
Mean (SD)			
White	n = 93 ; % = 69.9	n = 99 ; % = 75	n = 103 ; % = 77.4
No of events			
Asian	n = 19; % = 14.3	n = 13 ; % = 9.8	n = 17 ; % = 12.8
No of events			
Other	n = 21; % = 15.8	n = 20 ; % = 15.2	n = 13 ; % = 9.8
No of events			
Comorbidities	NR	NR	NR
Nominal			
Presence of frailty	NR	NR	NR
Nominal			

Characteristic	Empagliflozin 25 mg once daily (N = 133)	Empagliflozin 10 mg once daily (N = 132)	Linagliptin 5 mg once daily (N = 133)
≤1 years	n = 48 ; % = 36.1	n = 43 ; % = 32.6	n = 50 ; % = 37.6
No of events			
>1 to 5 years	n = 48 ; % = 36.1	n = 60 ; % = 45.5	n = 57 ; % = 42.9
No of events			
>5 to 10 years	n = 25 ; % = 18.8	n = 15; % = 11.4	n = 22 ; % = 16.5
No of events			
10 years	n = 12; % = 9	n = 14; % = 10.6	n = 4; % = 3
No of events	7.00 (0.07)		
HbA1c (%)	7.99 (0.97)	8.05 (1.03)	8.05 (0.89)
Mean (SD)	ND		
Heart rate Nominal	NR	NR	NR
	ND		
Smoking status Nominal	NR	NR	NR
	NR		
Alcohol consumption Nominal	INIX	NR	NR
Presence of severe	NR		
mental illness		NR	NR
Nominal			
People with significant cognitive impairment	NR	NR	NR
Nominal			
People with a learning disability	NR	NR	NR
Nominal			
Number of people with obesity	NR	NR	NR
Nominal			
Other antidiabetic medication used	NR	NR	NR

Characteristic	Empagliflozin 25 mg once daily (N = 133)	Empagliflozin 10 mg once daily (N = 132)	Linagliptin 5 mg once daily (N = 133)
Nominal			
Blood pressure- lowering medication used	NR	NR	NR
Other treatment being received	NR	NR	NR
Nominal			

70. Li, 2016

Bibliographic Reference

Li, Feng-Fei; Gao, Gu; Li, Qian; Zhu, Hong-Hong; Su, Xiao-Fei; Wu, Jin-Dan; Ye, Lei; Ma, Jian-Hua; Influence of Dapagliflozin on Glycemic Variations in Patients with Newly Diagnosed Type 2 Diabetes Mellitus.; Journal of diabetes research; 2016; vol. 2016; 5347262

70.1.	oludy details		
Secondary publication of another included study- see primary study for details	Wang, L. Xu, L. Yuan et al., "Sodium-glucose co-transporter-2 inhibitors suppress atrial natriuretic peptide secretion in patients with newly diagnosed Type 2 diabetes," Diabetic Medicine, 2016		
Other publications associated with this study included in review	None.		
Trial name / registration number	Not reported		
Study type	Randomised controlled trial (RCT)		
Study location	China		
Study setting	Hospital		
Study dates	07/2010 to 03/2012		
Sources of funding	Funded by Nanjing Public Health Bureau Project (no. YKK11110), Jiangsu Provincial Department of Science and Technology Project (no. BL2014010), and project funded by China Postdoctoral Science Foundation (2015M581829).		
Inclusion criteria	 Patients with newly diagnosed or drug-naive T2DM Patients receiving 8 weeks of lifestyle management counselling, those who continued to experience inadequate glycemic control, as defined by HbA1c levels of 7.5–10.5%, were recruited. 		
Exclusion criteria	 History of diabetes insipidus Severe uncontrolled hypertension (systolic blood pressure ≥ 180 mmHg and/or diastolic blood pressure ≥ 110 mmHg) and use of any renin-angiotensin system blocker Replacement or chronic systemic corticosteroid treatment History or current diagnosis of significant comorbid diseases, such as cardiovascular, hepatic, and renal diseases 		

	 Positive test for islet cell autoantibodies (such as glutamic acid decarboxylase autoantibodies, islet cell autoantibodies, or insulinoma-like antigen 2), indicating the possibility of type 1 diabetes mellitus.
Recruitment / selection of participants	Subjects were randomised to receive one of the following blinded treatment regimens in a 1:1:1 ratio: dapagliflozin 5 mg, once daily; dapagliflozin 10 mg, once daily; dapagliflozin 5 mg/10 mg matching placebo, once daily (distributed by Bristol-Myers Squibb, Lawrenceville, NJ), for 24 weeks, and, after 4 weeks of treatment, patients lacking glycemic control (fasting blood glucose > 11.1 mmol/L) were eligible to receive another antihyperglycemic drug, such as metformin, based on their particular symptoms. Scheduled visits were at weeks 1 and 24.
Intervention(s)	Dapagliflozin 5 mg or dapagliflozin 10 mg or dapagliflozin 5 mg/10 mg once daily.
Comparator	Placebo
Number of participants	28
Duration of follow-up	24 weeks
Indirectness	
Additional comments	

70.2.1. Dapagliflozin 5 mg or dapagliflozin 10 mg or dapagliflozin 5 mg/10 mg daily (N = 18)

Taken orally

70.2.2. Placebo daily (N = 10)

Taken orally

71. Li, 2019

Bibliographic Reference

Li, J; Zhang, P.; Fan, B; Guo, X; Zheng, Z; The efficacy of saxagliptin in T2DM patients with non-alcoholic fatty liver disease: preliminary data; Rev Assoc Med Bras; 2019; vol. 65 (no. 1); 33-37

	ludy details	
Secondary publication of another included study- see primary study for details	NA	
Other publications associated with this study included in review	None	
Trial name / registration number	Not reported	
Study type	Randomised controlled trial (RCT)	
Study location	Shandong, China	
Study setting	Community	
Study dates	07/204 to 12/2016	
Sources of funding	Not reported	
Inclusion criteria	 Aged ≥30 to ≤60 years Diagnosis of type 2 diabetes (WHO 1999 classification) or Newly diagnosed or diagnosed <2 years of enrolment Naive to hypoglycaemic drug treatment HbA1c level ≥7 to ≤9% Diagnosis of non-alcoholic fatty liver disease (Guidelines for management of non-alcoholic fatty liver disease [2010 revised edition], Chinese Society of Hepatology, Chinese Medical Association) No liver protection treatment 	
Exclusion criteria	 Acute complications and severe chronic complications of diabetes Viral hepatitis, drug hepatitis, auto-immune liver disease, other liver diseases caused by clear damage factors, hepatolenticular degeneration, and total parenteral nutrition 	

	 Liver cirrhosis, severe liver and kidney insufficiency, cardio- cerebrovascular diseases, acute infection, and genetic diseases
Recruitment / selection of participants	Recruited from Department of Endocrinology, Qilu Hospital of Shandong University, Qingdao, China. Randomisation according to computer-generated random number table one of 3 arms. All participants received information on diet and exercise.
Intervention(s)	 Saxagliptin 5 mg once daily Oral saxagliptin tablets 5 mg once daily for 24 weeks.
Strata 1: People with type 2 diabetes mellitus and heart failure	Not stated/unclear
Strata 2: People with atherosclerotic cardiovascular diseases	Not stated/unclear
Strata 3: People with type 2 diabetes mellitus and chronic kidney disease	Not stated/unclear
Strata 4: People with type 2 diabetes mellitus and high cardiovascular risk	Not stated/unclear
Subgroup 1: People with frailty	Not stated/unclear
Subgroup 2: Onset of type 2 diabetes mellitus	Not stated/unclear
Subgroup 3: People with non-alcoholic fatty liver disease	People with non-alcoholic fatty liver disease

Subgroup 4: People with obesity	Not stated/unclear
Subgroup 5: eGFR category at baseline	Not stated/unclear
Subgroup 6: Albuminuria category at baseline	Not stated/unclear
Sensitivity	5) All treatment naïve
analysis category: Enrichment trial status	Inclusion criteria: did not receive hypoglycaemic drug treatment
Comparator	 Glimepiride 2mg once daily Glimepiride 2mg once daily and Polyene phosphatidylcholine 456 mg thrice daily Oral glimepiride tablets 2 mg once daily in both of these groups for 24
	weeks with glimepiride dose adjusted based on blood glucose measurements.
Number of participants	N=64 (two arms only, third arm N=31 excluded from data extraction)
Duration of follow-up	24 weeks
Method of analysis	Other
anaiysis	Not explicitly reported but results reported for all randomised participants who completed trial (that is, excluding participants lost to follow up).

71.2.1. Saxagliptin 5 mg once daily (N = 31)

Oral saxagliptin tablets 5 mg once daily for 24 weeks.

71.2.2. Glimepiride 2 mg once daily (N = 33)

Oral glimepiride tablets 2 mg once daily for 24 weeks.

71.2.3. Glimepiride 2 mg once daily + Polyene phosphatidylcholine 456 mg thrice daily (N = 31)

Oral glimepiride tablets 2 mg once daily + Polyene phosphatidylcholine 456 mg thrice daily for 24 weeks.

71.3. Characteristics

71.3.1. Arm-level characteristics

/ 1.3.1. F	Allii-level Cilala	Otor lotioe	
Characteristic	Saxagliptin 5 mg once daily (N = 31)	Glimepiride 2 mg once daily (N = 33)	Glimepiride 2 mg once daily + Polyene phosphatidylcholine 456 mg thrice daily (N = 31)
% Male	n = 15; % = 48.4	n = 17 ; % = 51.5	n = 16; % = 51.6
Sample size		01.0	
Mean age (SD) (years)	46.6 (8.2)	47.4 (9.4)	49.3 (8.9)
Mean (SD)			
Ethnicity	NR	NR	NR
Nominal	–		
Comorbidities Nominal	NR	NR	NR
	ND		
Presence of frailty Nominal	NR	NR	NR
	11 2 (6 6)		
Time since type 2 diabetes diagnosis (years)	11.3 (6.6)	10.2 (7.6)	9.6 (5.4)
Mean (SD)			
HbA1c (%)	7.79 (0.52)	7.82 (0.61)	7.85 (0.57)
Mean (SD)			
Blood pressure	NR	NR	NR
Nominal			
Heart rate	NR	NR	NR
Nominal			
Smoking status	NR	NR	NR
Nominal			
Alcohol	NR		
consumption		NR	NR
Nominal			
Presence of severe mental illness	NR	NR	NR

NR				
People with significant cognitive impairment Nominal People with a learning disability Nominal Weight NR NR NR NR NR NR NR NR NR N	Characteristic	mg once daily	mg once daily	Polyene phosphatidylcholine
significant cognitive impairment Nominal People with a learning disability Nominal Weight NR NR NR NR NR NR NR NR NR N	Nominal			
People with a learning disabilityNRNRNRNominalNRNRNRWeightNRNRNRNominalSMI (kg/m2)27.2 (4.1)26.5 (3.2)26 (2.9)Mean (SD)NRNRNRNominalNA (NA)NA (NA)NA (NA)Cholesterol and lipid levels (mmol/L)NA (NA)NA (NA)NA (NA)Mean (SD)5.6 (0.8)5.6 (1.2)Trial cholesterol5.4 (0.9)5.6 (0.8)5.6 (1.2)Mean (SD)2.1 (0.8)2.1 (1)Albumin creatinine ratioNRNRNRNominaleGFR (mL/min/1.73m2)NRNRNRNominalOther antidiabetic medication usedNRNRNR	significant cognitive	NR	NR	NR
Iearning disability NR	Nominal			
Weight NR NR NR Nominal NR NR NR BMI (kg/m2) 27.2 (4.1) 26.5 (3.2) 26 (2.9) Mean (SD) NR NR NR Number of people with obesity NR NR NR Nominal NA (NA) NA (NA) NA (NA) Nominal NA (NA) NA (NA) NA (NA) Mean (SD) 5.6 (0.8) 5.6 (1.2) Mean (SD) 2.1 (0.8) 2.1 (1) Mean (SD) NR NR NR Albumin creatinine ratio NR NR NR Nominal R NR NR Nominal NR NR NR Other antidiabetic medication used NR NR NR		NR	NR	NR
NR	Nominal			
BMI (kg/m2) 27.2 (4.1) 26.5 (3.2) 26 (2.9) Mean (SD) NR NR NR Nominal NA (NA) NA (NA) NA (NA) Mean (SD) 5.4 (0.9) 5.6 (0.8) 5.6 (1.2) Mean (SD) 2 (1.1) 2.1 (0.8) 2.1 (1) Mean (SD) NR NR NR Albumin creatinine ratio NR NR NR Nominal R NR NR Nominal Other antidiabetic medication used NR NR NR	Weight	NR	NR	NR
Mean (SD)		07.0 (4.4)		
Number of people with obesity Nominal Cholesterol and lipid levels (mmol/L) Mean (SD) Total cholesterol Mean (SD) Triglycerides 2 (1.1) Albumin creatinine ratio NR NR NR NR NR NR NR NR NR N	, - ,	27.2 (4.1)	26.5 (3.2)	26 (2.9)
Cholesterol and lipid levels (mmol/L) Mean (SD) Total cholesterol Mean (SD) Triglycerides 2 (1.1) Albumin creatinine ratio Nominal eGFR (mL/min/1.73m2) Nominal Other antidiabetic medication used NA (NA) NA (NR	NR	NR
Cholesterol and lipid levels (mmol/L) Mean (SD) Total cholesterol Mean (SD) Triglycerides 2 (1.1) Albumin creatinine ratio Nominal eGFR (mL/min/1.73m2) Nominal Other antidiabetic medication used NA (NA) NA (Nominal			
Total cholesterol 5.4 (0.9) 5.6 (0.8) 5.6 (1.2) Mean (SD) 2 (1.1) 2.1 (0.8) 2.1 (1) Mean (SD) NR NR NR Albumin creatinine ratio NR NR NR Nominal NR NR NR Nominal NR NR NR Nominal NR NR NR	Cholesterol and lipid	NA (NA)	NA (NA)	NA (NA)
Total cholesterol 5.4 (0.9) 5.6 (0.8) 5.6 (1.2) Mean (SD) 2 (1.1) 2.1 (0.8) 2.1 (1) Mean (SD) NR NR NR Albumin creatinine ratio NR NR NR Nominal NR NR NR Nominal NR NR NR Nominal NR NR NR	Mean (SD)			
Section Sect	. ,	5.4 (0.9)		
Triglycerides 2 (1.1) Mean (SD) Albumin creatinine ratio NR NR NR NR NR NR NR NR NR N	M (OD)	, ,	5.6 (0.8)	5.6 (1.2)
Mean (SD) Albumin creatinine ratio NR NR NR NR NR NR NR NR NR N		0 (4.4)		
Albumin creatinine ratio NR NR NR NR NR NR NR NR NR N	riigiycendes	2 (1.1)	2.1 (0.8)	2.1 (1)
NR NR Nominal eGFR (mL/min/1.73m2) NR NR NR NR NR NR NR NR NR NR	Mean (SD)			
eGFR (mL/min/1.73m2) NR NR NR NR NR NR NR NR NR N		NR	NR	NR
(mL/min/1.73m2) NR NR Nominal Other antidiabetic MR medication used NR NR NR	Nominal			
Other antidiabetic NR NR NR		NR	NR	NR
medication used NR NR	Nominal			
Nominal		NR	NR	NR
	Nominal			

Characteristic	Saxagliptin 5 mg once daily (N = 31)	Glimepiride 2 mg once daily (N = 33)	Glimepiride 2 mg once daily + Polyene phosphatidylcholine 456 mg thrice daily (N = 31)
Blood pressure- lowering medication used Nominal	NR	NR	NR
Statins/lipid- lowering medication used Nominal	NR	NR	NR
Other treatment being received Nominal	NR	NR	NR

72. Liu, 2020

Bibliographic Reference

Liu, Lin; Yan, Hongmei; Xia, MingFeng; Zhao, Lin; Lv, Minzhi; Zhao, Naiqin; Rao, Shengxiang; Yao, Xiuzhong; Wu, Weiyun; Pan, Baishen; Bian, Hua; Gao, Xin; Efficacy of exenatide and insulin glargine on nonalcoholic fatty liver disease in patients with type 2 diabetes.; Diabetes/metabolism research and reviews; 2020; vol. 36 (no. 5); e3292

12.1.	luuy uelalis
Secondary publication of another included study- see primary study for details	NA
Other publications associated with this study included in review	None
Trial name / registration number	NCT02303730
Study type	Randomised controlled trial (RCT)
	Open-label trial
Study location	China
Study setting	Community
Study dates	05/2017 to 11/2017
Sources of funding	Funded by: 3SBio Inc; AstraZeneca China; National Key R&D Program of China, Grant/Award Numbers: 2017YFC1309800,2017YFC1309801, 2017YFC1309804.
Inclusion criteria	 Aged ≥18 to ≤70 years Newly diagnosed type 2 diabetes mellitus (WHO Diagnostic criteria 1999) Patients with non-alcoholic fatty disease (NAFLD) with magnetic resonance spectroscopy measurement of liver fat content>10%. HbA1c level ≥7% to ≤10% No heavy drinking history≤5 years (alcohol intake: male<20 g/d; female<10 g/d) HBsAg (-), hepatitis C virus antibody (HCV-Ab) (-) BMI ≥24 kg/m2

Exclusion Pregnancy, lactation, intended pregnancy, or failure to take criteria adequate contraceptive measures (e.g. sterilization, intrauterine device, oral contraceptives, and persistent use of condoms) Diagnosis of type 1 diabetes mellitus, gestational diabetes mellitus or other special types of diabetes Liver and renal dysfunction (ALT or AST) 2.5 times higher than the upper limit of normal, or total bilirubin 1.5 times higher than the upper limit of normal, or $Cr \ge 115 \mu mol/L$) Increased amylase (blood amylase is 2.5 times higher than the upper limit of normal) or presence of gastrointestinal disease Use of drugs that may affect liver fat content≤1 one month before or during the trial period, such as glucocorticoids, thyroid hormone, Use of GLP-1RA, DPP-4 inhibitors, or insulin ≤3 months before enrolment Presence of serious dyslipidaemia or other endocrine diseases (hypothyroidism, hypothalamic-pituitary dysfunction, etc) Fatty liver caused by viral hepatitis, drug, alcohol, Wilson disease or total parenteral nutrition Presence of liver cancer, infection, biliary tract disease or recently increased liver enzyme due to medication Participation in strenuous exercise or administration of any drugs that affect glucose metabolism History of pancreatitis, alcohol abuse, metal disorders or history of allergy to investigational drug Congestive heart failure defined as New York Heart Association (NYHA) class III or IV, unstable angina or myocardial infarction in recent 6 months Any situation that may affect the implementation or results of the study Recruitment / Participants recruited from 4 hospitals in China. After 2 week screening selection of period, participants randomised 1:1 to exenatide or insulin glargine for 24 participants weeks. Central randomisation list used generated by Department of Biostatistics, School of Public Health, Fudan University. Visits at weeks 4, 8, 12, 16, 20 and 24 with telephone calls at weeks 1, 2 and 6. Intervention(s) Exenatide 10 mcg twice daily Participants received 5 mcg twice daily for 4 weeks and then 10 mcg twice daily for 20 weeks. Not stated/unclear Strata 1: People with type 2 Exclusion criteria: New York Heart Association (NYHA) class III or IV. May diabetes be some participants with NYHA class II congestive heart failure. mellitus and heart failure Not stated/unclear Strata 2: People with atherosclerotic Exclusion criteria: Unstable angina or myocardial infarction in recent 6 cardiovascular months. According to this, trial population may include participants with diseases other types of atherosclerotic heart disease.

Strata 3: People with type 2 diabetes mellitus and chronic kidney disease	Not stated/unclear
Strata 4: People with type 2 diabetes mellitus and high cardiovascular risk	Not stated/unclear
Subgroup 1: People with frailty	Not stated/unclear
Subgroup 2: Onset of type 2 diabetes mellitus	Not stated/unclear
Subgroup 3: People with non-alcoholic fatty liver disease	People with non-alcoholic fatty liver disease Inclusion criteria: Participants with non-alcoholic fatty disease with magnetic resonance spectroscopy measurement of liver fat content>10%
Subgroup 4: People with obesity	Not stated/unclear
Subgroup 5: eGFR category at baseline	Not stated/unclear
Subgroup 6: Albuminuria category at baseline	Not stated/unclear
Sensitivity analysis category: Enrichment trial status	5) All treatment naïve All participants had managed diabetes using diet and exercise only.
Population subgroups	
Comparator	 Insulin glargine Participants received subcutaneous insulin glargine once daily for 24 weeks, starting at dose of 0.1 to 0.3 IU/kg (0.1-0.2 IU/kg for HbA1c<8%; 0.2-0.3 IU/kg for HbA1c>8% and then dose up-titrated to achieve FPG<7.0 mmol/L.

Number of participants	N=76
Duration of follow-up	24 weeks
Method of analysis	Modified ITT Full analysis set for all efficacy outcomes included all randomised and treated patients with more than efficacy result. Safety set included all participants who received at least one dose of study drug
Additional comments	

72.2.1. Exenatide 10 mcg twice daily (N = 38)

Subcutaneous injection of exenatide 5 mcg twice daily for 4 weeks then exenatide 10 mcg twice daily for 20 weeks.

72.2.2. Insulin glargine 0.1-0.3IU/kg once daily (N = 38)

Subcutaneous injection of insulin glargine once daily, gradually up-titrated to achieve FPG<7.0 mmol/L.

72.3. Characteristics

72.3.1. Arm-level characteristics

Characteristic	Exenatide 10 mcg twice daily (N = 38)	Insulin glargine 0.1-0.3IU/kg once daily (N = 38)
% Male	n = 19; % = 54.3	n = 19; % = 52.8
Sample size		
Mean age (SD)	47.63 (10.14)	50.56 (11.78)
Mean (SD)		
Ethnicity	NR	NR
Custom value		
Comorbidities	NR	NR
Custom value		
Presence of frailty	NR	NR

Custom value Time since type 2 diabetes diagnosis Mean (SD) HbA1c (%) Blood pressure (mmHg) NA (NA) Mean (SD) Systolic blood pressure 129.6 (14.48) Mean (SD) Diastolic blood pressure NR Mean (SD) Heart rate NR Custom value Smoking status NR Custom value Presence of severe mental illness Custom value People with significant cognitive impairment Custom value People with a learning disability NR O.52 (1.25) 0.52 (1.25)	Characteristic		Insulin glargine 0.1-0.3IU/kg
Time since type 2 diabetes diagnosis 0.32 (0.79) 0.52 (1.25) Mean (SD) 8.32 (0.94) 8.58 (0.91) Mean (SD) NA (NA) NA (NA) Mean (SD) NA (NA) NA (NA) Systolic blood pressure 129.6 (14.48) 127.97 (11.3) Mean (SD) 78.64 (8.57) 78.64 (8.67) Mean (SD) NR NR Custom value NR NR Smoking status NR NR Custom value NR NR Alcohol consumption NR NR Custom value NR NR Presence of severe mental illness NR NR Custom value NR NR People with significant cognitive impairment NR NR Custom value NR NR	Custom value	daily (N = 38)	once daily (N = 38)
Mean (SD)		0.22 (0.70)	
HbA1c (%) 8.32 (0.94) 8.58 (0.91) Mean (SD) NA (NA) NA (NA) Mean (SD) 129.6 (14.48) 127.97 (11.3) Mean (SD) 80.46 (8.57) 78.64 (8.67) Mean (SD) NR NR Mear rate NR NR Custom value NR NR People with significant cognitive impairment NR NR Custom value NR NR		0.32 (0.79)	0.52 (1.25)
Mean (SD) Blood pressure (mmHg) NA (NA) NA (NA) NA (NA) Mean (SD) Systolic blood pressure 129.6 (14.48) 127.97 (11.3) Mean (SD) Diastolic blood pressure 80.46 (8.57) 78.64 (8.67) Mean (SD) Heart rate NR Custom value Smoking status NR Custom value Alcohol consumption NR Custom value Presence of severe mental illness Custom value People with significant cognitive impairment Custom value People with a learning NR	Mean (SD)		
Blood pressure (mmHg) Mean (SD) Systolic blood pressure 129.6 (14.48) 127.97 (11.3) Mean (SD) Diastolic blood pressure 80.46 (8.57) Mean (SD) Heart rate NR Custom value Smoking status NR Custom value Alcohol consumption NR Custom value Presence of severe mental illness Custom value People with significant cognitive impairment Custom value People with a learning NR NR NR NR NR NR NR NR NR N	, ,	8.32 (0.94)	8.58 (0.91)
Mean (SD) Systolic blood pressure 129.6 (14.48) 127.97 (11.3) Mean (SD) Diastolic blood pressure 80.46 (8.57) 78.64 (8.67) Mean (SD) Heart rate NR Custom value Smoking status NR Custom value Alcohol consumption NR Custom value Presence of severe mental illness Custom value People with significant cognitive impairment Custom value People with a learning NR NR NR NR NR NR NR	· · ·		
Systolic blood pressure Mean (SD) Diastolic blood pressure 80.46 (8.57) Mean (SD) Heart rate NR Custom value Smoking status NR Custom value Alcohol consumption NR Custom value Presence of severe mental illness Custom value People with significant cognitive impairment Custom value People with a learning NR NR 127.97 (11.3) NR NR NR NR NR NR NR NR NR N	Blood pressure (mmHg)	NA (NA)	NA (NA)
Mean (SD) Diastolic blood pressure 80.46 (8.57) Mean (SD) Heart rate NR Custom value Smoking status NR Custom value Alcohol consumption NR Custom value Presence of severe mental illness Custom value People with significant cognitive impairment Custom value People with a learning NR NR NR NR NR NR NR	Mean (SD)		
Diastolic blood pressure 80.46 (8.57) Mean (SD) Heart rate NR Custom value Smoking status NR Custom value Alcohol consumption NR Custom value Presence of severe mental illness Custom value People with significant cognitive impairment Custom value People with a learning NR NR	·	129.6 (14.48)	127.97 (11.3)
Mean (SD) Heart rate NR Custom value Smoking status NR Custom value Alcohol consumption NR Custom value Presence of severe mental illness Custom value People with significant cognitive impairment Custom value People with a learning NR NR NR NR NR NR NR NR	• •	00.46 (0.57)	
Custom value Smoking status NR Custom value Alcohol consumption NR Custom value Presence of severe mental illness Custom value People with significant cognitive impairment Custom value People with a learning NR NR NR NR NR NR NR	•	60.40 (6.57)	78.64 (8.67)
Smoking status Custom value Alcohol consumption NR Custom value Presence of severe mental illness Custom value People with significant cognitive impairment Custom value People with a learning NR NR NR NR NR	` '	NR	NR
Custom value Alcohol consumption NR Custom value Presence of severe mental illness Custom value People with significant cognitive impairment Custom value People with a learning NR NR NR NR NR NR	Custom value		
Alcohol consumption Custom value Presence of severe mental illness Custom value People with significant cognitive impairment Custom value People with a learning NR NR NR NR NR	-	NR	NR
Custom value Presence of severe mental illness Custom value People with significant cognitive impairment Custom value People with a learning NR NR NR NR			
Presence of severe mental illness Custom value People with significant cognitive impairment Custom value People with a learning NR NR NR	•	NR	NR
illness Custom value People with significant cognitive impairment Custom value People with a learning NR NR NR NR		NR	
People with significant cognitive impairment Custom value People with a learning NR	illness	TVIX	NR
Custom value People with a learning NR NR			
People with a learning NR		NR	NR
	Custom value		
		NR	NR
Custom value	Custom value		
Weight (kg) 79.28 (9.64) 77.63 (13.7)	Weight (kg)	79.28 (9.64)	77.63 (13.7)
Mean (SD)	Mean (SD)		
BMI (kg/m2) 28.49 (3.02) 27.84 (3.1)	BMI (kg/m2)	28.49 (3.02)	27.84 (3.1)
Mean (SD)	Mean (SD)		

	-	L
Characteristic	Exenatide 10 mcg twice daily (N = 38)	Insulin glargine 0.1-0.3IU/kg once daily (N = 38)
Number of people with obesity	NR	NR
Custom value		
Cholesterol and lipid levels (mmol/L)	NA (NA)	NA (NA)
Mean (SD)		
Total cholesterol	4.96 (0.94)	4.92 (0.89)
Mean (SD)		,
HDL-cholesterol	1.11 (0.19)	1.08 (0.22)
Mean (SD)		1.00 (0.22)
LDL-cholesterol	3.01 (0.76)	0.04 (0.70)
Mean (SD)		2.81 (0.72)
Triglyceride	2.01 (1.23)	
	2.0 ((2 0)	2.41 (1.59)
Mean (SD)		
Albumin creatinine ratio	NR	NR
Custom value		
eGFR (mL/min/1.73m2)	NR	NR
Custom value		
Other antidiabetic medication	NR	NR
used		INIX
Custom value		
Blood pressure-lowering medication used	NR	NR
Custom value		
Statins/lipid-lowering medication used	NR	NR
Custom value		
Other treatment being received	NR	NR
Custom value		1111
Data for group baseline charact	enistics are for the following	ing purpler of porticipants.

Data for group baseline characteristics are for the following number of participants: Exenatide, n=35; Insulin glargine, n=36.

73. Mari, 2008

Bibliographic Reference

Mari, A.; Scherbaum, W. A.; Nilsson, P. M.; Lalanne, G.; Schweizer, A.; Dunning, B. E.; Jauffret, S.; Foley, J. E.; Characterization of the influence of vildagliptin on model-assessed -cell function in patients with type 2 diabetes and mild hyperglycaemia; J Clin Endocrinol Metab; 2008; vol. 93 (no. 1); 103-9

13.1.	luuy uelalis	
Secondary publication of another included study- see primary study for details	NA	
Other publications associated with this study included in review	 Additional outcomes reported in: Scherbaum, W. A., Schweizer, A., Mari, A., Nilsson, P. M., Lalanne, G., Jauffret, S., & Foley, J. E. (2008). Efficacy and tolerability of vildagliptin in drug-naive patients with type 2 diabetes and mild hyperglycaemia. Diabetes, Obesity and Metabolism, 10(8), 675-682. 	
Trial name / registration number	NCT00101712	
Study type	Randomised controlled trial (RCT)	
Study location	International (69 sites in Finland, France, Germany, Romania, Spain and Sweden)	
Study setting	Community	
Study dates	10/2004 to 05/2006	
Sources of funding	Funded by Novartis Pharmaceuticals Corporation	
Inclusion criteria	 Aged ≥18 years Diagnosis of type 2 diabetes≥8 weeks before enrolment Naive to treatment with oral antidiabetic drug (defined as no such drug≥12 weeks before screening and no such drug≥3 consecutive months at any time HbA1c level ≥6.2 to ≤7.5% at screening (≤7.0% for participating centres in Finland and Spain) BMI ≥22 to ≤45 kg/m2 If female then non-fertile or of childbearing potential using medically-approved birth control method 	

Exclusion criteria	 History of type 1 or secondary form of diabetes Acute metabolic diabetic complications ≤6 months of enrolment Evidence of significant diabetic complications History of significant cardiac arrhythmia, congestive heart failure or New York Heart Association class III or IV Liver disease (e.g. cirrhosis or chronic active hepatitis Any significant laboratory abnormality
Recruitment / selection of participants	Recruited from 69 sites in 6 countries with 2 week screening period, 52 week treatment period followed by 4 week treatment-free washout period. All participants received individualized lifestyle counselling regarding weight loss, diet and physical activity.
Intervention(s)	Vildagliptin 50 mg once daily
	Oral vildagliptin tablets 50 mg once daily, 30 min before breakfast, for 52 weeks.
Strata 1:	Not stated/unclear
People with type 2 diabetes mellitus and heart failure	Exclusion criteria: NYHA class 3 and 4. Trial may include participants with NYHA class 2.
Strata 2: People with atherosclerotic cardiovascular diseases	Not stated/unclear
Strata 3: People with type 2 diabetes mellitus and chronic kidney disease	Not stated/unclear
Strata 4: People with type 2 diabetes mellitus and high cardiovascular risk	Not stated/unclear
Subgroup 1: People with frailty	Not stated/unclear
Subgroup 2: Onset of type 2 diabetes mellitus	Not stated/unclear

Subgroup 3: People with non-alcoholic fatty liver disease	Not stated/unclear
Subgroup 4: People with	Mixed population
obesity	46.2% of vildagliptin group and 46.7% of placebo group were obese (BMI>=30 kg/m2)
Subgroup 5: eGFR category at baseline	Not stated/unclear
Subgroup 6: Albuminuria category at baseline	Not stated/unclear
Sensitivity analysis	7) Mixed population
category: Enrichment trial status	Inclusion criteria: participants who had not taken oral antidiabetic drug in previous 12 weeks and had not taken any such drug for than 3 consecutive months at any time.
Comparator	• Placebo
	Placebo for vildagliptin for 52 weeks
Number of participants	N= 306
Duration of follow-up	52 weeks + 4 week washout period
Method of analysis	Not stated/unclear
anaryoro	Population used for analysis not reported; also reports results for completer population/

73.2.1. Vildagliptin 50 mg once daily (N = 156)

Oral vildagliptin tablets 50 mg once daily for 52 weeks.

73.2.2. Placebo (N = 150)

Placebo to vildagliptin for 52 weeks.

73.3. Characteristics

73.3.1. Arm-level characteristics

75.5.1. Allii-level Cilala		
Characteristic	Vildagliptin 50 mg once daily (N = 156)	Placebo (N = 150)
% Male Sample size	n = 93; % = 59.6	n = 89 ; % = 59.3
Mean age (SD)	63.3 (10.2)	62.8 (11)
Mean (SD)		
Ethnicity	n = NA ; % = NA	n = NA ; % = NA
Sample size		INA
Caucasian Sample size	n = 155 ; % = 99.4	n = 149 ; % = 99.3
Other	n = 1; % = 0.6	
Sample size	11 - 1 , 70 - 0.0	n = 1; % = 0.7
Comorbidities	NR	
		NR
Nominal	ND	
Presence of frailty	NR	NR
Nominal		
Time since type 2 diabetes diagnosis (years)	2.5 (2.9)	2.7 (3.2)
Mean (SD)		
HbA1c (%) Significant difference between groups, p=0.0403	6.7 (0.4)	6.8 (0.4)
Mean (SD)		
Blood pressure	NR	ND
Nominal		NR
Heart rate	NR	NR
Nominal		
Smoking status Nominal	NR	NR
	ND	
Alcohol consumption	NR	NR

Characteristic	Vildagliptin 50 mg once daily (N = 156)	Placebo (N = 150)
Nominal		
Presence of severe mental illness	NR	NR
Nominal		
People with significant cognitive impairment	NR	NR
Nominal		
People with a learning disability	NR	NR
Nominal		
Weight	NR	NR
Nominal		
BMI (kg/m2) Mean (SD)	30.4 (4.9)	30 (4.9)
Number of people with obesity	NR	
Nominal		NR
Cholesterol and lipid levels	NR	
•		NR
Nominal		
Albumin creatinine ratio	NR	NR
Nominal		
eGFR (mL/min/1.73m2)	NR	NR
Nominal		
Other antidiabetic medication used	NR	NR
Nominal		
Blood pressure-lowering medication used	NR	NR
Nominal		
Statins/lipid-lowering medication used	NR	NR
Nominal		
Other treatment being received	NR	NR
Nominal		

74. Miyagawa, 2015

Bibliographic Reference

Miyagawa, J.; Odawara, M.; Takamura, T.; Iwamoto, N.; Takita, Y.; Imaoka, T.; Once-weekly glucagon-like peptide-1 receptor agonist dulaglutide is non-inferior to once-daily liraglutide and superior to placebo in Japanese patients with type 2 diabetes: a 26-week randomized phase III study; Diabetes Obes Metab; 2015; vol. 17 (no. 10); 974-83

14.1. 3	luuy uelalis	
Secondary publication of another included study- see primary study for details	No	
Other publications associated with this study included in review	 Odawara, M., Miyagawa, J., Iwamoto, N., Takita, Y., Imaoka, T., & Takamura, T. (2016). Once-weekly glucagon-like peptide-1 receptor agonist dulaglutide significantly decreases glycated haemoglobin compared with once-daily liraglutide in Japanese patients with type 2 diabetes: 52 weeks of treatment in a randomized phase III study. <i>Diabetes, Obesity and Metabolism</i>, 18(3), 249-257. 26-week health-related quality of life outcomes reported in: Suzuki, S., Oura, T., Takeuchi, M., & Boye, K. S. (2017). Evaluation of the impact of once weekly dulaglutide on patient-reported outcomes in Japanese patients with type 2 diabetes: comparisons with liraglutide, insulin glargine, and placebo in two randomized studies. <i>Health and Quality of Life Outcomes</i>, 15(1), 1-10. 	
Trial name / registration number	NCT01558271	
Study type	Randomised controlled trial (RCT)	
Study location	Japan (33 sites in 14 cities)	
Study setting	Community	
Study dates	04/2012 to 10/2013	
Sources of funding	Eli Lilly Japan K.K., Kobe, Japan.	
Inclusion criteria	Diagnosis of type 2 diabetes before screening	

- Management with diet and exercise only or treatment with oral antidiabetic monotherapy (except for thiazolidinedione) and willing to discontinue medication with 8 week washout period before randomisation
 HbA1c level 7.0-10.0% inclusive at screening and randomisation for those on diet and exercise only; or HbA1c level 6.5-9.0% inclusive at screening and HbA1c level 7.0-10.0% inclusive at
- randomisation for those on oral antidiabetic monotherapy

 BMI ≥18.5 to ≥35.0 kg/m2

Exclusion criteria

- Diagnosis of type 1 diabetes.
- Previous treatment with any other GLP-1 analogue.
- Receiving more than half of maximum dose of sulfonylureas at screening
- Currently taking insulin or thiazolidinediones (TZD), or previous insulin or TZD treatment ≤3 months screening.
- Obvious clinical signs or symptoms of pancreatitis, history of chronic pancreatitis, or acute pancreatitis at screening, as determined by investigator
- Serum amylase concentration ≥3 times upper limit of reference range and/or a serum lipase concentration ≥2 times upper limit of reference range, as determined by central laboratory at screening
- History (personal or family) of medullary C-cell hyperplasia, focal hyperplasia, or medullary thyroid carcinoma

Recruitment / selection of participants

Participants recruited from 33 sites in Japan. Initial 2 week screening period, then 2 week lead-in period for treatment-naive participants and 8 week wash out period for participants on monotherapy. Eligible participants randomized to treatment in 4:2:1 ratio (dulaglutide; liraglutide; placebo) using computer-generated random sequence with interactive voice response system and stratified by pre-study oral antidiabetic medication status (yes/no), BMI group (<25; ≥25 kg/m2), and HbA1c (≤8.5; >8.5%). Participants and investigators masked to assignment to dulaglutide and placebo treatment but not masked to assignment to liraglutide treatment. At end of 26 weeks, participants in placebo group switched to dulaglutide 0.75 once weekly for remaining 26 weeks. Participants not tolerating study drugs discontinued them but remained in study to collect safety data.

Intervention(s)

Dulaglutide 0.75 mg once weekly

Subcutaneous injection of dulaglutide 0.75 mg once weekly for 26 weeks, provided in non-identifiable solution in prefilled syringe and initiated at full dose. Subsequent 26 weeks was open-label dulaglutide.

Strata 1: People with type 2 diabetes mellitus and heart failure

Not stated/unclear

Strata 2: People with atherosclerotic cardiovascular diseases	Not stated/unclear
Strata 3: People with type 2 diabetes mellitus and chronic kidney disease	Not stated/unclear
Strata 4: People with type 2 diabetes mellitus and high cardiovascular risk	Not stated/unclear
Subgroup 1: People with frailty	Not stated/unclear
Subgroup 2: Onset of type 2 diabetes mellitus	Not stated/unclear
Subgroup 3: People with non-alcoholic fatty liver disease	Not stated/unclear
Subgroup 4: People with obesity	Not stated/unclear
Subgroup 5: eGFR category at baseline	Not stated/unclear
Subgroup 6: Albuminuria category at baseline	Not stated/unclear
Sensitivity	7) Mixed population
analysis category:	Participants were treatment-naive (diabetes managed with diet and
Enrichment trial status	exercise only) or if on oral antidiabetic monotherapy had 8-week washout period
Comparator	Liraglutide 0.9 mg once weeklyPlacebo

	Open label subcutaneous liraglutide up-titrated from 0.3 mg/day during week 1 to 0.6 mg/day during week 2 and 0.9 mg/day at start of week 3 for remaining 49 weeks. Participants in placebo arm received placebo for 26 weeks then were switched over to dulaglutide 0.75 mg once weekly for remaining 26 weeks.
Number of participants	N=492
Duration of follow-up	26 weeks + 52 weeks Participants in dulaglutide and liraglutide arms received treatment for 52 weeks. Participants in placebo arm switched to dulaglutide 0.75 mg once daily at 26 weeks for 26 weeks.
Method of analysis	Modified ITT mITT analysis for efficacy analysis - all randomised participants who took at least one dose of study drug. Other Safety analysis conducted on as-treated population according to actual treatments received.
Additional comments	

74.2.1. Dulaglutide 0.75 mg once weekly (N = 281)

Subcutaneous injection of dulaglutide 0.75 mg once weekly for 52 weeks.

74.2.2. Liraglutide 0.9 mg once daily (N = 141)

Subcutaneous injection of liraglutide 0.9 mg once daily for 52 weeks.

74.2.3. Placebo once weekly for 26 weeks then dulalgutide 0.75 mg once weekly for 26 weeks (N = 79)

Placebo injection once weekly using non-identifiable prefilled syringe (same as used for dulaglutide) for 26 weeks, then switched to dulaglutide 0.75 mg once weekly for 26 weeks.

74.3.1. Arm-level characteristics

74.3.1. AIII	i-level Characteri	51103	
Characteristic	Dulaglutide 0.75 mg once weekly (N = 281)		Placebo once weekly for 26 weeks then dulaglutide 0.75 mg once weekly for 26 weeks (N = 79)
% Male Sample size	n = 228 ; % = 81	n = 113 ; % = 83	n = 55 ; % = 79
	57.0 (0.0)		
Mean age (SD) Mean (SD)	57.2 (9.6)	57.9 (10.4)	57.7 (8.3)
Ethnicity	NR		
		NR	NR
Nominal			
Comorbidities	NR	ND	ND
		NR	NR
Nominal			
Presence of frailty	NR	NR	NR
Nominal			
Time since type 2	6.8 (5.6)		
diabetes diagnosis (years)	0.0 (0.0)	6.3 (6)	6.3 (5.1)
Mean (SD)			
HbA1c (%)	8.15 (0.77)	8.08 (0.89)	8.2 (0.83)
Mean (SD)	NB		
Blood pressure	NR	NR	NR
Nominal			
Heart rate	NR		
		NR	NR
Nominal			
Smoking status	NR	NR	NR
Nominal			
Alcohol consumption	NR	NR	NR
Nominal			
Presence of severe mental illness	NR	NR	NR

Characteristic	Dulaglutide 0.75 mg once weekly (N = 281)	_	Placebo once weekly for 26 weeks then dulaglutide 0.75 mg once weekly for 26 weeks (N = 79)
Nominal			
People with significant cognitive impairment	NR	NR	NR
Nominal			
People with a learning disability	NR	NR	NR
Nominal			
Weight (kg)	71.3 (12.5)	70.2 (12.5)	69.3 (11.6)
Mean (SD)	25.6 (2.6)		
BMI (kg/m2) Mean (SD)	25.6 (3.6)	25.5 (3.5)	25.2 (3.2)
, ,	ND		
Number of people with obesity	NR	NR	NR
Nominal			
Cholesterol and lipid levels	NR	NR	NR
Nominal			
Albumin creatinine ratio Nominal	NR	NR	NR
	ND		
eGFR (mL/min/1.73m2) Nominal	NR	NR	NR
Other antidiabetic medication used Sample size	n = NA ; % = NA	n = NA ; % = NA	n = NA ; % = NA
Naive to oral	n = 186 ; % = 66		
antihyperglycaemic medication Sample size		n = 89 ; % = 65	n = 48 ; % = 69
·	$p = 0.4 \cdot 0.4 = 2.4$		
Pre-study oral antihyperglycaemic medication	n = 94 ; % = 34	n = 48; % = 35	n = 22 ; % = 31

Characteristic	Dulaglutide 0.75 mg once weekly (N = 281)	_	Placebo once weekly for 26 weeks then dulaglutide 0.75 mg once weekly for 26 weeks (N = 79)
Sample size			
Blood pressure- lowering medication used	NR	NR	NR
Nominal			
Statins/lipid-lowering medication used	NR	NR	NR
Nominal			
Other treatment being received	NR	NR	NR
Nominal			

All baseline characteristics data are for the following number of participants: Dulaglutide, n=280; Liraglutide, n=137; Placebo, n=70.

75. Miyazaki, 2002

Bibliographic Reference

Miyazaki, Y.; Matsuda, M.; DeFronzo, R. A.; Dose-response effect of pioglitazone on insulin sensitivity and insulin secretion in type 2 diabetes; Diabetes Care; 2002; vol. 25 (no. 3); 517-23

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Secondary publication of another included study- see primary study for details	
Other publications associated with this study included in review	
Trial name / registration number	NR
Study type	Randomised controlled trial (RCT)
Study location	NR
Study setting	Multicentre
Study dates	NR
Sources of funding	Support for the multicentre trial was provided by Takeda
Inclusion criteria	 - HbA1c >= 7% - fasting plasma glucose>= 140mg/dl - fasting c-peptide >1 ng/ml
Exclusion criteria	-Use of insulin -Unstable proliferative retinopathy - Impaired liver function (aspartate aminotransferase or alanine aminotransferase >2.5 * upper limit of normal

- Impaired kidney function (serum creatinine >1.8 mg/dl)
- Anemia
Patients taking previous antidiabetic therapy (sulfonylureas or metformin) underwent a 6-
to 8-week single-blind washout period
before the baseline OGTT was performed.
After the washout period, only patients
with HbA1c 7.0% were enrolled. Patients were randomized to one of five parallel treatment groups: pioglitazone 7.5,
15, 30, or 45 mg/day or placebo. During
the double-blind period, patients were
seen every 2 weeks for the first 6 weeks
and every 4 weeks for the remaining 20
weeks. At 26 weeks, all subjects underwent repeat 75-g OGTT. To minimize the
confounding effect of weight loss on metabolic changes, no specific dietary modifications were recommended during the
study
Pioglitazone 7.5/15/30/45 mg/day
Not stated/unclear
Not stated/unclear
Not stated/unclear

Strata 4: People with type 2 diabetes mellitus and high cardiovascular risk	Not stated/unclear
Subgroup 1: People with frailty	Not stated/unclear
Subgroup 2: Onset of type 2 diabetes mellitus	Not stated/unclear
Subgroup 3: People with non-alcoholic fatty liver disease	Not stated/unclear
Subgroup 4: People with obesity	Not stated/unclear
Subgroup 5: eGFR category at baseline	Not stated/unclear
Subgroup 6: Albuminuria category at baseline	Not stated/unclear
Sensitivity analysis category: Enrichment trial status	3) Selection of specific populationOnly people with >7.0% enrolled, excluded responders.
Number of participants	54
Duration of follow-up	26 weeks
Indirectness	Direct
Method of analysis	Not stated/unclear

75.2.1. Pioglitazone 7.5mg/day (N = 13)

- **75.2.2.** Pioglitazone 15mg/day (N = 12)
- **75.2.3.** Pioglitazone 30mg/day (N = 11)
- **75.2.4.** Pioglitazone 45mg/day (N = 11)
- 75.2.5. Placebo (N = 11)

75.3.1. Study-level characteristics

Characteristic	Study (N =)
% Male	n = 34; % = 58.6
Sample size	
Mean age (SD)	54 (1)
Mean (SD)	
ВМІ	31.5 (0.6)
Mean (SD)	

75.3.2. Arm-level characteristics

Characteristic	Pioglitazone 7.5mg/day (N = 13)	Pioglitazone 15mg/day (N = 12)	Pioglitazone 30mg/day (N = 11)	Pioglitazone 45mg/day (N = 11)	Placebo (N = 11)
Age	51 (3)	57 (4)	51 (2)	51 (2)	58 (3)
Mean (SD)					
Caucasian	10	10	6	7	9
Nominal					
African- American	1	1	0	1	1
Nominal					

Characteristic	Pioglitazone 7.5mg/day (N = 13)		Pioglitazone 30mg/day (N = 11)	Pioglitazone 45mg/day (N = 11)	Placebo (N = 11)
Mexican- American	0	1	4	2	1
Nominal					
Asian	2	0	0	0	0
Nominal					
Others	0	0	1	1	0
Nominal					
Male Sample size	n = 10; % = 76.9	n = 8; % = 66.6	n = 8 ; % = 72.7	n = 5 ; % = 45.5	n = 3; % = 27.2

76. Moretto, 2008

Bibliographic Reference

Moretto, T. J.; Milton, D. R.; Ridge, T. D.; Macconell, L. A.; Okerson, T.; Wolka, A. M.; Brodows, R. G.; Efficacy and tolerability of exenatide monotherapy over 24 weeks in antidiabetic drug-naive patients with type 2 diabetes: a randomized, double-blind, placebo-controlled, parallel-group study; Clin Ther; 2008; vol. 30 (no. 8); 1448-60

70.1. 3	tudy details		
Secondary publication of another included study- see primary study for details	NA		
Other publications associated with this study included in review	None		
Trial name / registration number	NCT00381342		
Study type	Randomised controlled trial (RCT)		
Study location	International (23 centres in India, Puerto Rico, Romania, Russia, USA)		
Study setting	Community		
Study dates	09/2006 to 09/2007		
Sources of funding	Funded by Amylin Pharmaceuticals, Inc, San Diego, CA, USA and Eli Lilly and Co. Indiana, IN, USA. Publication funded by Eli Lilly and Co.		
Inclusion criteria	 Aged ≥18 years If female, either using contraceptives for >12 weeks before screening and continuing throughout study, or postmenopausal or surgically sterile Diagnosis of type 2 diabetes BMI 25-45 kg/m2 inclusive Managing their type 2 diabetes with diet and exercise consistent with the local standards of medical care in the opinion of the investigator HbA1c 6.5-10.0% inclusive at screening 		
Exclusion criteria	 Previous treatment with antidiabetic agent Blood pressure ≥160/≥110 mmHg 		

	 History or presence of clinically significant cardiac disease ≤1 year before participation in study History of renal transplant or active renal or hepatic disease Previous weight loss medication ≤12 weeks prior to screening
Recruitment / selection of participants	At screening, each patient received next consecutive patient number from block of patient numbers assigned to investigator. One week after screening, a 2-week, single-blind lead-in period in which participants received subcutaneous placebo injections twice per day (b.i.d.) to acclimatize them to self-administering injections. Participants were then randomly assigned 1:1:1 ratio, using computer-generated randomization sequence generated by interactive voice-response system to one of the 3 double-blind treatment arms, stratified by screening HbA1c (≤8%, >8%) within each site. Participants discontinued trial due to if (i) they had increase of 1% HbA1c level from baseline at any study visit, (ii) HbA1c>10.5% at or after week 12, or (iii) had ≥4 fasting serum glucose concentrations >260 mg/dL for 7 consecutive days according to self-monitored blood glucose tests.
Intervention(s)	 Exenatide 10 mcg twice daily Exenatide 5 mcg twice daily Participants instructed to maintain individualised diet and exercise regimen during trial and self-administered exenatide in upper arm, thought or abdomen twice daily (15 min before both breakfast and dinner). Participants in exenatide 10 mcg arm received 5 mcg twice daily for 4 weeks then 10 mcg twice daily for 20 weeks. Participants in exenatide 5 mcg arm received 5 mcg twice daily for 24 weeks.
Strata 1: People with type 2 diabetes mellitus and heart failure	Not stated/unclear
Strata 2: People with atherosclerotic cardiovascular diseases	People without other cardiovascular diseases Exclusion criteria: history or presence of clinically significant cardiac disease within 1 year prior to study
Strata 3: People with type 2 diabetes mellitus and chronic kidney disease	People without chronic kidney disease Exclusion criteria: history of renal transplant or active renal or hepatic disease
Strata 4: People with type 2 diabetes mellitus and high	Not stated/unclear

cardiovascular risk	
Subgroup 1: People with frailty	Not stated/unclear
Subgroup 2: Onset of type 2 diabetes mellitus	Not stated/unclear
Subgroup 3: People with non-alcoholic fatty liver disease	Not stated/unclear
Subgroup 4: People with obesity	Not stated/unclear
Subgroup 5: eGFR category at baseline	Not stated/unclear
Subgroup 6: Albuminuria category at baseline	Not stated/unclear
Sensitivity analysis category: Enrichment trial status	5) All treatment naïve All participants managed using only diet and exercise (exclusion criteria: previous treatment with anti-diabetic agent).
Population subgroups	
Comparator	 Placebo Participants in this arm received volume equivalent to 5 mcg exenatide twice daily for 4 weeks then volume equivalent to exenatide 5 mcg or 10 mcg twice daily for 20 weeks.
Number of participants	N=233
Duration of follow-up	24 weeks
Method of analysis	Modified ITT mTT - all randomised participants who received ≥1 dose of study drug - for all outcomes. Primary endpoint was HbA1c at week 24 or LOCF.

76.2.1. Exenatide 10 mcg twice daily (N = 78)

Subcutaneous injection of exenatide 5mcg twice per day (15 min before morning and evening meals) for 4 weeks then 10 mcg twice per day for 20 weeks.

76.2.2. Exenatide 5 mcg twice daily (N = 77)

Subcutaneous injection of exenatide 5mcg twice per day (15 min before morning and evening meals) for 24 weeks.

76.2.3. Placebo (N = 78)

Volume-matched placebo injections equivalent to exenatide 5 mcg twice daily for 4 weeks, then volume equivalent to exenatide 5 mcg or 10 mcg twice daily for 20 weeks.

76.3.1. Arm-level characteristics

Characteristic	Exenatide 10 mcg twice daily (N = 78)	Exenatide 5 mcg twice daily (N = 77)	Placebo (N = 78)
% Male	n = 40; % = 52	n = 48 ; % = 62	n = 42 ; % = 55
Sample size			00
Mean age (SD) (years)	55 (10)	54 (10)	53 (9)
Mean (SD)			
Asian	n = 18; % = 23	n = 22 ; % = 29	n = 21 ; % =
Sample size			27
Black	n = 3; % = 4	n = 0; % = 0	n = 3 ; % =
Sample size			4
Hispanic	n = 1; % = 1	n = 5; % = 6	n = 2 ; % =
Sample size			3
White	n = 56 ; % = 72	n = 50 ; % = 65	n = 51 ; % =
Sample size			66
Comorbidities	NR	NR	NR
Custom value			

Characteristic	Exenatide 10 mcg twice daily (N = 78)	Exenatide 5 mcg twice daily (N = 77)	Placebo (N = 78)
Presence of frailty	NR	NR	NR
Custom value			
Time since type 2 diabetes diagnosis (years)	2 (3)	2 (3)	1 (2)
Mean (SD)			
HbA1c (%)	7.8 (1)	7.9 (1)	7.8 (0.9)
Mean (SD)			- ()
Systolic blood pressure	130 (12)	129 (11)	129 (12)
Mean (SD)	70 (0)		
Diastolic blood pressure	79 (8)	78 (7)	78 (7)
Mean (SD)			
Heart rate	NR	NR	NR
Custom value			
Smoking status	NR	NR	NR
Custom value			
Alcohol consumption	NR	NR	NR
Custom value			
Presence of severe mental illness	NR	NR	NR
Custom value			
People with significant cognitive impairment	NR	NR	NR
Custom value			
People with a learning disability	NR	NR	NR
Custom value			
Weight (kg)	86 (16)	85 (15)	86 (16)
Mean (SD)	0.4.45)		
BMI (kg/m2)	31 (5)	32 (5)	32 (5)
Mean (SD)			

Characteristic	Exenatide 10 mcg twice daily (N = 78)	Exenatide 5 mcg twice daily (N = 77)	Placebo (N = 78)
Number of people with obesity	NR	NR	NR
Custom value			
Cholesterol and lipid levels	NR	NR	NR
Custom value			
Albumin creatinine ratio	NR	NR	NR
Custom value			
eGFR (mL/min/1.73m2)	NR	NR	NR
Custom value			
Other antidiabetic medication used	None	None	None
Custom value			
Blood pressure-lowering medication used	NR	NR	NR
Custom value			
Statins/lipid-lowering medication used	NR	NR	NR
Custom value			
Other treatment being received	NR	NR	NR
Custom value			

77. Mu, 2017

Bibliographic Reference

Mu, Y.; Pan, C.; Fan, B.; Hehnke, U.; Zhang, X.; Zhang, X.; Wang, X.; Liu, J.; Zhang, Y.; Du, J.; Ma, J.; Gong, Y.; Efficacy and safety of linagliptin/metformin single-pill combination as initial therapy in drug-naive Asian patients with type 2 diabetes; Diabetes Res Clin Pract; 2017; vol. 124; 48-56

11.1.	luuy uelans
Secondary publication of another included study- see primary study for details	No additional information.
Other publications associated with this study included in review	No additional information.
Trial name / registration number	NCT01708902.
Study type	Randomised controlled trial (RCT)
Study location	Multicenter. China, Malaysia, Philippines and Vietnam.
Study setting	Outpatient follow-up.
Study dates	October 9th 2012 to April 8th 2014.
Sources of funding	Supported by Boehringer Ingelheim and Eli Lilly and Company.
Inclusion criteria	Age 18-80; diagnosis of T2DM; never received any antidiabetes drugs (or no more than 30 cumulative days of antidiabetes therapy 12 weeks prior to randomisation and no antidiabetes therapy within these 12 weeks); HbA1c 7.5%-11.0% at screening for the main group and >11.0% at screening for the additional parallel group.
Exclusion criteria	History of acute coronary syndrome, stroke or transient ischaemic attacks within 3 months of randomisation; impaired hepatic function (serum levels of alanine aminotransferase, aspartate aminotransferase or alkaline phosphatase >3x upper limit of normal or bilirubin >1.5 x upper limit of normal); renal failure or renal impairment (eGFR <60mL/min); known history of pancreatitis.
Recruitment / selection of participants	No additional information.

Intervention(s)	Linagliptin and metformin (study A) N=294
	Two groups. Linagliptin 2.5mg once a day and metformin 500mg twice a day (n=147) and linagliptin 2.5mg once a day and metformin 1000mg twice a day (n=147).
	Concomitant therapy: People whose glycaemia was not adequately controlled received hypoglycaemic rescue therapy to ensure their safety. The use of rescue therapy (with sulfonylureas, thiazolidinediones or insulin), was permitted only during the randomised treatment period and was administered only if the person had hyperglycaemia on fasting plasma testing confirmed by a second glucose determination performed on a different day. If this could not be controlled then the person was to be discontinued from the trial.
Strata 1: People with type 2 diabetes mellitus and heart failure	People without heart failure
Strata 2: People with atherosclerotic cardiovascular diseases	People without other cardiovascular diseases
Strata 3: People with type 2 diabetes mellitus and chronic kidney disease	People without chronic kidney disease
Strata 4: People with type 2 diabetes mellitus and high cardiovascular risk	Not stated/unclear
Subgroup 1: People with frailty	Not stated/unclear
Subgroup 2: Onset of type 2 diabetes mellitus	Not stated/unclear
Subgroup 3: People with	People without non-alcoholic fatty liver disease

non-alcoholic fatty liver disease	
Subgroup 4: People with obesity	Not stated/unclear
Subgroup 5: eGFR category at baseline	Not stated/unclear
Subgroup 6: Albuminuria category at baseline	Not stated/unclear
Sensitivity analysis category: Enrichment trial status	7) Mixed population
Population subgroups	No additional information.
Comparator	Metformin (study A) N=289
	Two groups. Metformin 500mg twice a day (n=145) and metformin 1000mg twice a day (n=144).
	Concomitant therapy: People whose glycaemia was not adequately controlled received hypoglycaemic rescue therapy to ensure their safety. The use of rescue therapy (with sulfonylureas, thiazolidinediones or insulin), was permitted only during the randomised treatment period and was administered only if the person had hyperglycaemia on fasting plasma testing confirmed by a second glucose determination performed on a different day. If this could not be controlled then the person was to be discontinued from the trial.
	Linagliptin (study A) N=147
	Linagliptin 5mg once a day.
	Concomitant therapy: People whose glycaemia was not adequately controlled received hypoglycaemic rescue therapy to ensure their safety. The use of rescue therapy (with sulfonylureas, thiazolidinediones or insulin), was permitted only during the randomised treatment period and was administered only if the person had hyperglycaemia on fasting plasma testing confirmed by a second glucose determination performed on a different day. If this could not be controlled then the person was to be discontinued from the trial.
	testing confirmed by a second glucose determination performed on a different day. If this could not be controlled then the person was to be

Number of participants	730 in study 1 (the main trial), 163 in study 2 (the side trial) - only the main trial is included as the side trial was only followed up for 12 weeks for the continuous outcomes.
Duration of follow-up	26 weeks.
Indirectness	No additional information.
Method of analysis	Other
	Full case analysis with last observation carried forward
Additional comments	No additional information.

77.2.1. Linagliptin + metformin (study A) (N = 294)

Two groups. Linagliptin 2.5mg once a day and metformin 500mg twice a day (n=147) and linagliptin 2.5mg once a day and metformin 1000mg twice a day (n=147). Concomitant therapy: People whose glycaemia was not adequately controlled received hypoglycaemic rescue therapy to ensure their safety. The use of rescue therapy (with sulfonylureas, thiazolidinediones or insulin), was permitted only during the randomised treatment period and was administered only if the person had hyperglycaemia on fasting plasma testing confirmed by a second glucose determination performed on a different day. If this could not be controlled then the person was to be discontinued from the trial.

77.2.2. Metformin (study A) (N = 289)

Two groups. Metformin 500mg twice a day (n=145) and metformin 1000mg twice a day (n=144). Concomitant therapy: People whose glycaemia was not adequately controlled received hypoglycaemic rescue therapy to ensure their safety. The use of rescue therapy (with sulfonylureas, thiazolidinediones or insulin), was permitted only during the randomised treatment period and was administered only if the person had hyperglycaemia on fasting plasma testing confirmed by a second glucose determination performed on a different day. If this could not be controlled then the person was to be discontinued from the trial.

77.2.3. Linagliptin (study A) (N = 147)

Linagliptin 5mg once a day. Concomitant therapy: People whose glycaemia was not adequately controlled received hypoglycaemic rescue therapy to ensure their safety. The use of rescue therapy (with sulfonylureas, thiazolidinediones or insulin), was permitted only during the randomised treatment period and was administered only if the person had hyperglycaemia on fasting plasma testing confirmed by a second glucose determination performed on a different day. If this could not be controlled then the person was to be discontinued from the trial.

77.3.1. Arm-level characteristics

77.3.1. Arm-lev	el characteristics		
Characteristic	Linagliptin + metformin (study A) (N = 294)	Metformin (study A) (N = 289)	Linagliptin (study A) (N = 147)
% Male	n = 179 ; % = 61	n = 182 ; % = 63	n = 76 ; % = 52
Sample size			
Mean age (SD) (years)	51.1 (9.8)	51.8 (10)	50.8 (10.5)
Mean (SD)			
Ethnicity	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
Sample size			
Comorbidities	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
Sample size			
Presence of frailty	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
Sample size			
Time since type 2 diabetes diagnosis	n = NA ; % = NA	n = NA ; % = NA	n = NA ; % = NA
Sample size			
Less than and equal to 1 year	n = 236 ; % = 80	n = 223 ; % = 77	n = 110 ; % = 78
Sample size			
Greater than 1 to less than and equal to 5 years	n = 35 ; % = 12	n = 37 ; % = 13	n = 23 ; % = 16
Sample size			
Greater than 5 and less than and equal to 10 years	n = 11; % = 4	n = 14 ; % = 5	n = 7; % = 5
Sample size			
Greater than 10 years	n = 1; % = 0.03	n = 3 ; % = 1	n = 1; % = 1
Sample size			
HbA1c (%)	8.7 (1)	8.7 (1.1)	8.7 (0.9)
Mean (SD)			
Blood pressure	NR (NR)	NR (NR)	NR (NR)
Mean (SD)			

Characteristic	Linagliptin + metformin (study A) (N = 294)	Metformin (study A) (N = 289)	Linagliptin (study A) (N = 147)
Heart rate	NR (NR)	NR (NR)	NR (NR)
Mean (SD)			
Smoking status	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
Sample size			
Alcohol consumption	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
Sample size			
Presence of severe mental illness	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
Sample size	ND - 0/ ND		
People with significant cognitive impairment	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
Sample size	ND 0/ ND		
People with a learning disability Sample size	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
	70.7 (12)		
Weight (kg) Mean (SD)	70.7 (12)	70.1 (11.5)	70.2 (13.5)
BMI (kg/m2)	26 (3.65)		
Mean (SD)	20 (0.00)	26 (3.3)	26.2 (3.9)
Number of people with obesity	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
Sample size			
Cholesterol and lipid levels Mean (SD)	NR (NR)	NR (NR)	NR (NR)
Albumin creatinine ratio	NR (NR)		
Mean (SD)	INIX (INIX)	NR (NR)	NR (NR)
eGFR (mL/min/1.73m2)	NR (NR)	NR (NR)	NR (NR)
Mean (SD)			(,
Other antidiabetic medication used	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
Sample size			

Characteristic	Linagliptin + metformin (study A) (N = 294)	Metformin (study A) (N = 289)	Linagliptin (study A) (N = 147)
Blood pressure-lowering medication used Sample size	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
Statins/lipid-lowering	n = NR ; % = NR		
medication used		n = NR ; % = NR	n = NR ; % = NR
Sample size			
Other treatment being received	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
Sample size			

78. Nauck, 2016

Bibliographic Reference

Nauck, M. A.; Di Domenico, M.; Patel, S.; Kobe, M.; Toorawa, R.; Woerle, H. J.; Linagliptin and pioglitazone combination therapy versus monotherapy with linagliptin or pioglitazone: A randomised, double-blind, parallel-group, multinational clinical trial; Diabetes Vasc Dis Res; 2016; vol. 13 (no. 4); 286-298

70.1.	tudy details					
Secondary publication of another included study- see primary study for details	NA					
Other publications associated with this study included in review	None					
Trial name / registration number	NCT01183013					
Study type	Randomised controlled trial (RCT)					
Study location	International (132 sites in Estonia, Germany, Latvia, Spain, UK and USA)					
Study setting	Community					
Study dates	08/2010 to 02/2013					
Sources of funding	Funded by Boehringer Ingelheim and Eli Lilly and Co.					
Inclusion criteria	 Aged ≥18 to ≤80 years at screening Diagnosis of type 2 diabetes insufficient glycaemic control on diet and exercise alone No use of oral antidiabetic drug≤10 weeks of screening BMI≤45 kg/m2 at screening 					
Exclusion criteria	 Pregnancy, nursing baby, or of child-bearing potential and not practicing effective birth control/not planning to practice such control for duration of study Uncontrolled hyperglycaemia with confirmed glucose level > 240 mg/dl (> 13.3 mmol/l) after overnight fast during screening or placebo run-in period Myocardial infarction ≤6 months, stroke or TIA ≤ 3 months prior to informed consent 					

- Clinical evidence of active liver disease (e.g. jaundice) or the ALT level > 2.5 times the upper limit of normal (according to pioglitazone label)
- Bariatric surgery, performed ≤2 years prior to informed consent or planned at the time of informed consent
- Gastrointestinal surgeries prior to informed consent that induce chronic malabsorption
- Known hypersensitivity or allergy to the investigational products (linagliptin and/or pioglitazone) or their excipients (including matching placebos)
- Any contraindication or restriction for the use of pioglitazone according to local prescribing information, including diagnosis of heart failure or history of heart failure, and haemodialysis patients
- Treatment with rosiglitazone, pioglitazone, GLP-1 analogues, or insulin ≤3 months prior to informed consent
- Treatment with anti-obesity drugs (e.g. sibutramine, orlistat) ≤3 months prior to informed consent
- Alcohol or drug abuse≤3 months prior to informed consent or history of alcoholism
- Current treatment with systemic corticosteroids at time of informed consent or change in dosage of thyroid hormones within 6 weeks prior to informed consent
- History of pancreatitis or bladder cancer

Recruitment / selection of participants

Participants recruited from 132 sites across the world for 2 part trial (part A and part B). Initial washout period of >10 weeks for participants on oral antidiabetic monotherapy, followed by 2 week placebo run-in period for all participants. Part A was 30 week treatment period with randomisation using equal allocation to one of 7 treatment groups: Fixed-dose combination linagliptin 5 mg/pioglitazone 15, 30 or 45 mg once daily; linagliptin 5 mg once daily; pioglitazone 15, 30 or 45 mg once daily. Randomisation by computer-generated random sequence using interactive voice response system or web response system. Treatments masked using double-blind and double-dummy design. Part B was a planned extension period up to 54 weeks with 5 treatment groups: participants in the 2 groups who received pioglitazone 15 mg in part A (as monotherapy or fixed-dose combination) and who continued to part B were to receive pioglitazone 30 mg (either as monotherapy or fixed-dose combination, according to assignment in part A). Participants in 5 other part A treatment groups were to remain same in part B.

However, due to USFDA and EMA safety concerns about pioglitazone (increased risk of bladder cancer) trial was attenuated so that all participants ended trial participation at 30 weeks and no new participants were entered into part B. Participants already in part B concluded participation by completing final visit procedures.

Intervention(s)

- Linagliptin 5 mg/Pioglitazone 15 mg SPC once daily
- Linagliptin 5 mg/Pioglitazone 30 mg SPC once daily
- Linagliptin 5 mg/Pioglitazone 45 mg SPC once daily

All participants in these groups received treatment drugs for 30 weeks (part A). Participants in Pioglitazone 45 mg single-pill combination group initially received pioglitazone 30 mg, up-titrated to 45 mg over 6-week

	forced titration period. Participants in all groups received daily placebo pills.
Cointervention	Placebo. Reports double-dummy design but no further details reported so assume that all arms received two pills (one active treatment, one placebo).
Strata 1: People with	People without heart failure
type 2 diabetes mellitus and heart failure	Exclusion criteria: Any contraindication or restriction for the use of pioglitazone according to local prescribing information, including diagnosis of heart failure or history of heart failure, and haemodialysis patients
Strata 2: People with	People without other cardiovascular diseases
atherosclerotic	Exclusion criteria: Myocardial infarction \leq 6 months, stroke or TIA \leq 3 months prior to informed consent
Strata 3: People with type 2 diabetes mellitus and chronic kidney disease	Not stated/unclear
Strata 4: People with type 2 diabetes mellitus and high cardiovascular risk	Not stated/unclear
Subgroup 1: People with frailty	Not stated/unclear
Subgroup 2: Onset of type 2 diabetes mellitus	Not stated/unclear
Subgroup 3: People with non-alcoholic fatty liver disease	Not stated/unclear
Subgroup 4: People with obesity	Not stated/unclear
Subgroup 5: eGFR category at baseline	eGFR ≥30mL/min/1.73m2

	All but one participant in trial (assigned to Linagliptin 5 mg/Pioglitazone 30 mg FDC group) had eGFR≥30mL/min/1.73 m2. This participant had eGFR of 27.85 mL/min.					
Subgroup 6: Albuminuria category at baseline	Not stated/unclear					
Sensitivity analysis	7) Mixed population					
category: Enrichment trial status	nclusion criteria: No use of oral antidiabetic drug≤10 weeks of screening					
Population subgroups						
Comparator	 Pioglitazone 15 mg once daily Pioglitazone 30 mg once daily Pioglitazone 45 mg once daily Linagliptin 5 mg once daily All participants in these groups received treatment drugs for 30 weeks (part A). Participants randomised to pioglitazone 45 mg group initially received pioglitazone 30 mg, up-titrated to 45 mg during 6 week forced titration period. Participants in all groups received daily placebo pills.					
Number of participants	N=936					
Duration of follow-up	30 weeks					
Indirectness	No concerns					
Method of analysis	ACA					
	Secondary efficacy analysis of change in HbA1c based on observed cases with those with missing data excluded.					
	Modified ITT					
	Primary efficacy analysis of mean change from baseline in HbA1c conducted on all randomised participants who received at least one dose of study drug and had baseline and at least one on-treatment HbA1c measurement (Full analysis set). Safety analysis conducted on all randomised participants who received at least one dose of study drug.					

78.2.1. Pioglitazone 45 mg + Linagliptin 5 mg SPC once daily (N = 133)

Pioglitazone 45 mg/Linagliptin 5 mg single-pill combination (SPC) oral tablets once daily for 6 weeks, followed by linagliptin 5 mg low dose/pioglitazone 45 mg fixed-dose combination tablets once daily for up to 87 weeks.

78.2.2. Pioglitazone 30 mg + Linagliptin 5 mg SPC once daily (N = 133)

Linagliptin 5 mg low dose/Pioglitazone 30 mg single-pill combination (SPC) oral tablets once daily for up to 84 weeks.

78.2.3. Pioglitazone 15 mg + Linagliptin 5 mg SPC once daily (N = 126)

Linagliptin 5 mg/Pioglitazone 15 mg single-pill combination (SPC) oral tablets once daily for 30 weeks, followed by linagliptin 5 mg low dose/pioglitazone 30 mg fixed dose combination once daily for up to 54 weeks.

78.2.4. Linagliptin 5 mg once daily (N = 135)

Linagliptin 5mg low dose oral tablets once daily for 30 weeks followed by linagliptin 5mg low dose/Pioglitazone 30 mg fixed dose combination oral tablets once daily for up to 54 weeks.

78.2.5. Pioglitazone 45 mg once daily (N = 138)

Pioglitazone 30 mg oral capsules once daily for 6 weeks followed by pioglitazone 45 mg once daily for up to 78 weeks.

78.2.6. Pioglitazone 30 mg once daily (N = 140)

Pioglitazone 30 mg oral capsules once daily for up to 84 weeks.

78.2.7. Pioglitazone 15 mg once daily (N = 131)

Pioglitazone 15 mg oral capsules once daily for 30 weeks followed by pioglitazone 30 mg once daily for up to 54 weeks.

78.3.1. Arm-level characteristics

78.3.1.	Am	i-ievei cha	aracteristi	US			
Characteristi c	one 45 mg + Linaglipti n 5 mg SPC once	one 30 mg +	one 15 mg + Linaglipti n 5 mg SPC once	tin 5 mg once	one 45	one 30 mg once	one 15 mg once
% Male Sample size	n = 73; % = 54.9	n = 68; % = 51.1	n = 71; % = 56.3	n = 83 ; % = 61.5	n = 72; % = 52.2	n = 73; % = 52.1	n = 73 ; % = 55.7
Mean age (SD) (years) Mean (SD)	59.8 (10.2)	56.7 (10.1)	57.1 (10)	56 (10.4)	56.5 (11)	57 (11.5)	56.3 (10.4)
Black/Africa n American Sample size	n = 16; % = 12	n = 11; % = 8.3	•	n = 20 ; % = 14.8	n = 17; % = 12.3	n = 17; % = 12.1	n = 20; % = 15.3
Other American Indian/Alaska Native, Asian or Hawaiian/Pac ific Islander	n = 4; % = 3.1	n = 3; % = 2.3	n = 4; % = 3.2	n = 4; % = 3	n = 2; % = 1.4	n = 3; % = 2.1	n = 3; % = 2.3
Sample size White Sample size	n = 113 ; % = 85	n = 119 ; % = 89.5		n = 111 ; % = 82.2	n = 119 ; % = 86.2	n = 120 ; % = 85.7	
Comorbiditie s Custom value	NR	NR	NR	NR	NR	NR	NR
Presence of frailty Custom value	NR	NR	NR	NR	NR	NR	NR
Time since type 2 diabetes diagnosis	n = NA ; % = NA				n = NA ; % = NA		n = NA ; % = NA

Characteristi c	one 45 mg + Linaglipti n 5 mg SPC once	Pioglitaz one 30 mg + Linaglipti n 5 mg SPC once daily (N = 133)	n 5 mg SPC once	tin 5 mg once	one 45 mg once	Pioglitaz one 30 mg once daily (N = 140)	one 15 mg once
(years) Full analysis set: LINA/PIOG 45, n=126; LINA/PIOG 30, n=125; LINA/PIOG 5, n=120; LINA 5, n=130; PIOG 45, n=134; PIOG 30, n=134; PIOG 15 ,n=124							
Sample size							
< or equal to 1 year	n = 37; % = 29.4	n = 35; % = 28	n = 32; % = 26.7	% =	n = 39; % = 29.1	n = 38; % = 28.4	n = 35; % = 28.2
Sample size				31.5			
>1 to < or equal to 5 years Sample size	n = 42; % = 33.3	n = 48; % = 38.4	n = 42; % = 35	n = 48; % = 36.9	n = 63; % = 47	n = 55; % = 41	n = 53; % = 42.7
>5 years to < or equal to 10 years	n = 26; % = 20.6		n = 33; % = 27.5			n = 30; % = 22.4	n = 26; % = 21
Sample size							
10 years Sample size	n = 21; % = 16.7	n = 10; % = 8	n = 13; % = 10.8	n = 16; % = 12.3		n = 11; % = 8.2	n = 10; % = 8.1
HbA1c (%) Full analysis set: LINA/PIOG 45, n=126; LINA/PIOG 30, n=125;	8 (0.8)	8.2 (1.1)	8.1 (0.9)	9 (0.9)	8.1 (0.9)	9 (0.9)	8.3 (0.9)

	one 45 mg + Linaglipti n 5 mg SPC once	Pioglitaz one 30 mg + Linaglipti n 5 mg SPC once daily (N = 133)	Pioglitaz one 15 mg + Linaglipti n 5 mg SPC once daily (N = 126)	tin 5 mg once	one 45 mg once	Pioglitaz one 30 mg once daily (N = 140)	Pioglitaz one 15 mg once daily (N = 131)
LINA/PIOG 5, n=120; LINA 5, n=130; PIOG 45, n=134; PIOG 30, n=134; PIOG 15 ,n=124 Mean (SD)							
	NR						
pressure		NR	NR	NR	NR	NR	NR
Custom value							
	NR						
		NR	NR	NR	NR	NR	NR
Custom value							
status	NR	NR	NR	NR	NR	NR	NR
Custom value							
Alcohol consumptio n	NR	NR	NR	NR	NR	NR	NR
Custom value							
Presence of severe mental illness	NR	NR	NR	NR	NR	NR	NR
Custom value							
People with significant cognitive impairment	NR	NR	NR	NR	NR	NR	NR
Custom value							

Characteristi c	one 45 mg + Linaglipti n 5 mg SPC once	Pioglitaz one 30 mg + Linaglipti n 5 mg SPC once daily (N = 133)	n 5 mg SPC once	tin 5 mg once	mg once	Pioglitaz one 30 mg once daily (N = 140)	Pioglitaz one 15 mg once daily (N = 131)
People with a learning disability Custom value	NR	NR	NR	NR	NR	NR	NR
	04.0						
Weight (kg) Mean (SD)	91.8 (18.5)	91.8 (19.3)	93.6 (19)	94.7 (19.3)	96.9 (19)	91.5 (19.9)	91.5 (20.7)
BMI (kg/m2)	32.4 (5.4)	32.6 (5.7)	32.4 (5.5)	32.7	33.9 (5.5)	32.2 (5.3)	32.3 (5.6)
Mean (SD)				(5.3)			
Number of people with obesity	NR	NR	NR	NR	NR	NR	NR
Custom value	ND						
Cholesterol and lipid levels	NR	NR	NR	NR	NR	NR	NR
Custom value							
Albumin creatinine ratio	NR	NR	NR	NR	NR	NR	NR
Custom value							
(mL/min/1.73 m2)	n = NA ; % = NA	,	n = NA ; % = NA	,	•	,	n = NA ; % = NA
Sample size							
Normal renal function (> or equal to 90)	n = 35; % = 26.3	n = 43; % = 32.3		n = 48; % = 35.6	n = 55; % = 39.9	n = 52 ; % = 37.1	n = 52; % = 39.7
Sample size							

Characteristi c	one 45 mg + Linaglipti n 5 mg SPC once	Pioglitaz one 30 mg + Linaglipti n 5 mg SPC once daily (N = 133)	n 5 mg SPC once	tin 5 mg once	mg once	Pioglitaz one 30 mg once daily (N = 140)	Pioglitaz one 15 mg once daily (N = 131)
Mild impairment (60 to <90)	n = 82; % = 61.7	n = 79 ; % = 59.4	n = 69; % = 54.8	n = 78 ; % = 57.8	n = 64; % = 46.4	n = 67; % = 47.9	n = 67; % = 51.1
Sample size							
Moderate impairment (30 to <60) Sample size	n = 16; % = 12				n = 19; % = 13.8	n = 20; % = 14.3	n = 12; % = 9.2
Severe impairment or end-stage renal disease (<30)	n = 0; % = 0	n = 1; % = 0.8	n = 0; % = 0	n = 0; % = 0	n = 0; % = 0	n = 0; % = 0	n = 0; % = 0
Sample size							
Other antidiabetic medication used Full analysis set: LINA/PIOG 45, n=126; LINA/PIOG 30, n=125; LINA/PIOG 5, n=120; LINA 5, n=130; PIOG 45, n=134; PIOG 30, n=134; PIOG 15, n=124 Sample size	n = NA ; % = NA		n = NA ; % = NA		n = NA ; % = NA	n = NA ; % = NA	n = NA ; % = NA
No prior antidiabetic drug used	n = 82; % = 65.1	n = 82; % = 65.6	n = 84; % = 70	n = 82 ; % = 63.1	n = 92 ; % = 68.7	n = 82 ; % = 61.2	n = 86; % = 69.4

Characteristi c	one 45 mg + Linaglipti n 5 mg SPC once	Pioglitaz one 30 mg + Linaglipti n 5 mg SPC once daily (N = 133)	n 5 mg SPC once	tin 5 mg once	one 45 mg once	Pioglitaz one 30 mg once daily (N = 140)	one 15 mg once
Sample size							
One prior antidiabetic drug used Sample size	n = 42; % = 34.1	n = 38; % = 30.4	n = 35; % = 29.2	n = 47; % = 36.2	n = 37; % = 27.6	n = 48; % = 35.8	n = 35; % = 28.2
Two prior	n = 1; %						
antidiabetic drugs used	= 0.8	n = 5; % = 4	n = 1; % = 0.8	n = 1; % = 0.8	•	n = 4; % = 3	n = 3; % = 2.4
Sample size							
Blood pressure- lowering medication used	NR	NR	NR	NR	NR	NR	NR
Statins/lipid- lowering medication used	NR	NR	NR	NR	NR	NR	NR
Other	NR						
treatment being received	INIX	NR	NR	NR	NR	NR	NR
Custom value							

79. Odawara, 2016

Bibliographic Reference

Odawara, M; Miyagawa, J; Iwamoto, N; Takita, Y; Imaoka, T; Takamura, T; Once-weekly glucagon-like peptide-1 receptor agonist dulaglutide significantly decreases glycated haemoglobin compared with once-daily liraglutide in Japanese patients with type 2 diabetes: 52 weeks of treatment in a randomized phase III study.; Diabetes, obesity & metabolism; 2016; vol. 18 (no. 3); 249-57

Secondary publication of another included study- see primary study for details Miyagawa, J., Odawara, M., Takamura, T., Iwamoto, N., Takita, Y., & Imaoka, T. (2015). Once-weekly glucagon-like peptide-1 receptor agonist dulaglutide is non-inferior to once-daily liraglutide and superior to placebo in Japanese patients with type 2 diabetes: a 26-week randomized phase III study. Diabetes, Obesity and Metabolism, 17(10), 974-983. Health-related quality of life outcomes reported in: Suzuki, S., Oura, T., Takeuchi, M., & Boye, K. S. (2017). Evaluation of the impact of once weekly dulagilutide on patient-reported outcomes in juncluded in paparese patients with type 2 diabetes: comparisons with liraglutide, insulin glargine, and placebo in two randomized studies. Health and Quality of Life Outcomes, 15(1), 1-10. NCT01558271 Study type Randomised controlled trial (RCT) Study dates Sources of funding Inclusion criteria Diagnosis of type 2 diabetes before screening Diagnosis of type 2 diabetes on proving treatment with oral antidiabetic monotherapy Diagnosis of type 1 diabetes. Diagnosis of type 1 diabetes. Previous treatment with any other GLP-1 analogue.	79.1. 3	tudy details						
mobilication of another included study-see primary study for details Other publications associated with this study included in review Trial name / registration number Study type Study dates Study dates Study dates Study dates Suruces of funding Inclusion criteria Miyagawa, J., Odawara, M., Takamura, T., Iwamoto, N., Takita, Y., & Imaoka, T. (2015). Once-weekly glucagon-like peptide-1 receptor agonist dulaglutide in non-inferior to once-daily liraglutide and superior to placebo in Japanese patients with type 2 diabetes: a 26-week randomized phase III study. Diabetes, Obesity and Metabolism, 17(10), 974-983. Health-related quality of life outcomes reported in: Suzuki, S., Oura, T., Takeuchi, M., & Boye, K. S. (2017). Evaluation of the impact of once weekly dulaglutide on patient-reported outcomes in sulin glargine, and placebo in two randomized studies. Health and Quality of Life Outcomes, 15(1), 1-10. NCT01558271 Trial name / registration number Study type Randomised controlled trial (RCT) Study dates Od/2012 to 10/2013 Eli Lilly Japan K.K., Kobe, Japan. Diagnosis of type 2 diabetes before screening Diagnosis of type 2 diabetes before screening Management with diet and exercise only or treatment with oral antidiabetic monotherapy (except for thiazolidinedione) and willing to discontinue medication with 8 week washout period before randomisation HbA1c level 7.0-10.0% inclusive at screening and randomisation for those on diet and exercise only; or HbA1c level 6.5-9.0% inclusive at screening and HbA1c level 7.0-10.0% inclusive at randomisation for those on oral antidiabetic monotherapy BMI ≥ 18.5 to ≥ 35.0 kg/m2 Exclusion	Secondary	Yes, see primary study (Miyagawa 2015) for details of trial:						
publications associated with this study included in review included in review insulin glargine, and placebo in two randomized studies. Health and Quality of Life Outcomes, 15(1), 1-10. Trial name / registration number Study type Randomised controlled trial (RCT) Study location Study setting Study dates O4/2012 to 10/2013 Eli Lilly Japan K.K., Kobe, Japan. Diagnosis of type 2 diabetes before screening • Management with diet and exercise only or treatment with oral antidiabetic monotherapy (except for thiazolidinedione) and willing to discontinue medication with 8 week washout period before randomisation for those on diet and exercise only; or HbA1c level 6.5-9.0% inclusive at screening and HbA1c level 7.0-10.0% inclusive at randomisation for those on oral antidiabetic monotherapy • BMI ≥18.5 to ≥35.0 kg/m2 Exclusion Suzuki, S., Oura, T., Takeuchi, M., & Boye, K. S. (2017). Evaluation of the impact of the with lingation patients. Will all patients—subject of the patie	publication of another included study- see primary study	naoka, T. (2015). Once-weekly glucagon-like peptide-1 receptor agonist ulaglutide is non-inferior to once-daily liraglutide and superior to placebo Japanese patients with type 2 diabetes: a 26-week randomized phase III						
Suzuki, S., Oura, T., Takeuchi, M., & Boye, K. S. (2017). Evaluation of the impact of once weekly dulaglutide on patient-reported outcomes in Japanese patients with type 2 diabetes: comparisons with liraglutide, insulin glargine, and placebo in two randomized studies. Health and Quality of Life Outcomes, 15(1), 1-10. Trial name / registration number Study type Randomised controlled trial (RCT) Study location Japan (33 sites in 14 cities) Study setting Community Study dates Sources of funding Inclusion criteria Diagnosis of type 2 diabetes before screening • Diagnosis of type 2 diabetes before screening • Management with diet and exercise only or treatment with oral antidiabetic monotherapy (except for thiazolidinedione) and willing to discontinue medication with 8 week washout period before randomisation • HbA1c level 7.0-10.0% inclusive at screening and randomisation for those on diet and exercise only; or HbA1c level 6.5-9.0% inclusive at screening and HbA1c level 7.0-10.0% inclusive at randomisation for those on oral antidiabetic monotherapy • BMI ≥18.5 to ≥35.0 kg/m2 Exclusion Suzuki, S., Oura, T., Takeuchi, M., & Boye, K. S. (2017). Evaluation of the impact of once weekly dulaglutide on patient-reported outcomes in Japanese patients with tipe 2 diabetes: comparisons with liraglutide, insulint patients. Health and Call patients. Hea		Health-related quality of life outcomes reported in:						
Tegistration number Study type Randomised controlled trial (RCT) Study location Japan (33 sites in 14 cities) Study setting Community Study dates 04/2012 to 10/2013 Sources of funding Inclusion criteria • Diagnosis of type 2 diabetes before screening • Management with diet and exercise only or treatment with oral antidiabetic monotherapy (except for thiazolidinedione) and willing to discontinue medication with 8 week washout period before randomisation • HbA1c level 7.0-10.0% inclusive at screening and randomisation for those on diet and exercise only; or HbA1c level 6.5-9.0% inclusive at screening and HbA1c level 7.0-10.0% inclusive at randomisation for those on oral antidiabetic monotherapy • BMI ≥18.5 to ≥35.0 kg/m2 Exclusion • Diagnosis of type 1 diabetes.	associated with this study included in	impact of once weekly dulaglutide on patient-reported outcomes in Japanese patients with type 2 diabetes: comparisons with liraglutide, insulin glargine, and placebo in two randomized studies. <i>Health and</i>						
Study location Japan (33 sites in 14 cities) Study setting Community Sources of funding Eli Lilly Japan K.K., Kobe, Japan. Inclusion criteria • Diagnosis of type 2 diabetes before screening • Management with diet and exercise only or treatment with oral antidiabetic monotherapy (except for thiazolidinedione) and willing to discontinue medication with 8 week washout period before randomisation • HbA1c level 7.0-10.0% inclusive at screening and randomisation for those on diet and exercise only; or HbA1c level 6.5-9.0% inclusive at screening and HbA1c level 7.0-10.0% inclusive at randomisation for those on oral antidiabetic monotherapy • BMI ≥18.5 to ≥35.0 kg/m2	registration	NCT01558271						
Study dates Community Sources of funding Eli Lilly Japan K.K., Kobe, Japan. Inclusion criteria • Diagnosis of type 2 diabetes before screening • Management with diet and exercise only or treatment with oral antidiabetic monotherapy (except for thiazolidinedione) and willing to discontinue medication with 8 week washout period before randomisation • HbA1c level 7.0-10.0% inclusive at screening and randomisation for those on diet and exercise only; or HbA1c level 6.5-9.0% inclusive at screening and HbA1c level 7.0-10.0% inclusive at randomisation for those on oral antidiabetic monotherapy • BMI ≥18.5 to ≥35.0 kg/m2 Exclusion	Study type	Randomised controlled trial (RCT)						
Study dates Sources of funding Inclusion criteria Diagnosis of type 2 diabetes before screening Management with diet and exercise only or treatment with oral antidiabetic monotherapy (except for thiazolidinedione) and willing to discontinue medication with 8 week washout period before randomisation HbA1c level 7.0-10.0% inclusive at screening and randomisation for those on diet and exercise only; or HbA1c level 6.5-9.0% inclusive at screening and HbA1c level 7.0-10.0% inclusive at randomisation for those on oral antidiabetic monotherapy BMI ≥18.5 to ≥35.0 kg/m2 Exclusion Output Diagnosis of type 1 diabetes.	Study location	Japan (33 sites in 14 cities)						
 Sources of funding Inclusion criteria Diagnosis of type 2 diabetes before screening Management with diet and exercise only or treatment with oral antidiabetic monotherapy (except for thiazolidinedione) and willing to discontinue medication with 8 week washout period before randomisation HbA1c level 7.0-10.0% inclusive at screening and randomisation for those on diet and exercise only; or HbA1c level 6.5-9.0% inclusive at screening and HbA1c level 7.0-10.0% inclusive at randomisation for those on oral antidiabetic monotherapy BMI ≥18.5 to ≥35.0 kg/m2 Exclusion 	Study setting	Community						
 Inclusion criteria Diagnosis of type 2 diabetes before screening Management with diet and exercise only or treatment with oral antidiabetic monotherapy (except for thiazolidinedione) and willing to discontinue medication with 8 week washout period before randomisation HbA1c level 7.0-10.0% inclusive at screening and randomisation for those on diet and exercise only; or HbA1c level 6.5-9.0% inclusive at screening and HbA1c level 7.0-10.0% inclusive at randomisation for those on oral antidiabetic monotherapy BMI ≥18.5 to ≥35.0 kg/m2 Exclusion 	Study dates	04/2012 to 10/2013						
 Management with diet and exercise only or treatment with oral antidiabetic monotherapy (except for thiazolidinedione) and willing to discontinue medication with 8 week washout period before randomisation HbA1c level 7.0-10.0% inclusive at screening and randomisation for those on diet and exercise only; or HbA1c level 6.5-9.0% inclusive at screening and HbA1c level 7.0-10.0% inclusive at randomisation for those on oral antidiabetic monotherapy BMI ≥18.5 to ≥35.0 kg/m2 Exclusion 		Eli Lilly Japan K.K., Kobe, Japan.						
J 71		 Management with diet and exercise only or treatment with oral antidiabetic monotherapy (except for thiazolidinedione) and willing to discontinue medication with 8 week washout period before randomisation HbA1c level 7.0-10.0% inclusive at screening and randomisation for those on diet and exercise only; or HbA1c level 6.5-9.0% inclusive at screening and HbA1c level 7.0-10.0% inclusive at randomisation for those on oral antidiabetic monotherapy 						

diabetes mellitus and chronic kidney

disease

Receiving more than half of maximum dose of sulfonylureas at screening Currently taking insulin or thiazolidinediones (TZD), or previous insulin or TZD treatment ≤3 months screening. Obvious clinical signs or symptoms of pancreatitis, history of chronic pancreatitis, or acute pancreatitis at screening, as determined by investigator Serum amylase concentration ≥3 times upper limit of reference range and/or a serum lipase concentration ≥2 times upper limit of reference range, as determined by central laboratory at screening History (personal or family) of medullary C-cell hyperplasia, focal hyperplasia, or medullary thyroid carcinoma Recruitment / Participants recruited from 33 sites in Japan. Initial 2 week screening selection of period, then 2 week lead-in period for treatment-naive participants and 8 participants week wash out period for participants on monotherapy. Eligible participants randomized to treatment in 4:2:1 ratio (dulaglutide; liraglutide; placebo) using computer-generated random sequence with interactive voice response system and stratified by pre-study oral antidiabetic medication status (yes/no), BMI group (<25; ≥25 kg/m2), and HbA1c (≤8.5; >8.5%). Participants and investigators masked to assignment to dulaglutide and placebo treatment but not masked to assignment to liraglutide treatment. At end of 26 weeks, participants in placebo group switched to dulaglutide 0.75 once weekly for remaining 26 weeks. Participants not tolerating study drugs discontinued them but remained in study to collect safety data. Intervention(s) Dulaglutide 0.75 mg once weekly Subcutaneous injection of dulaglutide 0.75 mg once weekly for 52 weeks, provided in non-identifiable solution in prefilled syringe and initiated at full dose. Strata 1: Not stated/unclear People with type 2 diabetes mellitus and heart failure Strata 2: Not stated/unclear People with atherosclerotic cardiovascular diseases Not stated/unclear Strata 3: People with type 2

Subgroup 1: People with frailty	Not stated/unclear
Subgroup 2: Onset of type 2 diabetes mellitus	Not stated/unclear
Subgroup 3: People with non-alcoholic fatty liver disease	Not stated/unclear
Subgroup 4: People with obesity	Not stated/unclear
Subgroup 5: eGFR category at baseline	Not stated/unclear
Subgroup 6: Albuminuria category at baseline	Not stated/unclear
Sensitivity analysis category: Enrichment trial status	7) Mixed population
Population subgroups	Inclusion criteria: treatment-naive (diabetes managed with diet and exercise only) or if on oral antidiabetic monotherapy had 8-week washout period
Comparator	 Liraglutide 0.9 mg once daily Open label subcutaneous liraglutide up-titrated from 0.3 mg/day during week 1 to 0.6 mg/day during week 2 and 0.9 mg/day at start of week 3 for remaining 49 weeks.
Number of participants	N=422 (Note that data in this article are only for the participants who were originally assigned to the dulaglutide or liraglutide; data does not include the participants who were assigned to placebo for 26 weeks and who subsequently switched to dulaglutide).
Duration of follow-up	52 weeks
Indirectness	Trial conducted in Japan, unlikely to be representative of UK population.
Method of analysis	Other Safety analysis conducted on as-treated population according to actual
	treatments received.

- **79.2.1. Dulaglutide 0.75 mg once weekly (N = 281)** Subcutaneous injection of dulaglutide 0.75 mg once weekly for 52 weeks.
- **79.2.2.** Liraglutide 0.9 mg once daily (N = 141) Subcutaneous injection of liraglutide 0.9 mg once daily for 52 weeks.

80. Pan, 2012

Bibliographic Reference

Pan, C. Y.; Yang, W.; Tou, C.; Gause-Nilsson, I.; Zhao, J.; Efficacy and safety of saxagliptin in drug-naïve Asian patients with type 2 diabetes mellitus: a randomized controlled trial; Diabetes Metab Res Rev; 2012; vol. 28 (no. 3); 268-75

Secondary publication of another included study- see primary study for details	NA	
Other publications associated with this study included in review	None	
Trial name / I registration number	NCT00698932	
Study type	Randomised controlled trial (RCT)	
Study location	International (40 sites in China, India, Philippines, and South Korea)	
Study setting (Community	
Study dates	06/2008 to 10/2009	
Sources of funding	Funded by Astra-Zeneca and Bristol-Myers Squibb.	
Inclusion criteria	 Men and non-pregnant, non-breastfeeding women aged≥18 years Diagnosis of type 2 diabetes Never received medication for type 2 diabetes (insulin or oral hypoglycaemic agents) or had received such for<6 months since diagnosis No anti-hyperglycaemic therapy for >3 consecutive days or 7 non-consecutive days during the 8 weeks (12 weeks for thiazolidinediones) before trial enrolment Fasting C-peptide level≥0.33 nmol/L and HbA1c 7.2–10.0% (55–86 mmol/mol) at lead-in and HbA1c 7.0–10.0% (53–86 mmol/mol) at randomization 	
Exclusion criteria	 Diagnosis of type 1 diabetes History of diabetic ketoacidosis or hyperosmolar non-ketonic coma Symptoms of poorly controlled diabetes 	

	 New York Heart Association class III or IV congestive heart failure or left ventricular ejection fraction of≤40% Significant cardiovascular history within 6-mo of visit 1 History of haemoglobinopathies Unstable or rapidly progressing renal disease based on investigator's clinical judgement Autoimmune skin disorder Gastro-intestinal surgery that could alter drug absorption Illegal drug or alcohol abuse in past year Any clinically significant abnormality that would compromise study participation Immunocompromised conditions (e.g. human immunodeficiency virus) Serum creatinine of≥1.4 mg/dL for women and ≥1.5 mg/dL for men at randomization Insulin therapy within 1 year of enrolment (with exception of its use during hospitalization or for gestational diabetes) Previous treatment with any DPP-4 inhibitor Currently receiving treatment with systemic glucocorticoids (other than replacement therapy or treatment with cytochrome P450 3A4 inducers)
Recruitment / selection of participants	Participants recruited from 40 centres (China, 19 sites; India, 8 sites; Philippines, 7 sites; South Korea, 6 sites). Initial enrolment period (visit 1), 4 week single-blind placebo-controlled lead-in period (during which laboratory measurements taken and diet/lifestyle counselling provided), and then, 24 week double-blind treatment period to visit 12. Diet/lifestyle modifications reinforced throughout trial period. Randomisation at visit 4 (week 0/baseline) by computer-generated allocation sequence, stratified by country.
Intervention(s)	Saxagliptin 5 mg once daily
	Oral saxagliptin 5 ng tablets once daily for 24 weeks.
Strata 1: People with type 2 diabetes mellitus and heart failure	Not stated/unclear Exclusion criteria: NYHA class 3 or 4. Trial could include participants with NYHA class 2 but no data provided.
Strata 2:	People without other cardiovascular diseases
People with atherosclerotic cardiovascular diseases	Exclusion criteria: significant cardiovascular history within 6 months of visit 1.
Strata 3: People with type 2 diabetes mellitus and chronic kidney disease	Not stated/unclear

Strata 4: People with type 2 diabetes mellitus and high cardiovascular risk	Not stated/unclear
Subgroup 1: People with frailty	Not stated/unclear
Subgroup 2: Onset of type 2 diabetes mellitus	Not stated/unclear
Subgroup 3: People with non-alcoholic fatty liver disease	Not stated/unclear
Subgroup 4: People with obesity	Not stated/unclear
Subgroup 5: eGFR category at baseline	Not stated/unclear
Subgroup 6: Albuminuria category at baseline	Not stated/unclear
Sensitivity analysis category: Enrichment trial status	7) Mixed population Inclusion criteria: Never received medication for type 2 diabetes (insulin or oral hypoglycaemic agents) or had received such for<6 months since diagnosis; and no anti-hyperglycaemic therapy for >3 consecutive days or 7 non-consecutive days during the 8 weeks (12 weeks for thiazolidinediones) before trial enrolment.
Population subgroups	
Comparator	 Placebo Matching placebo for 24 weeks. Identical tablets and packaging to saxagliptin treatment used.
Number of participants	N=568
Duration of follow-up	24 weeks
Method of analysis	Modified ITT

Efficacy analysis set included all randomised participants who received at least one dose of study drug, had baseline data, and data for at least one post-baseline efficacy endpoint, with data on or after rescue medication excluded. Safety analysis included all participants who received at least one dose of study drug and excluded data on or after rescue medication use.

80.2. Study arms

80.2.1. Saxagliptin 5 mg once daily (N = 284)

Oral saxagliptin 5 mg tablets once daily for 24 weeks.

80.2.2. Placebo (N = 284)

Matching oral placebo tablet for 24 weeks.

80.3. Characteristics

80.3.1. Arm-level characteristics

Characteristic	Saxagliptin 5 mg once daily (N = 284)	Placebo (N = 284)
% Male	n = 160; % = 56.3	n = 155; % = 54.6
Sample size		
Mean age (SD)	51.2 (10)	51.6 (10.3)
Mean (SD)		
Ethnicity Ethnicity not reported, only country of participation	n = NA ; % = NA	n = NA ; % = NA
Sample size		
China	n = 169; % = 59.5	n = 166; %
Sample size		= 58.5
India Sample size	n = 62; % = 21.8	n = 60 ; % = 21.1
•		
Philipines Sample size	n = 35 ; % = 12.3	n = 36 ; % = 12.7
South Korea	n = 18; % = 6.3	
Sample size	11 - 10 , 70 - 0.3	n = 22 ; % = 7.7
Sample Size		

Characteristic	Saxagliptin 5 mg once daily (N = 284)	Placebo (N = 284)
Comorbidities	NR	,
Custom value		NR
Presence of frailty	NR	MB
Custom value		NR
Time since type 2 diabetes diagnosis (years)	0.8 (1.4)	4.0.(0.0)
Mean (SD)		1.2 (2.6)
HbA1c (%)	8.1 (0.8)	0.0 (0.0)
Mean (SD)		8.2 (0.8)
Blood pressure	NR	ND
Custom value		NR
Heart rate	NR	ND
Custom value		NR
Smoking status	NR	ND
Custom value		NR
Alcohol consumption	NR	ND
Custom value		NR
Presence of severe mental illness	NR	ND
Custom value		NR
People with significant cognitive impairment	NR	ND
Custom value		NR
People with a learning disability	NR	NR
Custom value		INIX
Weight (kg)	69.2 (11.4)	60.2 (12.4)
Mean (SD)		69.2 (12.4)
BMI (kg/m2)	25.9 (3.4)	25.0 (2.7)
Mean (SD)		25.9 (3.7)
Number of people with obesity	NR	NR
Custom value		IVIX
Cholesterol and lipid levels (mmol/L) LDL-cholesterol data not represented in clinical	NA (NA)	NA (NA)

Characteristic	Saxagliptin 5 mg	Placebo (N
	once daily (N = 284)	= 284)
database for participants with triglyceride levels>400 mg/dL		
Mean (SD)		
Total cholesterol	5 (0.06)	5 (0.07)
Mean (SD)		,
HDL-cholesterol	1.2 (0.02)	1.1 (0.02)
Mean (SD)		
LDL-cholesterol	3 (0.05)	2.9 (0.06)
Mean (SD)		
Triglyceride	1.9 (0.08)	2.1 (0.11)
Mean (SD)		
Albumin creatinine ratio	NR	NR
Custom value		
eGFR (mL/min/1.73m2)	NR	NR
Custom value		
Other antidiabetic medication used	NR	NR
Custom value		
Blood pressure-lowering medication used	NR	NR
Custom value		
Statins/lipid-lowering medication used	NR	NR
Custom value		
Other treatment being received	NR	NR
Custom value		

81. Perez, 2009

Bibliographic Reference

Perez, A.; Zhao, Z.; Jacks, R.; Spanheimer, R.; Efficacy and safety of pioglitazone/metformin fixed-dose combination therapy compared with pioglitazone and metformin monotherapy in treating patients with T2DM; Curr Med Res Opin; 2009; vol. 25 (no. 12); 2915-23

Secondary publication of another included study- see primary study for details	No additional information.
Other publications associated with this study included in review	Perez, Alfonso, Jacks, Randal, Arora, Vipin et al. (2010) Effects of pioglitazone and metformin fixed-dose combination therapy on cardiovascular risk markers of inflammation and lipid profile compared with pioglitazone and metformin monotherapy in patients with type 2 diabetes. Journal of clinical hypertension (Greenwich, Conn.) 12(12): 973-82
Trial name / registration number	NCT00727857.
Study type	Randomised controlled trial (RCT)
Study location	Multicentre trial (134 sites in 6 countries).
Study setting	Outpatient follow-up.
Study dates	No additional information.
Sources of funding	Funded by Takeda Global Research & Development Center, Inc.
Inclusion criteria	At least 18 years of age with type 2 diabetes; baseline HbA1c 7.5%-10.0%; treatment naïve (had not received treatment with antidiabetic medication in the 12 weeks prior to screening other than short-term use of up to 15 days); body mass index no more than 45 kg/m2; had to have received counselling on lifestyle modification including diet and exercise.
Exclusion criteria	Type 1 diabetes; NYHA class III or IV heart failure; history of myocardial infarction, cerebrovascular accident, percutaneous coronary intervention, coronary artery bypass graft or transient ischaemic attack in the 6 months prior to screening; serum creatinine level equal to or more than 1.5mg/dL in men or equal to or more than 1.4mg/dL in women; a triglyceride level >500mg/dL; an ALT level greater than 2.5 times the upper limit of normal, active liver disease or jaundice; discontinuation from a thiazolidinedione or metformin therapy due to lack of efficacy, or clinical or laboratory signs of intolerance; pregnancy, intent to become pregnant, or lactation during the study in women.

Recruitment / selection of participants	People were withdrawn from the study and completed an early termination visit if they voluntarily withdrew; had an adverse event their imposed an unnecessary risk; major protocol deviation; loss to follow-up; pregnancy; lack of efficacy with FPG elevated from baseline and >230mg/dL from week 8-16; or lack of efficacy with HbA1c decreased <0.5% from baseline or level >8.5% at week 16 or later.
Intervention(s)	Pioglitazone + Metformin N=201
	Pioglitazone 15mg twice a day and metformin 850mg twice a day for 24 weeks.
	Concomitant therapy: Other antidiabetes medications were not allowed during the treatment period.
Strata 1: People with type 2 diabetes mellitus and heart failure	People without heart failure
Strata 2: People with atherosclerotic cardiovascular diseases	People without other cardiovascular diseases
Strata 3: People with type 2 diabetes mellitus and chronic kidney disease	People without chronic kidney disease
Strata 4: People with	People at higher risk of developing cardiovascular disease
type 2 diabetes mellitus and high cardiovascular risk	Based on BMI, presence of diabetes, age and smoking status.
Subgroup 1: People with frailty	Not stated/unclear
Subgroup 2: Onset of type 2 diabetes mellitus	Not stated/unclear
Subgroup 3: People with non-alcoholic	People without non-alcoholic fatty liver disease

fatty liver disease Subgroup 4: People with obesity Not stated/unclear Subgroup 5: GGRR category at baseline Not stated/unclear Subgroup 6: Albuminuria category at baseline Not stated/unclear Sensitivity analysis category: Enrichment trial status Population subgroups Comparator Pioglitazone N=189 Pioglitazone 15mg twice a day for 24 weeks. Concomitant therapy: Other antidiabetes medications were not allowed during the treatment period. Number of participants Concomitant therapy: Other antidiabetes medications were not allowed during the treatment period. Number of participants 600 Duration of follow-up Indirectness A weeks. Method of analysis No additional information. Method of analysis No additional information. Additional comments No additional information.		
People with obesity Subgroup 5: eGFR category at baseline Subgroup 6: Albuminuria category at baseline Sensitivity analysis category: Enrichment trial status Population subgroups Comparator Pioglitazone N=189 Pioglitazone 15mg twice a day for 24 weeks. Concomitant therapy: Other antidiabetes medications were not allowed during the treatment period. Metformin 850mg twice a day for 24 weeks. Concomitant therapy: Other antidiabetes medications were not allowed during the treatment period. Number of participants Duration of follow-up Indirectness Method of analysis Method of analysis Pioglitazone 15mg twice a day for 24 weeks.		
GEFR category at baseline Subgroup 6: Albuminuria category at baseline Sensitivity analysis category: Enrichment trial status Population subgroups Comparator Pioglitazone N=189 Pioglitazone 15mg twice a day for 24 weeks. Concomitant therapy: Other antidiabetes medications were not allowed during the treatment period. Metformin N=210 Metformin 850mg twice a day for 24 weeks. Concomitant therapy: Other antidiabetes medications were not allowed during the treatment period. Number of participants Duration of follow-up Indirectness Method of analysis Full case analysis - people had to receive at least one dose of the medication and have one analysis value. Last observation carried forward. Additional No additional information.	People with	Not stated/unclear
Albuminuria category at baseline Sensitivity analysis category: Enrichment trial status Population subgroups Comparator Pioglitazone N=189 Pioglitazone 15mg twice a day for 24 weeks. Concomitant therapy: Other antidiabetes medications were not allowed during the treatment period. Metformin N=210 Metformin 850mg twice a day for 24 weeks. Concomitant therapy: Other antidiabetes medications were not allowed during the treatment period. Number of participants Duration of follow-up Indirectness No additional information. Method of analysis Full case analysis - people had to receive at least one dose of the medication and have one analysis value. Last observation carried forward. Additional	eGFR category	Not stated/unclear
analysis category: Enrichment trial status Population subgroups Comparator Pioglitazone N=189 Pioglitazone 15mg twice a day for 24 weeks. Concomitant therapy: Other antidiabetes medications were not allowed during the treatment period. Metformin N=210 Metformin 850mg twice a day for 24 weeks. Concomitant therapy: Other antidiabetes medications were not allowed during the treatment period. Number of participants Duration of follow-up Indirectness Method of analysis Full case analysis - people had to receive at least one dose of the medication and have one analysis value. Last observation carried forward. Additional	Albuminuria category at	Not stated/unclear
Comparator Pioglitazone N=189 Pioglitazone 15mg twice a day for 24 weeks. Concomitant therapy: Other antidiabetes medications were not allowed during the treatment period. Metformin N=210 Metformin 850mg twice a day for 24 weeks. Concomitant therapy: Other antidiabetes medications were not allowed during the treatment period. Number of participants Duration of follow-up Indirectness No additional information. Other Full case analysis - people had to receive at least one dose of the medication and have one analysis value. Last observation carried forward. Additional	analysis category: Enrichment	
Pioglitazone 15mg twice a day for 24 weeks. Concomitant therapy: Other antidiabetes medications were not allowed during the treatment period. Metformin N=210 Metformin 850mg twice a day for 24 weeks. Concomitant therapy: Other antidiabetes medications were not allowed during the treatment period. Number of participants Duration of follow-up Indirectness No additional information. Method of analysis Full case analysis - people had to receive at least one dose of the medication and have one analysis value. Last observation carried forward. Additional	•	No additional information.
Concomitant therapy: Other antidiabetes medications were not allowed during the treatment period. Metformin N=210 Metformin 850mg twice a day for 24 weeks. Concomitant therapy: Other antidiabetes medications were not allowed during the treatment period. Number of participants Duration of follow-up Indirectness No additional information. Method of analysis Full case analysis - people had to receive at least one dose of the medication and have one analysis value. Last observation carried forward. Additional No additional information.	Comparator	Pioglitazone N=189
during the treatment period. Metformin N=210 Metformin 850mg twice a day for 24 weeks. Concomitant therapy: Other antidiabetes medications were not allowed during the treatment period. Number of participants Duration of follow-up Indirectness No additional information. Other Full case analysis - people had to receive at least one dose of the medication and have one analysis value. Last observation carried forward. Additional No additional information.		Pioglitazone 15mg twice a day for 24 weeks.
Metformin 850mg twice a day for 24 weeks. Concomitant therapy: Other antidiabetes medications were not allowed during the treatment period. Number of participants Duration of follow-up Indirectness No additional information. Method of analysis Full case analysis - people had to receive at least one dose of the medication and have one analysis value. Last observation carried forward. Additional No additional information.		· ·
Concomitant therapy: Other antidiabetes medications were not allowed during the treatment period. Number of participants Duration of follow-up Indirectness No additional information. Other Full case analysis - people had to receive at least one dose of the medication and have one analysis value. Last observation carried forward. Additional No additional information.		Metformin N=210
during the treatment period. Number of participants Duration of follow-up Indirectness No additional information. Method of analysis Full case analysis - people had to receive at least one dose of the medication and have one analysis value. Last observation carried forward. Additional		Metformin 850mg twice a day for 24 weeks.
Duration of follow-up Indirectness No additional information. Method of analysis Full case analysis - people had to receive at least one dose of the medication and have one analysis value. Last observation carried forward. Additional No additional information.		
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Additional No additional information.		Full case analysis - people had to receive at least one dose of the
		The state of the s

81.2.1. Pioglitazone + Metformin (N = 201)

Pioglitazone 15mg twice a day and metformin 850mg twice a day for 24 weeks. Concomitant therapy: Other antidiabetes medications were not allowed during the treatment period.

81.2.2. **Pioglitazone (N = 189)**

Pioglitazone 15mg twice a day for 24 weeks. Concomitant therapy: Other antidiabetes medications were not allowed during the treatment period.

81.2.3. Metformin (N = 210)

Metformin 850mg twice a day for 24 weeks. Concomitant therapy: Other antidiabetes medications were not allowed during the treatment period.

81.3. Characteristics

81.3.1. Arm-level characteristics

Characteristic	Pioglitazone + Metformin (N = 201)	Pioglitazone (N = 189)	Metformin (N = 210)
% Male	n = 90 ; % = 45	n = 66 ; % = 35	n = 98 ; % = 57
Sample size			
Mean age (SD) (years)	54.7 (12.2)	54 (12.1)	53.7 (12)
Mean (SD)			
Ethnicity	n = NA ; % = NA	n = NA ; % = NA	· ·
Sample size			NA
Hispanic or Latino	n = 49 ; % = 24.4	n = 49 ; % = 25.9	·
Sample size			26.2
Non-Hispanic and Non- Latino	n = 42; % = 20.9	n = 36 ; % = 19	n = 46 ; % = 21.9
Sample size			
American Indian or Alaska Native	n = 63; % = 31.3	n = 62 ; % = 32.8	n = 67 ; % = 31.9
Sample size			

Characteristic	Pioglitazone + Metformin (N = 201)	Pioglitazone (N = 189)	Metformin (N = 210)
Asian	n = 3; % = 1.5	n = 5; % = 2.6	n = 5 ; % = 2.4
Sample size			
Black or African American	n = 12; % = 6	n = 13; % = 6.9	n = 14 ; % = 6.7
Sample size	404 - 0/ 04 5		
White Sample size	n = 184 ; % = 91.5	n = 165; % = 87.3	n = 185 ; % = 88.1
Multiracial	n = 61; % = 30.3		
Sample size	01, 70 00.0	n = 56; % = 29.6	n = 61; % = 29
Comorbidities	n = NA ; % = NA		
Sample size	,	n = NA ; % = NA	n = NA ; % = NA
Hypertension	n = 91; % = 45.3	n = 95 ; % = 50	n = 94 ; % = 45
Sample size			
Hyperlipidaemia	n = 90 ; % = 44.8	n = 85; % = 44.7	n = 95 ; % = 45.5
Sample size			.0.0
Heart failure	n = 0; % = 0	n = 1; % = 0.5	n = 1; % = 0.5
Sample size	ND 0/ ND		
Presence of frailty Sample size	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
Time since type 2 diabetes	n = NR ; % = NR		
diagnosis	11 - IVIX , 70 - IVIX	n = NR ; % = NR	n = NR ; % = NR
Sample size			
HbA1c (%)	8.89 (0.07)	8.69 (0.07)	8.65 (0.07)
Mean (SE)			
Blood pressure	NR (NR)	NR (NR)	NR (NR)
Mean (SD)			
Heart rate	NR (NR)	NR (NR)	NR (NR)
Mean (SD)	ND 0/ ND		
Sample size	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
Sample size			

Characteristic	Pioglitazone + Metformin (N = 201)	Pioglitazone (N = 189)	Metformin (N = 210)
Alcohol consumption	n = NR ; % = NR	n = NR ; % = NR	· ·
Sample size			NR
Presence of severe mental illness	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
Sample size			
People with significant cognitive impairment	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
Sample size			
People with a learning disability	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
Sample size			
Weight	NR (NR)	NR (NR)	NR (NR)
Mean (SD)			
BMI (kg/m2)	30.8 (5.7)	31.2 (5.5)	30.8 (5.7)
Mean (SD)			
Number of people with obesity	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
Sample size			
Cholesterol and lipid levels (mg/dL)	NA (NA)	NA (NA)	NA (NA)
Mean (SE)			
Total cholesterol	195.5 (2.85)	198.4 (2.95)	196 (2.79)
Mean (SE)			
HDL cholesterol	41.8 (0.76)	43.4 (0.79)	42.5 (0.75)
Mean (SE)			
LDL cholesterol	118.3 (2.38)	118.9 (2.47)	117.1 (2.34)
Mean (SE)			
Triglycerides	202.3 (10.17)	204.2 (10.49)	213.7 (9.95)
Mean (SE)			
Albumin creatinine ratio	NR (NR)	NR (NR)	NR (NR)
Mean (SD)			

Characteristic	Pioglitazone + Metformin (N = 201)	Pioglitazone (N = 189)	Metformin (N = 210)
eGFR (mL/min/1.73m2)	NR (NR)	NR (NR)	NR (NR)
Mean (SD)		,	, ,
Other antidiabetic medication used	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
Sample size			
Blood pressure-lowering medication used	n = NA ; % = NA	n = NA ; % = NA	n = NA ; % = NA
Sample size			
ACEIs/ARBs	n = 57; % = 28.4	n = 69 ; % = 36.3	n = 68 ; % = 32.5
Sample size	07 0/ 40 4		02.0
Diuretics Sample size	n = 27; % = 13.4	n = 28 ; % = 14.7	n = 28 ; % = 13.4
Beta blockers	n = 19 ; % = 9.5		
Dota biookoro	10,70 0.0	n = 20 ; % = 10.5	· ·
Sample size			8.6
Calcium channel blockers Sample size	n = 17; % = 8.5	n = 16; % = 8.4	n = 22 ; % = 10.5
Statins/lipid-lowering	n = 30 ; % = 14.9		
medication used	11 - 30 , 70 - 14.3	n = 26 ; % = 13.7	n = 39 ; % = 18.7
Sample size			
Other treatment being received	n = NA ; % = NA	n = NA ; % = NA	n = NA ; % = NA
Sample size			
Antiplatelets	n = 33; % = 16.4	n = 29 ; % = 15.3	n = 34 ; % = 16.3
Sample size			10.0
NSAIDs	n = 23 ; % = 11.4	n = 23 ; % = 12.1	n = 25 ; % = 12
Sample size			

82. Perez, 2010

Bibliographic Reference

Perez, Alfonso; Jacks, Randal; Arora, Vipin; Spanheimer, Robert; Effects of pioglitazone and metformin fixed-dose combination therapy on cardiovascular risk markers of inflammation and lipid profile compared with pioglitazone and metformin monotherapy in patients with type 2 diabetes.; Journal of clinical hypertension (Greenwich, Conn.); 2010; vol. 12 (no. 12); 973-82

Secondary publication of another included study- see primary study for details	Perez, A., Zhao, Z., Jacks, R. et al. (2009) Efficacy and safety of pioglitazone/metformin fixed-dose combination therapy compared with pioglitazone and metformin monotherapy in treating patients with T2DM. Curr Med Res Opin 25(12): 2915-23
Other publications associated with this study included in review	No additional information.

83. Pfützner, 2011

Bibliographic Reference

Pfützner, A.; Paz-Pacheco, E.; Allen, E.; Frederich, R.; Chen, R.; Initial combination therapy with saxagliptin and metformin provides sustained glycaemic control and is well tolerated for up to 76 weeks; Diabetes Obes Metab; 2011; vol. 13 (no. 6); 567-76

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Secondary publication of another included study- see primary study for details	No additional information.
Other publications associated with this study included in review	Jadzinsky, M, Pfutzner, A, Paz-Pacheco, E et al. (2009) Saxagliptin given in combination with metformin as initial therapy improves glycaemic control in patients with type 2 diabetes compared with either monotherapy: a randomized controlled trial. Diabetes, obesity & metabolism 11(6): 611-22
Trial name / registration number	CV181-039. NCT00327015.
Study type	Randomised controlled trial (RCT)
Study location	Multicentre trial.
Study setting	Outpatient follow-up.
Study dates	No additional information.
Sources of funding	Funded, designed and supervised by Bristol-Myers Squibb and AstraZeneca.
Inclusion criteria	People who were treatment-naïve (people who have never received treatment for diabetes or who had received medical treatment for less than 1 month since original diagnosis and who had not received therapy for more than three consecutive days or seven non-consecutive days during the 8 weeks before screening) with type 2 diabetes; 18-77 years of age; inadequate glycaemic control (HbA1c 8.0-12.0%); BMI no more than 40kg/m2; fasting C-peptide concentration at least 1.0ng/mL.
Exclusion criteria	Symptoms of poorly controlled diabetes; history of diabetic ketoacidosis or hyperosmolar nonketotic coma; insulin therapy within 1 year of screening; cardiovascular event within the prior 6 months or NYHA stage III/IV congestive heart failure and/or known LVEF less than or equal to 40%; significant history of renal or hepatic disease or a psychiatric disorder; alcohol or drug abuse within the previous year; treatment with potent CYP3A4 inhibitors or inducers; immunocompromised individuals; active liver disease or clinically significant abnormal hepatic, renal, endocrine, metabolic or haematological screening tests.

Recruitment / People who completed all visits during the 24-week period or who met progressively strict glycaemic rescue criteria (FPG >240mg/dL at week 6, selection of >220mg/dL at week 8, >200mg/dL at week 12 onwards) entered the longparticipants term period. People who were rescued in the 24 week phase were advanced directly to the 52 week extension period and received open label pioglitazone 15mg once daily (titratable to 45mg once daily) added to their blinded study medication. No changes in metformin dose was permitted. People with HbA1c >8.0% at week 30, 37 or 50 or >7.5% at week 63 were similarly rescued with pioglitazone. People who did not have a reduction in FPG of at least 30mg/dL within 8 weeks of starting rescue therapy were discontinued from the study and referred for additional antihyperglycaemic intervention. Intervention(s) Saxagliptin + Metformin N=643 Two groups: Saxagliptin 10mg + metformin initially 500mg daily (n=323) and saxagliptin 5mg + metformin initially 500mg daily (n=320). Metformin was titrated to 1000mg/day at week 1, if FPG was >110mg/day and it was tolerated then it was increased in 500mg/day increments during weeks 2-5 to a maximum of 2000mg/day. Saxagliptin was taken once daily before the morning meal, metformin was taken in divided doses with the morning and evening meals. Concomitant therapy: Pioglitazone 15mg once daily (titratable to 45mg once daily) was provided as rescue therapy to anyone who required it. Strata 1: People without heart failure People with type 2 diabetes mellitus and heart failure Strata 2: People without other cardiovascular diseases People with atherosclerotic cardiovascular diseases Strata 3: People without chronic kidney disease People with type 2 diabetes mellitus and chronic kidney disease Strata 4: People at higher risk of developing cardiovascular disease People with type 2 Based on BMI and presence of diabetes diabetes mellitus and high cardiovascular risk

Subgroup 1: People with frailty	Not stated/unclear
Subgroup 2: Onset of type 2 diabetes mellitus	Not stated/unclear
Subgroup 3: People with non-alcoholic fatty liver disease	People without non-alcoholic fatty liver disease
Subgroup 4: People with obesity	Not stated/unclear
Subgroup 5: eGFR category at baseline	Not stated/unclear
Subgroup 6: Albuminuria category at baseline	Not stated/unclear
Sensitivity analysis category: Enrichment trial status	7) Mixed population
Population subgroups	No additional information.
Comparator	Saxagliptin N=335
	Saxagliptin 10mg daily with placebo. Saxagliptin was taken once daily before the morning meal.
	Concomitant therapy: Pioglitazone 15mg once daily (titratable to 45mg once daily) was provided as rescue therapy to anyone who required it.
	Metformin N=328
	Metformin initially 500mg daily with placebo. Metformin was titrated to 1000mg/day at week 1, if FPG was >110mg/day and it was tolerated then it was increased in 500mg/day increments during weeks 2-5 to a maximum of 2000mg/day. Metformin was taken in divided doses with the morning and evening meals.

	Concomitant therapy: Pioglitazone 15mg once daily (titratable to 45mg once daily) was provided as rescue therapy to anyone who required it.
Number of participants	1306
Duration of follow-up	72 weeks.
Indirectness	No additional information.
Method of analysis	Other Full case analysis - people who received at least one dose of treatment and had at least one efficacy measurement taken before rescue medication was used. Last observation carried forward.
Additional comments	No additional information.

83.2.1. Saxagliptin + Metformin (N = 643)

Two groups: Saxagliptin 10mg + metformin initially 500mg daily (n=323) and saxagliptin 5mg + metformin initially 500mg daily (n=320). Metformin was titrated to 1000mg/day at week 1, if FPG was >110mg/day and it was tolerated then it was increased in 500mg/day increments during weeks 2-5 to a maximum of 2000mg/day. Saxagliptin was taken once daily before the morning meal, metformin was taken in divided doses with the morning and evening meals. Concomitant therapy: Pioglitazone 15mg once daily (titratable to 45mg once daily) was provided as rescue therapy to anyone who required it.

83.2.2. Saxagliptin (N = 335)

Saxagliptin 10mg daily with placebo. Saxagliptin was taken once daily before the morning meal. Concomitant therapy: Pioglitazone 15mg once daily (titratable to 45mg once daily) was provided as rescue therapy to anyone who required it.

83.2.3. Metformin (N = 328)

Metformin initially 500mg daily with placebo. Metformin was titrated to 1000mg/day at week 1, if FPG was >110mg/day and it was tolerated then it was increased in 500mg/day increments during weeks 2-5 to a maximum of 2000mg/day. Metformin was taken in divided doses with the morning and evening meals. Concomitant therapy: Pioglitazone 15mg once daily (titratable to 45mg once daily) was provided as rescue therapy to anyone who required it.

83.3. Characteristics

83.3.1. Arm-level characteristics

83.3.1. Arm-level	characteristics		
Characteristic	Saxagliptin + Metformin (N = 643)	Saxagliptin (N = 335)	Metformin (N = 328)
% Male Sample size	n = 311 ; % = 48	n = 169 ; % = 50	n = 163 ; % = 50
	FO 4 (44)		
Mean age (SD) (years)	52.1 (11)	52.1 (10.2)	51.8 (10.7)
Mean (SD)			
Ethnicity Sample size	n = NA ; % = NA	n = NA ; % = NA	n = NA ; % = NA
	400 0/ 70		
White Sample size	n = 489 ; % = 76	n = 255 ; % = 76	n = 251 ; % = 77
	405 - 0/ 40		
Asian	n = 105 ; % = 16	n = 56 ; % = 17	n = 52 ; % = 16
Sample size			
Black/African American	n = 14; % = 2	n = 6; % = 2	n = 4 ; % = 1
Sample size			
Other	n = 35; % = 5	n = 18 ; % = 5	n = 21 ; % = 6
Sample size			
Comorbidities	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
Sample size			Turk
Presence of frailty Sample size	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
Time since type 2 diabetes	1.7 (3.1)		
diagnosis (years)	(6.1)	1.7 (2.8)	1.7 (3.1)
Mean (SD)			
HbA1c (%)	9.5 (1.2)	9.6 (1.3)	9.4 (1.3)
Mean (SD)			
Blood pressure	NR (NR)	NR (NR)	NR (NR)
Mean (SD)			
Heart rate	NR (NR)	NR (NR)	NR (NR)
Mean (SD)			

Characteristic	Saxagliptin + Metformin (N = 643)	Saxagliptin (N = 335)	Metformin (N = 328)
Smoking status	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
Sample size			INIX
Alcohol consumption	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
Sample size			IVIX
Presence of severe mental illness	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
Sample size			
People with significant cognitive impairment Sample size	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
·	- ND : 0/ - ND		
People with a learning disability	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
Sample size	ND (ND)		
Weight	NR (NR)	NR (NR)	NR (NR)
Mean (SD)			
BMI (kg/m2)	30.2 (4.7)	30.2 (4.9)	30.2 (4.9)
Mean (SD)			
Number of people with obesity	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
Sample size			
Cholesterol and lipid levels Mean (SD)	NR (NR)	NR (NR)	NR (NR)
Albumin creatinine ratio	NR (NR)		
Mean (SD)	INIX (INIX)	NR (NR)	NR (NR)
eGFR (mL/min/1.73m2)	NR (NR)	NR (NR)	NR (NR)
Mean (SD)			
Other antidiabetic medication used	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
Sample size			
Blood pressure-lowering medication used	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
Sample size			

Characteristic	Saxagliptin + Metformin (N = 643)	Saxagliptin (N = 335)	Metformin (N = 328)
Statins/lipid-lowering medication used Sample size	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
Other treatment being received Sample size	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR

84. Pistrosch, 2013

Bibliographic Reference

Pistrosch, F.; Köhler, C.; Schaper, F.; Landgraf, W.; Forst, T.; Hanefeld, M.; Effects of insulin glargine versus metformin on glycemic variability, microvascular and beta-cell function in early type 2 diabetes; Acta Diabetol; 2013; vol. 50 (no. 4); 587-95

0 4 .1. 3	tudy details
Secondary publication of another included study- see primary study for details	No additional information.
Other publications associated with this study included in review	No additional information.
Trial name / registration number	NCT00857870.
Study type	Randomised controlled trial (RCT)
Study location	Multicentre trial.
Study setting	Outpatient follow-up.
Study dates	No additional information.
Sources of funding	Funded by Sanofi-Aventis, Germany.
Inclusion criteria	Drug naïve type 2 diabetes mellitus with <5 years after diagnosis and a HbA1c between 6.5 and 8%.
Exclusion criteria	Renal dysfunction with a calculated glomerular filtration rate below 60mL/min; acute or chronic diseases which could lead to tissue hypoxia; the use of intravascular contrast agents throughout the study; increase in serum transaminases more than 2.5 times the upper limit of normal; systemic corticosteroid treatment.
Recruitment / selection of participants	No additional information.
Intervention(s)	Metformin N=36
	Metformin 1000mg twice a day.

	Concomitant therapy: No additional information.
Strata 1: People with type 2 diabetes mellitus and heart failure	Not stated/unclear
Strata 2: People with atherosclerotic cardiovascular diseases	Not stated/unclear
Strata 3: People with type 2 diabetes mellitus and chronic kidney disease	Not stated/unclear
People with type 2	People at higher risk of developing cardiovascular disease Based on BMI, systolic blood pressure, age and presence of type 2 diabetes.
Subgroup 1: People with frailty	Not stated/unclear
Subgroup 2: Onset of type 2 diabetes mellitus	Not stated/unclear
Subgroup 3: People with non-alcoholic fatty liver disease	Not stated/unclear
Subgroup 4: People with obesity	Not stated/unclear
Subgroup 5: eGFR category at baseline	Not stated/unclear
Subgroup 6: Albuminuria	Not stated/unclear

84.2.1. Metformin (N = 36)

Metformin 1000mg twice a day. Concomitant therapy: No additional information.

84.2.2. Insulin (N = 39)

Insulin glargine at bedtime for 36 weeks. Insulin dose was titrated stepwise to a target fasting glucose of no more than 5.6 mmol/L according to a standardized titrated schedule. Concomitant therapy: No additional information.

84.3. Characteristics

84.3.1. Arm-level characteristics

Characteristic	Metformin (N = 36)	Insulin (N = 39)
% Male	n = 18; % = 50	n = 26 ; % = 66.7
Sample size		
Mean age (SD) (years)	62.03 (9.4)	60 (9.3)
Mean (SD)		
Ethnicity	n = NR ; % = NR	n = NR ; % = NR
Sample size		
Comorbidities Sample size	n = NR ; % = NR	n = NR ; % = NR
Sample size	n - ND · 0/ - ND	
Presence of frailty Sample size	n = NR ; % = NR	n = NR ; % = NR
Time since type 2 diabetes diagnosis (years)	2.6 (1.6)	2.8 (1.4)
Mean (SD)		
HbA1c (%)	6.9 (0.4)	6.36 (0.4)
Mean (SD)		
Blood pressure (mmHg) Mean (SD)	NA (NA)	NA (NA)
Systolic blood pressure	141.5 (14.8)	
Mean (SD)	141.5 (14.0)	141 (15.7)
Diastolic blood pressure	81.2 (10.4)	
Mean (SD)	01.2 (10.4)	85.3 (9.8)
Heart rate	NR (NR)	
Mean (SD)	TVIC (IVIC)	NR (NR)
Smoking status	n = NR ; % = NR	n = NR ; % = NR
Sample size		
Alcohol consumption	n = NR ; % = NR	n = NR ; % = NR
Sample size	N.B. 67	
Presence of severe mental illness	n = NR ; % = NR	n = NR ; % = NR

Characteristic	Metformin (N = 36)	Insulin (N = 39)
Sample size		
People with significant cognitive impairment	n = NR ; % = NR	n = NR ; % = NR
Sample size		
People with a learning disability	n = NR ; % = NR	n = NR ; % = NR
Sample size		
Weight (kg)	87.6 (17.9)	87.6 (15.1)
Mean (SD)	00.0 (5.0)	
BMI (kg/m2) Mean (SD)	29.9 (5.3)	29.2 (4.6)
Number of people with obesity	n = NR ; % = NR	
Number of people with obesity	11 - 1410, 70 - 1410	n = NR; % = NR
Sample size		
Cholesterol and lipid levels	NR (NR)	NR (NR)
Mean (SD)		
Albumin creatinine ratio	NR (NR)	NR (NR)
Mean (SD)		
eGFR (mL/min/1.73m2)	NR (NR)	NR (NR)
Mean (SD)	ND - 0/ - ND	
Other antidiabetic medication used Sample size	n = NR ; % = NR	n = NR ; % = NR
Blood pressure-lowering medication used	n = NR ; % = NR	n = NR ; % = NR
Sample size		
Statins/lipid-lowering medication used	n = NR ; % = NR	n = NR ; % = NR
Sample size		
Other treatment being received	n = NR ; % = NR	n = NR ; % = NR
Sample size		

85. Pi-Sunyer, 2007

Bibliographic Reference

Pi-Sunyer, F. X.; Schweizer, A.; Mills, D.; Dejager, S.; Efficacy and tolerability of vildagliptin monotherapy in drug-naïve patients with type 2 diabetes; Diabetes Res Clin Pract; 2007; vol. 76 (no. 1); 132-8

85.1. S	tudy details		
Secondary publication of another included study- see primary study for details	NA		
Other publications associated with this study included in review	None		
Trial name / registration number	NCT00120536, CLAF237A2384		
Study type	Randomised controlled trial (RCT)		
Study location	International (98 centres in US, India, and Slovakia)		
Study setting	Community		
Study dates	06/2005 to 07/2006		
Sources of funding	Funded by Novartis Pharmaceuticals Corporation.		
Inclusion criteria	 Aged 18-80 years If female, non-fertile or of childbearing potential using medically-approved birth control method Oral antidiabetic drug-naive (Not receiving oral antidiabetic drugs≥12 weeks prior to screening and not received oral antidiabetic drug>3 consecutive months at any time in the past) HbA1c level 7.5-10% inclusive BMI 22-45 kg/m2 inclusive FPG<15 mmol/L 		
Exclusion criteria	 History of type 1 or secondary forms of diabetes Acute metabolic diabetic complications Serious cardiovascular events in previous 6 months (e.g. Myocardial infarction, unstable angina, or coronary artery bypass surgery) Congestive heart failure, NYHA Class III or IV Liver disease (e.g. cirrhosis or chronic active hepatitis) 		

	 ALT or AST >3 times the upper limit of normal (ULN) Direct bilirubin>1.3 times the ULN Serum creatinine levels>220mmol/L Clinically significant abnormal thyroid stimulating hormone Fasting triglycerides>7.9 mmol/L
Recruitment / selection of participants	Participants recruited from 98 centres in USA (88 centres), India (4 centres) and Slovakia (6 centres). Two-week screening period, and 24-week treatment period. No information about randomisation method provided. Efficacy and tolerability assessed at weeks 4, 8, 12, 16 and 24.
Intervention(s)	 Vildagliptin 50 mg once daily Vildagliptin 50 mg twice daily Vildagliptin 100 mg once daily Oral vildagliptin for 24 weeks.
Strata 1: People with type 2 diabetes mellitus and heart failure	Not stated/unclear Exclusion criteria: NYHA class 3 or 4. Trial may have included some participants who were NYHA class 2.
Strata 2: People with atherosclerotic cardiovascular diseases	People without other cardiovascular diseases Exclusion criteria: Serious cardiovascular events in previous 6 months (e.g. Myocardial infarction, unstable angina, or coronary artery bypass surgery)
Strata 3: People with type 2 diabetes mellitus and chronic kidney disease	Not stated/unclear
Strata 4: People with type 2 diabetes mellitus and high cardiovascular risk	Not stated/unclear
Subgroup 1: People with frailty	Not stated/unclear
Subgroup 2: Onset of type 2 diabetes mellitus	Not stated/unclear

Not stated/unclear
Not stated/unclear
Not stated/unclear
Not stated/unclear
7) Mixed population Inclusion criteria: as no oral antidiabetic drug for at least 12-mo prior to screening and no OAD for >3 consecutive months at any time in past.
 Placebo No details reported but double-blind trial so presumably matching placebo for 24 weeks.
N=354
24 weeks
None
Modified ITT Primary and secondary efficacy analysis conducted on all randomised participants who received at least one dose of study drug and had baseline and at least one post-baseline HbA1c assessment.

85.2.1. Vildagliptin 100 mg once daily (N = 91)

Oral vildagliptin tablet 100 mg once daily for 24 weeks.

85.2.2. Vildagliptin 50 mg twice daily (N = 83)

Oral vildagliptin tablet 50 mg once twice daily for 24 weeks.

85.2.3. Vildagliptin 50 mg once daily (N = 88)

Oral vildagliptin tablet 50 mg once daily for 24 weeks.

85.2.4. Placebo (N = 92)

Matching placebo for 24 weeks.

85.3. Characteristics

85.3.1. Arm-level characteristics

85.3.1. Al	rm-level characte	ristics		
Characteristic	Vildagliptin 100 mg once daily (N = 91)		Vildagliptin 50 mg once daily (N = 88)	Placebo (N = 92)
% Male	n = 49 ; % = 53.8	n = 47 ; % = 56.6	n = 49 ; % = 55.7	n = 50; % = 54.3
Sample size				01.0
Mean age (SD) (year)	52 (11.7)	50.2 (12.7)	50.6 (10.4)	52 (12)
Mean (SD)				
Ethnicity Sample size	n = NA ; % = NA	n = NA ; % = NA	n = NA ; % = NA	n = NA ; % = NA
	· - 45 · 0/ - 40 5			
Asian (Indian subcontinent)	n = 15; % = 16.5	n = 15 ; % = 18.1	n = 14; % = 15.9	n = 15; % = 16.3
Sample size				
Asian (non-Indian subcontinent)	n = 1; % = 1.1	n = 1 ; % = 1.2	n = 3; % = 3.4	n = 1; % = 1.1
Sample size				
Black	n = 11 ; % = 12.1	n = 5; % = 6	n = 7; % = 8	n = 12; %
Sample size				= 13
Caucasian	n = 53 ; % = 58.2	n = 44 ; % = 53	n = 48 ; % = 54.5	
Sample size				= 51.1
Hispanic or Latino	n = 11; % = 12.1	n = 18 ; % = 21.7	n = 16 ; % = 18.2	
Sample size				= 18.5
Comorbidities	NR	NR	NR	NR
Custom value				
Presence of frailty	NR	NR	NR	NR

Characteristic	Vildagliptin 100 mg once daily (N = 91)	Vildagliptin 50 mg twice daily (N = 83)	Vildagliptin 50 mg once daily (N = 88)	Placebo (N = 92)
Custom value				
Time since type 2 diabetes diagnosis (years)	2.1 (2.9)	2.4 (3.2)	1.8 (2.7)	2.5 (3.7)
Mean (SD)				
HbA1c (%)	8.3 (0.8)	8.4 (0.9)	8.4 (0.9)	8.5 (0.8)
Mean (SD)				
Blood pressure	NR	NR	NR	NR
Custom value				
Heart rate	NR	NR	NR	NR
Custom value				
Smoking status	NR	NR	NR	NR
Custom value				
Alcohol consumption	NR	NR	NR	NR
Custom value				
Presence of severe mental illness	NR	NR	NR	NR
Custom value				
People with significant cognitive impairment	NR	NR	NR	NR
Custom value				
People with a learning disability	NR	NR	NR	NR
Custom value				
Weight (kg)	90.8 (19.9)	89.9 (18.5)	90.5 (22.3)	93 (23.2)
Mean (SD)	04.0 (=)			
BMI (kg/m2)	31.9 (5)	32.3 (6)	31.9 (5.4)	32.7 (6.4)
Mean (SD)	ND			
Number of people with obesity	NR	NR	NR	NR
Custom value				

Characteristic	Vildagliptin 100 mg once daily (N = 91)	Vildagliptin 50 mg twice daily (N = 83)	Vildagliptin 50 mg once daily (N = 88)	Placebo (N = 92)
Cholesterol and lipid levels	NR	NR	NR	NR
Custom value				
Albumin creatinine ratio Custom value	NR	NR	NR	NR
	ND			
eGFR (mL/min/1.73m2)	NR	NR	NR	NR
Custom value				
Other antidiabetic medication used	NR	NR	NR	NR
Custom value				
Blood pressure- lowering medication used Custom value	NR	NR	NR	NR
	NR			
Statins/lipid-lowering medication used	INK	NR	NR	NR
Custom value				
Other treatment being received	NR	NR	NR	NR
Custom value				

86. Pratley, 2014

Bibliographic Reference

Pratley, R. E.; Fleck, P.; Wilson, C.; Efficacy and safety of initial combination therapy with alogliptin plus metformin versus either as monotherapy in drug-naive patients with type 2 diabetes: A randomized, double-blind, 6-month study; Diabetes Obes Metab; 2014; vol. 16 (no. 7); 613-621

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Secondary publication of another included study- see primary study for details	No additional information.
Other publications associated with this study included in review	No additional information.
Trial name / registration number	NCT01023581.
Study type	Randomised controlled trial (RCT)
Study location	Multisite trial (198 sites worldwide).
Study setting	Outpatient follow-up.
Study dates	No additional information.
Sources of funding	Sponsored by Takeda Development Center Americas, Inc., Deerfield, IL, USA, and Takeda Development Centre Europe Ltd., London, UK.
Inclusion criteria	Aged 18-80 years with inadequately controlled T2DM (HbA1c 7.5-10%) following diet/exercise therapy alone for at least 2 months prior to screening; had taken fewer than 7 days of any antidiabetic medication within 2 months of screening; had a body mass index of 23-45 kg/m2 (20-35 kg/m2 for Asian participants) and a minimum fasting C-peptide of 0.8ng/mL. Just prior to randomisation (week -1 of the placebo run-in/stabilisation period), additional requirements were: HbA1c 7.5-10%, inclusive (weekly repeat tests permitted up to 2 weeks); 75-125% study drug compliance during run-in/stabilisation; no use of oral or systemically injected corticosteroids or weight-loss drugs.
Exclusion criteria	Low haemoglobin; elevated blood pressure; heart failure (NYHA class III-IV); coronary angioplasty, stent placement, bypass surgery or myocardial infarction within 3 months of screening; elevated ALT or serum creatinine or low creatinine clearance.

Recruitment / selection of participants	No additional information.
Intervention(s)	Alogliptin + Metformin N=225
	Two groups: Alogliptin 12.5mg twice a day and metformin 1000mg twice a day (n=114) and alogliptin 12.5mg twice a day and metformin 500mg twice a day (n=111).
	Concomitant therapy: Rescue medication was a sulfonylurea (specific type/dose to be chosen by the investigator), which was added to the double-blind study medication regimen, or another medication if it was contraindicated or inappropriate.
Strata 1: People with type 2 diabetes mellitus and heart failure	People without heart failure
Strata 2: People with atherosclerotic cardiovascular diseases	People without other cardiovascular diseases
Strata 3: People with type 2 diabetes mellitus and chronic kidney disease	People without chronic kidney disease
Strata 4: People with	People at higher risk of developing cardiovascular disease
type 2 diabetes mellitus and high	Based on BMI, triglycerides and presence of diabetes.
cardiovascular risk	
Subgroup 1: People with frailty	Not stated/unclear
Subgroup 2: Onset of type 2 diabetes mellitus	Not stated/unclear
Subgroup 3: People with non-alcoholic	People without non-alcoholic fatty liver disease

fatty liver disease	
Subgroup 4: People with obesity	Not stated/unclear
Subgroup 5: eGFR category at baseline	Not stated/unclear
Subgroup 6: Albuminuria category at baseline	Not stated/unclear
Sensitivity analysis category: Enrichment trial status	7) Mixed population
Population subgroups	No additional information.
Comparator	Metformin N=225
	Two groups: Metformin 1000mg twice a day (n=111) and metformin 500mg twice a day (n=114). Concomitant therapy: Rescue medication was a sulfonylurea (specific type/dose to be chosen by the investigator), which was added to the double-blind study medication regimen, or another medication if it was contraindicated or inappropriate. Alogliptin N=225 Two groups: Alogliptin 25mg once a day (n=112) and alogliptin 12.5mg twice a day (n=113).
	Concomitant therapy: Rescue medication was a sulfonylurea (specific type/dose to be chosen by the investigator), which was added to the double-blind study medication regimen, or another medication if it was contraindicated or inappropriate. Placebo N=109 Matching placebo. Concomitant therapy: Rescue medication was a sulfonylurea (specific type/dose to be chosen by the investigator), which was added to the
	double-blind study medication regimen, or another medication if it was contraindicated or inappropriate.

Number of participants	784
Duration of follow-up	26 weeks.
Indirectness	No additional information.
Method of analysis	Other
	Full analysis set - only people who received at least one dose of the intervention and were followed up with at least one measurement.
Additional comments	No additional information.

86.2.1. Alogliptin + Metformin (N = 225)

Two groups: Alogliptin 12.5mg twice a day and metformin 1000mg twice a day (n=114) and alogliptin 12.5mg twice a day and metformin 500mg twice a day (n=111). Concomitant therapy: Rescue medication was a sulfonylurea (specific type/dose to be chosen by the investigator), which was added to the double-blind study medication regimen, or another medication if it was contraindicated or inappropriate.

86.2.2. Metformin (N = 225)

Two groups: Metformin 1000mg twice a day (n=111) and metformin 500mg twice a day (n=114). Concomitant therapy: Rescue medication was a sulfonylurea (specific type/dose to be chosen by the investigator), which was added to the double-blind study medication regimen, or another medication if it was contraindicated or inappropriate.

86.2.3. Alogliptin (N = 225)

Two groups: Alogliptin 25mg once a day (n=112) and alogliptin 12.5mg twice a day (n=113). Concomitant therapy: Rescue medication was a sulfonylurea (specific type/dose to be chosen by the investigator), which was added to the double-blind study medication regimen, or another medication if it was contraindicated or inappropriate.

86.2.4. Placebo (N = 109)

Matching placebo. Concomitant therapy: Rescue medication was a sulfonylurea (specific type/dose to be chosen by the investigator), which was added to the double-blind study medication regimen, or another medication if it was contraindicated or inappropriate.

86.3. Characteristics

86.3.1. Arm-level characteristics

86.3.1. Arm-le	evel characteristic	CS		
Characteristic	Alogliptin + Metformin (N = 225)	Metformin (N = 225)	Alogliptin (N = 225)	Placebo (N = 109)
% Male	n = 110; % = 49	n = 98 ; % = 44	n = 111 ; % = 49	n = 55 ; % = 51
Sample size			10	0.1
Mean age (SD) (years)	54.1 (11)	53.6 (10.8)	53.2 (9.6)	53.1 (9.6)
Mean (SD)				
Ethnicity	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
Sample size	10.0/.00			
Asian Sample size	n = 46 ; % = 20	n = 39 ; % = 17	n = 38 ; % = 17	n = 20 ; % = 18
Black or African American	n = 11 : % = 5			
Sample size	, //	n = 12; % = 5	n = 6; % = 3	n = 8; % = 7
White	n = 154 ; % = 68	n = 164 ; % = 73	n = 167 ; % = 74	n = 76 ; % = 70
Sample size		73	7-4	70
Other	n = 14; % = 6	n = 10; % = 4	n = 14 ; % = 6	n = 5 ; % = 5
Sample size				
Comorbidities Sample size	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
Presence of frailty	n = NR ; % = NR			
Sample size	11 1417, 70 1417	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
Time since type 2 diabetes diagnosis (years)	4.2 (4.9)	3.6 (4.3)	3.7 (4)	4.3 (4.78)
Mean (SD)				
HbA1c (%)	NR (NR)	NR (NR)	NR (NR)	NR (NR)
Mean (SD)				
Blood pressure (mmHg)	NA (NA)	NA (NA)	NA (NA)	NA (NA)
Mean (SD)				
Systolic blood pressure	126.2 (12.85)	126.6 (12.08)	126.4 (12.82)	127.6 (12.7)
Mean (SD)				

Characteristic	Alogliptin + Metformin (N = 225)	Metformin (N = 225)	Alogliptin (N = 225)	Placebo (N = 109)
Diastolic blood pressure	77 (7.7)	75.2 (7.94)	77.6 (7.8)	79.1 (7.36)
Mean (SD)				
Heart rate	NR (NR)	NR (NR)	NR (NR)	NR (NR)
Mean (SD)				
Smoking status	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
Sample size		INIX	INFX	- INIX
Alcohol consumption	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
Sample size		IVIX	IVIX	- IVIX
Presence of severe mental illness	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
Sample size				
People with significant cognitive impairment	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
Sample size				
People with a learning disability	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
Sample size				
Weight	NR (NR)	NR (NR)	NR (NR)	NR (NR)
Mean (SD)	24 (5.4)			
BMI (kg/m2) Mean (SD)	31 (5.4)	30.4 (4.9)	30.6 (5.2)	31.2 (5.3)
Number of people with obesity	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
Sample size				
Cholesterol and lipid levels (mg/dL)	NA (NA)	NA (NA)	NA (NA)	NA (NA)
Mean (SD)				
Total cholesterol	195.3 (45)	198.8 (42.3)	197.9 (46.5)	196.5 (39.7)
Mean (SD)				
HDL cholesterol	44.7 (10.4)	46.2 (11.2)	45.3 (11.5)	44.2 (9.2)
Mean (SD)				

Characteristic	Alogliptin + Metformin (N = 225)	Metformin (N = 225)	Alogliptin (N = 225)	Placebo (N = 109)
LDL cholesterol	112.5 (31.2)	116.9 (35.8)	112.8 (36.9)	115 (35)
Mean (SD)				
Triglycerides	193.9 (148)	186.1 (115.2)	207.6 (164.9)	197.8 (148.4)
Mean (SD)				(1-101)
Albumin creatinine ratio	NR (NR)	NR (NR)	NR (NR)	NR (NR)
Mean (SD)	`			
eGFR (mL/min/1.73m2)	NR (NR)	NR (NR)	NR (NR)	NR (NR)
Mean (SD)				
Other antidiabetic medication used	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
Sample size				
Blood pressure-lowering medication used	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
Sample size				
Statins/lipid-lowering medication used	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
Sample size				
Other treatment being received	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
Sample size				

87. Roden, 2005

Bibliographic Reference

Roden, M; Laakso, M; Johns, D; Widel, M; Urquhart, R; Richardson, C; Mariz, S; Tan, M H; Long-term effects of pioglitazone and metformin on insulin sensitivity in patients with Type 2 diabetes mellitus.; Diabetic medicine: a journal of the British Diabetic Association; 2005; vol. 22 (no. 8); 1101-6

	tudy details
Secondary publication of another included study- see primary study for details	No additional information.
Other publications associated with this study included in review	No additional information.
Trial name / registration number	No additional information.
Study type	Randomised controlled trial (RCT)
Study location	Europe (study A and B) and Canada (study B only).
Study setting	Outpatient follow-up.
Study dates	No additional information.
Sources of funding	Sponsored by Eli Lilly and Company and Takeda Europe R & D.
Inclusion criteria	Male and females with type 2 diabetes; aged 35-75 years; HbA1c between 7.5-11%.
Exclusion criteria	Previous treatment with metformin, pioglitazone or other thiazolidinediones.
Recruitment / selection of participants	No additional information.
Intervention(s)	Study A intervention: Pioglitazone N=597 Pioglitazone initially 30mg once a day titrated up to a maximum of 45mg per day for 52 weeks. Concomitant therapy: No additional information.

	Study B intervention:
	Pioglitazone (background sulfonylurea) N=316
	Pioglitazone initially 30mg once a day titrated up to a maximum of 45mg per day for 52 weeks. In addition people in this arm received a regular sulfonylurea that remained unchanged throughout the study. Concomitant therapy: No additional information.
Cointervention	For people in study A - none.
	For people in study B - all people received a sulfonylurea throughout the study.
Strata 1: People with type 2 diabetes mellitus and heart failure	Not stated/unclear
Strata 2: People with atherosclerotic cardiovascular diseases	Not stated/unclear
Strata 3: People with type 2 diabetes mellitus and chronic kidney disease	Not stated/unclear
Strata 4: People with type 2 diabetes mellitus and high cardiovascular risk	Not stated/unclear
Subgroup 1: People with frailty	Not stated/unclear
Subgroup 2: Onset of type 2 diabetes mellitus	Not stated/unclear
Subgroup 3: People with non-alcoholic	Not stated/unclear

fatty liver disease	
Subgroup 4: People with obesity	Not stated/unclear
Subgroup 5: eGFR category at baseline	Not stated/unclear
Subgroup 6: Albuminuria category at baseline	Not stated/unclear
Sensitivity analysis category: Enrichment trial status	5) All treatment naïve
Population subgroups	No additional information.
Comparator	Study A comparator:
	Metformin N=597 Metformin initially 850mg once a day titrated up to a maximum of 2550mg per day for 52 weeks. Concomitant therapy: No additional information. Study B comparator:
	Metformin N=597
	Metformin initially 850mg once a day titrated up to a maximum of 2550mg per day for 52 weeks. Concomitant therapy: No additional information.
Number of participants	1830
Duration of follow-up	52 weeks.
Indirectness	No additional information.
Method of analysis	Not stated/unclear
Additional comments	No additional information.

87.2.1. **Pioglitazone (N = 597)**

Pioglitazone initially 30mg once a day titrated up to a maximum of 45mg per day for 52 weeks. Concomitant therapy: No additional information.

87.2.2. Metformin (N = 597)

Metformin initially 850mg once a day titrated up to a maximum of 2550mg per day for 52 weeks. Concomitant therapy: No additional information.

87.2.3. Pioglitazone (background sulfonylurea) (N = 316)

Pioglitazone initially 30mg once a day titrated up to a maximum of 45mg per day for 52 weeks. In addition people in this arm received a regular sulfonylurea that remained unchanged throughout the study. Concomitant therapy: No additional information.

87.2.4. Metformin (background sulfonylurea) (N = 320)

Metformin initially 850mg once a day titrated up to a maximum of 2550mg per day for 52 weeks. In addition people in this arm received a regular sulfonylurea that remained unchanged throughout the study. Concomitant therapy: No additional information.

87.3. Characteristics

Characteristic	Pioglitazone (N = 597)	Metformin (N = 597)	Pioglitazone (background sulfonylurea) (N = 316)	Metformin (background sulfonylurea) (N = 320)
% Male Sample size	n = NR ; % = 52.6	n = NR ; % = 57.8	n = NR ; % = 53.6	n = NR ; % = 54.7
Mean age (SD) (years) Mean (SD)	57 (9.4)	56 (9.3)	60 (8.8)	60 (8)
Ethnicity Sample size	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR

Characteristic	Pioglitazone (N = 597)	Metformin (N = 597)	Pioglitazone (background sulfonylurea) (N = 316)	Metformin (background sulfonylurea) (N = 320)
Comorbidities	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
Sample size		- INIX		
Presence of frailty	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
Sample size				
Time since type 2 diabetes diagnosis (years)	3.4 (4.3)	3.1 (3.8)	7 (5.6)	7.1 (5.6)
Mean (SD)				
HbA1c (%)	8.7 (1)	8.7 (1)	8.8 (0.98)	8.8 (0.97)
Mean (SD)				
Smoking status Sample size	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
Alcohol	n = NR ; % =			
consumption	NR	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
Sample size				
Presence of severe mental illness	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
Sample size				
People with significant cognitive impairment	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
Sample size				
People with a learning disability	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
Sample size				
Number of people with obesity	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
Sample size				
Other antidiabetic medication used	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
Sample size				

Characteristic	Pioglitazone (N = 597)	Metformin (N = 597)	Pioglitazone (background sulfonylurea) (N = 316)	Metformin (background sulfonylurea) (N = 320)
Blood pressure- lowering medication used Sample size	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
Statins/lipid- lowering medication used Sample size	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
Other treatment being received Sample size	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR

88. Roden, 2015

Bibliographic Reference

Roden, Michael; Merker, Ludwig; Christiansen, Anita Vedel; Roux, Flavien; Salsali, Afshin; Kim, Gabriel; Stella, Peter; Woerle, Hans J; Broedl, Uli C; Safety, tolerability and effects on cardiometabolic risk factors of empagliflozin monotherapy in drug-naive patients with type 2 diabetes: a double-blind extension of a Phase III randomized controlled trial.; Cardiovascular diabetology; 2015; vol. 14; 154

ludy details			
NA			
Roden, M.; Weng, J.; Eilbracht, J.; Delafont, B.; Kim, G.; Woerle, H. J.; Broedl, U. C.; Empagliflozin monotherapy with sitagliptin as an active comparator in patients with type 2 diabetes: A randomised, double-blind, placebo-controlled, phase 3 trial; Lancet Diabetes Endocrinol; 2013; vol. 1 no. 3); 208-219			
EMPA-REG EXTEND MONO/NCT01289990			
Randomised controlled trial (RCT)			
International (124 academic medical centres, hospitals, or private practices in 9 countries: Belgium, Canada, China, Germany, India, Ireland, Japan, Switzerland, and USA).			
Community			
02/2011 to 05/2013			
Funded by Boehringer Ingelheim and Eli Lilly & Co.			
 Aged≥18 years (≥20 years in Japan) Diagnosis of type 2 diabetes Drug-naive (no oral or injectable anti-diabetes therapy for ≥12 weeks prior to randomisation) Insufficient glycaemic control on diet and exercise HbA1c 7-10% inclusive (7-9% inclusive in Germany) BMI≤45 kg/m2 			
 Uncontrolled hyperglycaemia (glucose concentration >13.3 mmol/l following overnight fast, confirmed by second measurement) 			

- Acute coronary syndrome (non-STEMI, STEMI and unstable angina pectoris), stroke or transient ischaemic attach within 3 months prior to informed consent
- Indication of liver disease, either ALT, AST, or alkaline phosphatase above 3 x upper normal limit
- Impaired renal function (eGFR<50 ml/min)
- Bariatric surgery within the past two years or other gastrointestinal surgeries
- Medical history of cancer
- Contraindications to sitagliptin
- Blood dyscrasias or any disorders causing haemolysis or unstable red blood cell
- Treatment with any anti-diabetes drug within 12 weeks prior to randomisation
- Treatment with anti-obesity drugs or any other treatment leading to unstable body weight
- Current treatment with systemic steroids or change in dosage of thyroid hormones within 6 weeks prior to informed consent or any other uncontrolled endocrine disorder except T2DM
- Pre-menopausal women who are nursing or pregnant or are of child-bearing potential and not practicing an acceptable method of birth control
- Alcohol or drug abuse
- Intake of an investigational drug in another trial within 30 days prior to intake of study medication in this trial
- Any other clinical condition that would jeopardize patients safety while participating in this clinical trial

Recruitment / selection of participants

899 participants were originally recruited for 24 week RCT (NCT01177813) examining interventions. Participants in this extension trial were included if they had completed the original 24 week trial, were still eligible according to original criteria, and consented to continue for a further 52 weeks (total 76 weeks treatment). Participants remained in groups assigned in original trial, continued to receive diet and exercise counselling based on local recommendations. Participants who received rescue medication in original 24 week trial and were still on it continued on it for duration of this extension trial. Rescue medication (choice at investigator's discretion but GLP-1 RA and DPP-4 inhibitor not allowed) could be initiated in extension period if there was confirmed plasma glucose level >10 mmol/L after overnight fast or HbA1c>8%. Participants with uncontrolled hyper- or hypoglycaemia discontinued trial.

Note that original trial also included a non-randomised open-label empagliflozin 25 mg arm (n=87; data not extracted).

Intervention(s)

- Empagliflozin 25 mg once daily
- Empagliflozin 10 mg once daily

All participants received 3 tablets for 76 weeks: Empagliflozin 25 mg or 10 mg once daily plus matching placebo tablets for the other active arms (placebo for sitagliptin, placebo for other empagliflozin dose).

Cointervention Placebo in triple-dummy design with matching placebo tables for other active arms.

Strata 1: People with type 2 diabetes mellitus and heart failure	Not stated/unclear
Strata 2:	People without other cardiovascular diseases
People with	
atherosclerotic cardiovascular diseases	Exclusion criteria: Acute coronary syndrome (non-STEMI, STEMI and unstable angina pectoris), stroke or transient ischaemic attach within 3 months prior to informed consent.
Strata 3:	Not stated/unclear
People with type 2 diabetes mellitus and chronic kidney	
disease	
Strata 4: People with type 2	Not stated/unclear
diabetes mellitus and high cardiovascular risk	
Subgroup 1: People with frailty	Not stated/unclear
Subgroup 2: Onset of type 2 diabetes mellitus	Not stated/unclear
Subgroup 3: People with non-alcoholic fatty liver disease	Not stated/unclear
Subgroup 4: People with obesity	Not stated/unclear
Subgroup 5:	eGFR ≥30mL/min/1.73m2
eGFR category at baseline	Exclusion criteria: Impaired renal function (eGFR<50 ml/min)
Subgroup 6: Albuminuria category at baseline	Not stated/unclear
Sensitivity analysis	7) Mixed population

category: Inclusion criteria: Drug-naive (no oral or injectable anti-diabetes therapy for ≥12 weeks prior to randomisation) Population subgroups • Sitagliptin 100 mg once daily Comparator • Sitagliptin 100 mg once daily Placebo All participants received 3 tablets for 76 weeks. For sitagliptin, sitagliptin 100 mg tablet, placebo for empagliflozin 25 mg and placebo for empagliflozin 10 mg. For placebo arm, participants received 3 placebo tablets (one for each of the 3 active arms: empagliflozin 25 mg, empagliflozin 10 mg, sitagliptin 100 mg). Number of participants N=899 in original 24 week trial N=615 in 52 week extension trial (completers of original trial, still meets original eligibility criteria, and consented to participate in extension period) Duration of follow-up None Method of analysis Modified ITT Full analysis set used for HbA1c, FPG, weight, and blood pressure included all randomised participants who received one or more study drug dose and had baseline HbA1c measurement in initial 24 week study. Safety analysis used all randomised participants who received one or more study drug dose.		
Comparator Sitagliptin 100 mg once daily Placebo All participants received 3 tablets for 76 weeks. For sitagliptin, sitagliptin 100 mg tablet, placebo for empagliflozin 25 mg and placebo for empagliflozin 10 mg. For placebo arm, participants received 3 placebo tablets (one for each of the 3 active arms: empagliflozin 25 mg, empagliflozin 10 mg, sitagliptin 100 mg). Number of participants N=899 in original 24 week trial N=615 in 52 week extension trial (completers of original trial, still meets original eligibility criteria, and consented to participate in extension period) Duration of follow-up Indirectness None Method of analysis Modified ITT Full analysis set used for HbA1c, FPG, weight, and blood pressure included all randomised participants who received one or more study drug dose and had baseline HbA1c measurement in initial 24 week study. Safety analysis used all randomised participants who received one or more study drug dose.	Enrichment	•
All participants received 3 tablets for 76 weeks. For sitagliptin, sitagliptin 100 mg tablet, placebo for empagliflozin 25 mg and placebo for empagliflozin 10 mg. For placebo arm, participants received 3 placebo tablets (one for each of the 3 active arms: empagliflozin 25 mg, empagliflozin 10 mg, sitagliptin 100 mg). Number of participants N=899 in original 24 week trial N=615 in 52 week extension trial (completers of original trial, still meets original eligibility criteria, and consented to participate in extension period) Duration of follow-up Indirectness None Method of analysis Full analysis set used for HbA1c, FPG, weight, and blood pressure included all randomised participants who received one or more study drug dose and had baseline HbA1c measurement in initial 24 week study. Safety analysis used all randomised participants who received one or more study drug dose.	•	
participants N=615 in 52 week extension trial (completers of original trial, still meets original eligibility criteria, and consented to participate in extension period) To weeks None Method of analysis Modified ITT Full analysis set used for HbA1c, FPG, weight, and blood pressure included all randomised participants who received one or more study drug dose and had baseline HbA1c measurement in initial 24 week study. Safety analysis used all randomised participants who received one or more study drug dose.	Comparator	 Placebo All participants received 3 tablets for 76 weeks. For sitagliptin, sitagliptin 100 mg tablet, placebo for empagliflozin 25 mg and placebo for empagliflozin 10 mg. For placebo arm, participants received 3 placebo tablets (one for each of the 3 active arms: empagliflozin 25 mg,
Indirectness None Method of analysis Full analysis set used for HbA1c, FPG, weight, and blood pressure included all randomised participants who received one or more study drug dose and had baseline HbA1c measurement in initial 24 week study. Safety analysis used all randomised participants who received one or more study drug dose.		N=615 in 52 week extension trial (completers of original trial, still meets
Method of analysis Full analysis set used for HbA1c, FPG, weight, and blood pressure included all randomised participants who received one or more study drug dose and had baseline HbA1c measurement in initial 24 week study. Safety analysis used all randomised participants who received one or more study drug dose.		76 weeks
Full analysis set used for HbA1c, FPG, weight, and blood pressure included all randomised participants who received one or more study drug dose and had baseline HbA1c measurement in initial 24 week study. Safety analysis used all randomised participants who received one or more study drug dose.	Indirectness	None
Additional		Full analysis set used for HbA1c, FPG, weight, and blood pressure included all randomised participants who received one or more study drug dose and had baseline HbA1c measurement in initial 24 week study. Safety analysis used all randomised participants who received one or
comments	Additional comments	

88.2.1. Empagliflozin 25 mg once daily (N = 224)

Oral empagliflozin tablets 25 mg once daily for 24 weeks. Those consented to extension period continued for a further 52 weeks

88.2.2. Empagliflozin 10 mg once daily (N = 224)

Oral empagliflozin tablets 10 mg once daily for 24 weeks. Those consented to extension period continued for a further 52 weeks

88.2.3. Sitagliptin 100 mg once daily (N = 223)

Oral sitagliptin tablets 100 mg once daily for 24 weeks. Those consented to extension period continued for a further 52 weeks

88.2.4. Placebo (N = 228)

Matching placebo for 24 weeks. Those consented to extension period continued for a further 52 weeks

88.3. Characteristics

88.3.1. Arm-level characteristics

00.3.1. F	Arm-level characte	HISTICS		
Characteristic	Empagliflozin 25 mg once daily (N = 224)	Empagliflozin 10 mg once daily (N = 224)	Sitagliptin 100 mg once daily (N = 223)	
% Male Sample size	n = 145; % = 64.7	n = 142 ; % = 63.4	n = 141 ; % = 63.2	n = 123 ; % = 53.9
	50.0 (44.0)			
Mean age (SD) Mean (SD)	53.8 (11.6)	56.2 (11.6)	55.1 (9.9)	54.9 (10.9)
Ethnicity	n = NA ; % = NA			
Sample size	11 - 147 (, 70 - 147 (n = NA ; % = NA	n = NA ; % = NA	n = NA ; % = NA
Asian	n = 144; % = 64.3			
Sample size	,	n = 143 ; % = 63.8	n = 143 ; % = 64.1	n = 146 ; % = 64
Black/African- American	n = 7; % = 3.1	n = 3; % = 1.3	n = 3; % = 1.3	n = 6; % = 2.6
Sample size				
Other	n = 0; % = 0	n = 1; % = 0.1	n = 1; % = 0.4	n = 0; % = 0
Sample size				- 0
White Sample size	n = 73 ; % = 32.6	n = 77 ; % = 34.4	n = 76 ; % = 34.1	n = 76; % = 33.3
Comorbidities	NR			
oomorbianes	IVIX	NR	NR	NR
Custom value				
Presence of frailty	NR	NR	NR	NR
Custom value				
Time since type 2 diabetes diagnosis (years)	n = NA ; % = NA	n = NA ; % = NA	n = NA ; % = NA	n = NA ; % = NA
Sample size				

Characteristic	Empagliflozin 25 mg once daily (N = 224)	Empagliflozin 10 mg once daily (N = 224)	Sitagliptin 100 mg once daily (N = 223)	
< or equal to 1 year Sample size	n = 91; % = 40.6	n = 87 ; % = 38.8	n = 93 ; % = 41.7	n = 72; % = 31.6
•	00 0/ 07 4			
>1 to 5 years Sample size	n = 83 ; % = 37.1	n = 92 ; % = 41.1	n = 86 ; % = 38.6	n = 104 ; % = 45.6
>5 to 10 years	n = 37 ; % = 16.5	n = 29 ; % = 12.9	n = 32 ; % = 14.3	n = 33; % = 14.5
Sample size			14.3	- 14.5
More than 10 years Sample size	n = 13; % = 5.8	n = 16; % = 7.1	n = 12 ; % = 5.4	n = 19; % = 8.3
	7.86 (0.85)			
HbA1c (%) Mean (SD)	7.00 (0.03)	7.87 (0.88)	7.85 (0.79)	7.91 (0.78)
Blood pressure (mmHg)	NA (NA)	NA (NA)	NA (NA)	NA (NA)
Mean (SD)				
Systolic blood pressure	129.9 (17.5)	133 (16.6)	132.5 (15.8)	130.4 (16.3)
Mean (SD)				
Diastolic blood pressure	78.3 (9.4)	79.2 (9.6)	80.1 (10)	78.9 (9.6)
Mean (SD)				
Smoking status Custom value	NR	NR	NR	NR
Alcohol consumption	NR	NR	NR	NR
Custom value				
Presence of severe mental illness	NR	NR	NR	NR
Custom value				
People with significant cognitive impairment	NR	NR	empty data	NR
Custom value				

Characteristic	Empagliflozin 25 mg once daily (N = 224)	Empagliflozin 10 mg once daily (N = 224)	Sitagliptin 100 mg once daily (N = 223)	Placebo (N = 228)
People with a learning disability	NR	NR	NR	NR
Custom value				
Weight (kg)	77.8 (18)	78.4 (18.7)	79.3 (20.4)	78.2 (19.9)
Mean (SD)	00.0 (5.5)			(19.9)
BMI Nominal	28.2 (5.5)	28.3 (5.5)	28.2 (5.2)	28.7 (6.2)
Number of people	NR			
with obesity		NR	NR	NR
Custom value				
Cholesterol and lipid levels (mmol/L)	NA (NA)	NA (NA)	NA (NA)	NA (NA)
Mean (SE)				
Total cholesterol	5 (0.08)	5 (0.08)	4.95 (0.07)	5.03 (0.08)
Mean (SE) HDL-cholesterol	1.25 (0.02)			,
Mean (SE)	1.23 (0.02)	1.24 (0.02)	1.26 (0.02)	1.26 (0.02)
LDL-cholesterol	2.75 (0.07)	2.86 (0.07)	2.74 (0.05)	2.9 (0.06)
Mean (SE)	2.27 (0.2)			
Triglycerides Mean (SE)	2.37 (0.2)	2.08 (0.12)	2.2 (0.13)	2.01 (0.09)
eGFR (mL/min/1.73m2) Mean (SD)	87.6 (18.3)	87.7 (19.2)	87.6 (17.3)	86.8 (17.9)
Other antidiabetic medication used	NR	NR	NR	NR
Nominal				
Blood pressure- lowering medication used	NR	NR	NR	NR
Nominal				

Characteristic	Empagliflozin 25 mg once daily (N = 224)	Empagliflozin 10 mg once daily (N = 224)	Sitagliptin 100 mg once daily (N = 223)	
Statins/lipid-lowering medication used Custom value	NR	NR	NR	NR
Other treatment being received Custom value	NR	NR	NR	NR

All data are reported for the participants in the original 24 week trial. Article reports baseline characteristics for participants who continued trial were comparable with overall 24 week trial population.

89. Rosenstock, 2009

Bibliographic Reference

Rosenstock, J.; Aguilar-Salinas, C.; Klein, E.; Nepal, S.; List, J.; Chen, R.; Effect of saxagliptin monotherapy in treatment-naïve patients with type 2 diabetes; Curr Med Res Opin; 2009; vol. 25 (no. 10); 2401-11

	tudy details
Secondary publication of another included study- see primary study for details	No additional information
Other publications associated with this study included in review	No additional information
Trial name / registration number	CV181-011
Study type	Randomised controlled trial (RCT)
Study location	International
Study setting	No additional information
Study dates	No additional information
Sources of funding	Funded by Bristol Myers-Squibb and AstraZeneca
Inclusion criteria	Aged 18-77 years Type 2 diabetes inadequately controlled (HbA1c >7-10%) with exercise and diet alone Treatment naïve (defined as never receiving medical treatment for diabetes, insulin or other oral antihyperglycemic medication for >6 months since diagnosis, or antihyperglycemic medication for >3 consecutive, or >7 non-consecutive days in the past 2 months Fasting C-peptide ≥1 ng/mL BMI ≤40 kg/m2

Exclusion criteria	Symptoms of poorly controlled diabetes
Citteria	History of diabetic ketoacidosis or hyperosmolar nonketoic coma
	Cardiovascular event within 6 months or New York Heart Association stage III/IV congestive heart failure and/or known left ventricular ejection fraction ≤40%
	Significant renal, liver or psychiatric history
	History of drug or alcohol abuse within the past year
	Those who were immunocompromised
	Active liver disease or clinically significant comorbidities on screening tests of hepatic, renal, endocrine, metabolic or haematologic function
Recruitment / selection of participants	No additional information
Intervention(s)	Participants were randomised to one of three saxagliptin doses - 2.5, 5 or 10 mg per day. Participants who demonstrated a lack of adequate glucose control during the 24-week treatment period were eligible for the addition of open-label metformin as rescue therapy. The glycemic rescue criteria were fasting plasma glucose >240 mg/dL (13.3 mmol/L) at weeks 4 and 6, >220 mg/dL (12.2 mmol/L) at week 8, or >200 mg/dL (11.1 mmol/L) at weeks 12, 16, 20, and 24
	three study arms containing different doses of saxagliptin were combined for this review
Strata 1: People with	People without heart failure
type 2 diabetes mellitus and heart failure	Exclusion criteria: NYHA class III/IV or ejection fraction <40%
Strata 2: People with atherosclerotic cardiovascular diseases	Not stated/unclear
Strata 3: People with type 2 diabetes mellitus and	Not stated/unclear

chronic kidney disease	
Strata 4: People with type 2 diabetes mellitus and high cardiovascular risk	People at higher risk of developing cardiovascular disease
Subgroup 1: People with frailty	Not stated/unclear
Subgroup 2: Onset of type 2 diabetes mellitus	Not stated/unclear
Subgroup 3: People with non-alcoholic fatty liver disease	Not stated/unclear
Subgroup 4: People with obesity	Not stated/unclear
Subgroup 5: eGFR category at baseline	Not stated/unclear
Subgroup 6: Albuminuria category at baseline	Not stated/unclear
Sensitivity analysis category: Enrichment trial status	6) No response criteria
Population subgroups	No additional information
Comparator	Participants randomised to the placebo arm received a placebo tablet once daily. Participants who demonstrated a lack of adequate glucose control during the 24-week treatment period were eligible for the addition of openlabel metformin as rescue therapy. The glycemic rescue criteria were fasting plasma glucose >240 mg/dL (13.3 mmol/L) at weeks 4 and 6, >220 mg/dL (12.2 mmol/L) at week 8, or >200 mg/dL (11.1 mmol/L) at weeks 12, 16, 20, and 24
Number of participants	403 randomised

	306 received saxagliptin, 210 completed
	102 received 2.5 mg saxagliptin
	106 received 5 mg saxagliptin
	98 received 10 mg saxagliptin
	95 received placebo, 55 completed
	Study also includes a group of 66 participants who received non-randomised, open-label 10 mg saxagliptin - not included in this analysis
Duration of follow-up	24 weeks
Indirectness	None
Method of analysis	ITT
Additional comments	No additional information

89.2.1. Saxagliptin (N = 306)

2.5, 5 or 10 mg saxagliptin per day *three study arms combined for this review*

89.2.2. Placebo (N = 95)

Placebo once daily

89.3. Characteristics

Characteristic	Saxagliptin (N = 306)	Placebo (N = 95)
% Male	n = 157; % = 51	n = 47 ; % = 50
Sample size		
Mean age (SD)	53 (empty data)	53.91 (11.27)
Mean (SD)		
Ethnicity	n = NA ; % = NA	n = NA ; % = NA

Characteristic	Saxagliptin (N = 306)	Placebo (N = 95)
Sample size		
White	n = 262 ; % = 86	n = 79 ; % = 83
Sample size	40 - 0/ 5	
Black/African American	n = 16; % = 5	n = 6; % = 6
Sample size		
Asian	n = 15; % = 5	n = 3; % = 3
Sample size		
Other	n = 13; % = 4	n = 7; % = 7
Sample size	ND	
Comorbidities	NR	NR
Nominal		
Presence of frailty	NR	NR
Nominal		
Time since type 2 diabetes diagnosis (years)	2.6 (3.3)	2.3 (2.7)
Mean (SD)		
HbA1c (%)	7.9 (1)	7.9 (0.9)
Mean (SD)		
Blood pressure	NR	NR
Nominal		
Heart rate	NR	NR
Nominal		
Smoking status	NR	NR
Nominal		
Alcohol consumption	NR	NR
Nominal		1413
Presence of severe mental illness	NR	ND
Nominal		NR
People with significant cognitive impairment	NR	NR
Nominal		INIX

Characteristic	Saxagliptin (N = 306)	Placebo (N = 95)
People with a learning disability	NR	NR
Nominal		
Weight (kg)	90.78 (18.11)	86.56 (16.9)
Mean (SD)		
BMI (kg/m²)	31.99 (4.68)	30.93 (4.26)
Mean (SD)		
Number of people with obesity	NR	NR
Nominal		
Cholesterol and lipid levels	NR	NR
Nominal		
Albumin creatinine ratio	NR	NR
Nominal		
eGFR (mL/min/1.73m2)	NR	NR
Nominal		
Other antidiabetic medication used	NR	NR
Nominal		
Blood pressure-lowering medication used	NR	NR
Nominal		
Statins/lipid-lowering medication used	NR	NR
Nominal		
Other treatment being received	NR	NR
Nominal		

90. Rosenstock, 2016

Bibliographic Reference

Rosenstock, J.; Chuck, L.; Gonzalez-Ortiz, M.; Merton, K.; Craig, J.; Capuano, G.; Qiu, R.; Initial combination therapy with canagliflozin plus metformin versus each component as monotherapy for drug-naive type 2 diabetes; Diabetes Care; 2016; vol. 39 (no. 3); 353-362

30.1. 3	tudy details
Secondary publication of another included study- see primary study for details	No additional information.
Other publications associated with this study included in review	No additional information.
Trial name / registration number	NCT01809327.
Study type	Randomised controlled trial (RCT)
Study location	Multicentre trial (12 countries).
Study setting	Outpatient follow-up.
Study dates	16th May 2013 to 1st December 2014.
Sources of funding	Supported by Janssen Research & Development LLC. Researchers received grants from multiple pharmacological companies.
Inclusion criteria	18-75 years of age; drug naïve type 2 diabetes (not on antihyperglycaemic therapy or off for at least 12 weeks before screening) that was inadequately controlled with diet and exercise (HbA1c 7.5-12.0% at screening).
Exclusion criteria	History of type 1 diabetes; repeated fasting self-monitored blood glucose >300mg/dL; myocardial infarction, unstable angina, revascularisation procedure or cerebrovascular accident no less than 12 weeks before screening; NYHA III-IV cardiac disease; uncontrolled hypertension; eGFR <60mL/min/1.73m2 or serum creatinine more than or equal to 1.4mg/dL for men and more than or equal to 1.3mg/dL for women; were taking any antihyperglycaemic therapy within 12 weeks before screening or during the placebo run-in period.
Recruitment / selection of participants	People were discontinued from the study if they had fasting plasma glucose values meeting prespecified glycaemic withdrawal criteria (FPG >270mg/dL after day 1 through week 6; >240mg/dL after week 6 through week 12; >200mg/dL after week 12 through week 26), serum creatinine

	greater than or equal to 1.5mg/dL for men or greater than or equal to 1.4mg/dL for women or eGFR <50mL/min/1.73m2.
Intervention(s)	Canagliflozin + metformin N=474
	Two groups: Canagliflozin 300mg once a day and metformin extended release initially 500mg increased up to 2000mg once a day or the maximum tolerated dose over 9 weeks (n=237) or canagliflozin 100mg once a day and metformin extended release initially 500mg increased up to 2000mg once a day or the maximum tolerated dose over 9 weeks (n=237).
	Concomitant therapy: No additional information.
Strata 1: People with type 2 diabetes mellitus and heart failure	People without heart failure
Strata 2: People with atherosclerotic cardiovascular diseases	People without other cardiovascular diseases
Strata 3: People with type 2 diabetes mellitus and chronic kidney disease	People without chronic kidney disease
Strata 4:	People at higher risk of developing cardiovascular disease
People with type 2 diabetes mellitus and high cardiovascular risk	Based on BMI and presence of diabetes.
Subgroup 1: People with frailty	Not stated/unclear
Subgroup 2: Onset of type 2 diabetes mellitus	Not stated/unclear
Subgroup 3: People with non-alcoholic	Not stated/unclear

fatty liver disease	
Subgroup 4: People with obesity	Not stated/unclear
Subgroup 5: eGFR category at baseline	eGFR ≥30mL/min/1.73m2
Subgroup 6: Albuminuria category at baseline	Not stated/unclear
Sensitivity analysis category: Enrichment trial status	7) Mixed population
Population subgroups	No additional information.
Comparator	Canagliflozin N=475
	Two groups: Canagliflozin 300mg once a day and matching placebo (n=238) or canagliflozin 100mg once a day and matching placebo (n=237).
	Concomitant therapy: No additional information.
	Metformin N=237
	Metformin extended release initially 500mg increased up to 2000mg once a day or the maximum tolerated dose over 9 weeks and matching placebo.
	Concomitant therapy: No additional information.
Number of participants	1186
Duration of follow-up	26 weeks.
Indirectness	No additional information.
Method of analysis	Modified ITT All people who were randomised and received at least one dose of the
	double-blind study drug.

Additional	No additional information.
comments	

90.2.1. Canagliflozin + metformin (N = 474)

Two groups: Canagliflozin 300mg once a day and metformin extended release initially 500mg increased up to 2000mg once a day or the maximum tolerated dose over 9 weeks (n=237) or canagliflozin 100mg once a day and metformin extended release initially 500mg increased up to 2000mg once a day or the maximum tolerated dose over 9 weeks (n=237). Concomitant therapy: No additional information.

90.2.2. Canagliflozin (N = 475)

Two groups: Canagliflozin 300mg once a day and matching placebo (n=238) or canagliflozin 100mg once a day and matching placebo (n=237). Concomitant therapy: No additional information.

90.2.3. Metformin (N = 237)

Metformin extended release initially 500mg increased up to 2000mg once a day or the maximum tolerated dose over 9 weeks and matching placebo. Concomitant therapy: No additional information.

90.3. Characteristics

Characteristic	Canagliflozin + metformin (N = 474)	Canagliflozin (N = 475)	Metformin (N = 237)
% Male Sample size	n = 263 ; % = 56	n = 230 ; % = 48	n = 116 ; % = 49
Mean age (SD) (years) Mean (SD)	54.8 (9.7)	54.9 (10.2)	55.2 (9.8)
Ethnicity Sample size	n = NA ; % = NA	n = NA ; % = NA	n = NA ; % = NA
White Sample size	n = 376 ; % = 79	n = 400 ; % = 84	n = 192 ; % = 81
Black/African American	n = 14; % = 3	n = 20 ; % = 4	n = 9; % = 4

Characteristic	Canagliflozin +	Canagliflozin (N	Metformin (N
- Thurword Total o	metformin (N = 474)	= 475)	= 237)
Sample size			
Asian	n = 11; % = 2	n = 8; % = 2	n = 9; % = 4
Sample size			
Other	n = 73 ; % = 15	n = 47 ; % = 10	n = 27 ; % = 11
Sample size	ND 0/ ND		
Comorbidities	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
Sample size	ND 0/ ND		
Presence of frailty Sample size	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
•	2 1 (2 6)		
Time since type 2 diabetes diagnosis (years)	3.1 (3.6)	3.4 (4.4)	3.3 (4.5)
Mean (SD)			
HbA1c (%)	8.9 (1.2)	8.8 (1.2)	8.8 (1.2)
Mean (SD)			
Blood pressure (mmHg)	NA (NA)	NA (NA)	NA (NA)
Mean (SD)			
Systolic blood pressure Mean (SD)	127.8 (11.9)	129.5 (11.6)	129.4 (12)
Diastolic blood pressure	78.3 (8)		
Diactono bioca procedio	7 0.0 (0)	78.8 (7.8)	78.3 (7.8)
Mean (SD)			
Heart rate	NR (NR)	NR (NR)	NR (NR)
Mean (SD)			
Smoking status	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
Sample size			TVI V
Alcohol consumption	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
Sample size	ND - 0/ ND		
Presence of severe mental illness	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
Sample size			

Characteristic	Canagliflozin + metformin (N = 474)	Canagliflozin (N = 475)	Metformin (N = 237)
People with significant cognitive impairment	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
Sample size			
People with a learning disability	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
Sample size			
Weight (kg)	89.9 (19.7)	91.6 (19.3)	92.1 (20.1)
Mean (SD)	00.4 (0)		
BMI (kg/m2)	32.4 (6)	32.5 (5.6)	33 (6)
Mean (SD) Number of people with	n = NR ; % = NR		
obesity	11 - INIX , 70 - INIX	n = NR ; % = NR	n = NR ; % = NR
Sample size Chalacteral and linid levels	NIA (NIA)		
Cholesterol and lipid levels (mg/dL)	NA (NA)	NA (NA)	NA (NA)
Mean (SD)			
HDL cholesterol	44 (10.6)	43.6 (10.6)	43.7 (10.6)
Mean (SD)	110 7 (20 4)		
LDL cholesterol Mean (SD)	118.7 (39.4)	119.3 (38.2)	115.5 (36.3)
Triglycerides	172.7 (95.3)		
	,	180.2 (114)	188.6 (126.6)
Mean (SD)			
Albumin creatinine ratio Mean (SD)	NR (NR)	NR (NR)	NR (NR)
eGFR (mL/min/1.73m2) (88 (10)		
ml/min/1.73 m2)	88 (19)	87.5 (18.7)	87 (19)
Mean (SD)			
Other antidiabetic medication used	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
Sample size			
Blood pressure-lowering medication used	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR

Characteristic	Canagliflozin + metformin (N = 474)	Canagliflozin (N = 475)	Metformin (N = 237)
Sample size			
Statins/lipid-lowering medication used Sample size	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
Other treatment being received Sample size	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR

91. Rosenstock, 2007

Bibliographic Reference

Rosenstock, J.; Kim, S. W.; Baron, M. A.; Camisasca, R. P.; Cressier, F.; Couturier, A.; Dejager, S.; Efficacy and tolerability of initial combination therapy with vildagliptin and pioglitazone compared with component monotherapy in patients with type 2 diabetes; Diabetes Obes Metab; 2007; vol. 9 (no. 2); 175-85

91.1. S	tudy details	
Secondary publication of another included study- see primary study for details	NA	
Other publications associated with this study included in review	None	
Trial name / registration number	NCT00101803	
Study type	Randomised controlled trial (RCT)	
Study location	International (145 sites in 8 countries: Czech Republic, India, Italy, Slovakia, South Korea, Taiwan, UK, USA)	
Study setting	Community	
Study dates	01/2005 to 01/2006	
Sources of funding	Funded by Novartis Pharmaceuticals Corporation.	
Inclusion criteria	 Aged 18-80 years Diagnosis of type 2 diabetes mellitus HbA1c level 7.5-11% inclusive at screening No pharmacological treatment ≥12-mo before screening No oral antidiabetic medication≥3 consecutive months at any time BMI 22-45 kg/m2 inclusive Fasting plasma glucose <15 mmol/l 	
Exclusion criteria	 History of type 1 or secondary forms of diabetes Acute metabolic diabetic complications Myocardial infarction, unstable angina or coronary artery bypass surgery within the previous 6 months Congestive heart failure Liver disease (e.g. cirrhosis or chronic active hepatitis) 	

 Any contraindications and warnings according to the country-specific label for pioglitazone Alanine aminotransferase or aspartate aminotransferase>2.5 times upper limit of normal (ULN) Direct bilirubin>1.3 times the ULN Serum creatinine levels>220mmol/I Clinically significant abnormal TSH or fasting triglycerides>7.9 mmol/I
Participants recruited from 145 centres in 8 countries. Treatment blinding maintained using double-dummy technique. All participants given glucose monitoring devices/supplies and instructed how to use them. All lab assessments conducted by central laboratory with standardised/validated procedures.
 Vildagliptin 100 mg + Pioglitazone 30 mg once daily Vildagliptin 50 mg + Pioglitazone 15mg once daily All drugs were oral and taken for 24 weeks. Pioglitazone doses based on recommended doses in prescribing information. Reports double dummy design but no further information reported, so assumed that participants in these arms received two active treatment pills only.
Placebo pills for comparator arms only (vildagliptin 100 mg once daily, pioglitazone 30 mg once daily)
People without heart failure Exclusion criteria: Congestive heart failure
People without other cardiovascular diseases Exclusion criteria: Myocardial infarction, unstable angina or coronary artery bypass surgery within the previous 6 months
Not stated/unclear
Not stated/unclear

Subgroup 1: People with frailty	Not stated/unclear
Subgroup 2: Onset of type 2 diabetes mellitus	Not stated/unclear
Subgroup 3: People with non-alcoholic fatty liver disease	Not stated/unclear
Subgroup 4: People with obesity	Not stated/unclear
Subgroup 5: eGFR category at baseline	Not stated/unclear
Subgroup 6: Albuminuria category at baseline	Not stated/unclear
Sensitivity analysis category: Enrichment trial status	7) Mixed population Inclusion criteria: no pharmacological treatment for at least 12 weeks prior to screening and no OAD for more than three consecutive months at any time in the past
Population subgroups	
Comparator	 Vildagliptin 100 mg once daily Pioglitazone 130 mg once daily Oral vildagliptin or oral pioglitazone 30 mg once daily for 24 weeks. Participants in both groups received placebo pills.
Number of participants	N=607
Duration of follow-up	24 weeks
Indirectness	None
Method of analysis	Modified ITT Primary efficacy analysis (HbA1c) and secondary efficacy analysis conducted on all randomised participants who received at least one study drug and had baseline and at least one postbaseline efficacy assessment (HbA1c).

91.2.1. Pioglitazone 30 mg once daily (N = 161)

Oral pioglitazone 30 mg once daily for 24 weeks.

91.2.2. Pioglitazone 15 mg + Vildagliptin 50 mg once daily (N = 144) Oral vildagliptin 50 mg + oral pioglitazone 15 mg once daily for 24 weeks.

91.2.3. Pioglitazone 30 mg + Vildagliptin 100 mg once daily (N = 148)

Oral vildagliptin 100 mg + oral pioglitazone 30 mg once daily for 24 weeks.

91.2.4. Vildagliptin 100 mg once daily (N = 154)

Oral vildagliptin 100 mg once daily for 24 weeks.

91.3. Characteristics

31.3.1.	Anni-level characteristics			
Characteristic	Pioglitazone 30 mg once daily (N = 161)	Pioglitazone 15 mg + Vildagliptin 50 mg once daily (N = 144)	Pioglitazone 30 mg + Vildagliptin 100 mg once daily (N = 148)	Vildagliptin 100 mg once daily (N = 154)
% Male Sample size	n = 103 ; % = 64	n = 84 ; % = 58.3	n = 86 ; % = 58.1	n = 98 ; % = 63.6
	EQ 4 (40 2)			
Mean age (SD) (years)	52.4 (10.3)	51 (11)	51 (11.3)	51.4 (10.8)
Mean (SD)				
Ethnicity	n = NA ; % = NA	n = NA ; % = NA	n = NA ; % = NA	n = NA ; % =
Sample size				NA
Asian	n = 69 ; % = 42.9	n = 68 ; % = 47.2	n = 66 ; % = 44.7	n = 70 ; % =
Sample size				45.5
Caucasian	n = 71 ; % = 44.1	n = 52 ; % = 36.1	n = 56 ; % = 37.8	n = 60 ; % =
Sample size				39
Hispanic or Latino	n = 14 ; % = 8.7	n = 15 ; % = 10.4	n = 23 ; % = 15.5	·
Sample size				11

Characteristic	Pioglitazone	Pioglitazone 15	Pioglitazone 30	Vildagliptin
	30 mg once daily (N = 161)	_	mg + Vildagliptin 100 mg once daily (N = 148)	
Other	n = 7; % = 4.3	n = 9; % = 6.3	n = 3; % = 2	n = 7; % = 4.5
Sample size				
Comorbidities	NR	NR	NR	NR
Custom value				
Presence of frailty	NR	NR	NR	NR
Custom value				
Time since type 2 diabetes diagnosis (years)	2.2 (3.3)	2 (3.2)	2 (3.1)	1.9 (3.1)
Mean (SD)				
HbA1c (%)	8.7 (1)	8.8 (0.9)	8.8 (1.1)	8.6 (1)
Mean (SD)				
Blood pressure	NR	NR	NR	NR
Custom value	ND			
Heart rate Custom value	NR	NR	NR	NR
	NID			
Smoking status Custom value	NR	NR	NR	NR
Alcohol	NR			
consumption	IVIX	NR	NR	NR
Nominal				
Presence of severe mental illness	NR	NR	NR	NR
Nominal				
People with significant cognitive impairment	NR	NR	NR	NR
People with a	NR			
learning disability	IVIX	NR	NR	NR

Characteristic	Pioglitazone 30 mg once daily (N = 161)	Pioglitazone 15 mg + Vildagliptin 50 mg once daily (N = 144)	Pioglitazone 30 mg + Vildagliptin 100 mg once daily (N = 148)	Vildagliptin 100 mg once daily (N = 154)
Nominal				
Weight	NR	NR	NR	NR
Nominal				
BMI (kg/m2)	28.9 (5.5)	29 (5.4)	29.6 (5.8)	29.4 (5.8)
Mean (SD)				
Number of people with obesity	NR	NR	NR	NR
Nominal				
Cholesterol and lipid levels (mmol/L)	NA (NA)	NA (NA)	NA (NA)	NA (NA)
Mean (SD)				
Total cholesterol	5.3 (0.1)	5.2 (0.1)	5.2 (0.1)	5.4 (0.1)
Mean (SD)				
HDL-cholesterol Mean (SD)	1.13 (0.03)	1.1 (0.03)	1.09 (0.02)	1.09 (0.03)
LDL-cholesterol	3.2 (0.1)			
	0.2 (0.1)	3.1 (0.1)	3.1 (0.1)	3.2 (0.1)
Mean (SD)				
Non-HDL- cholesterol	4.1 (0.1)	4.1 (0.1)	4.1 (0.1)	4.3 (0.1)
Mean (SD)				
Triglycerides Mean (SD)	2.3 (0.1)	2.5 (0.2)	2.4 (0.1)	2.5 (0.1)
Albumin creatinine	NR			
ratio	W	NR	NR	NR
Nominal				
eGFR (mL/min/1.73m2)	NR	NR	NR	NR
Nominal				
Other antidiabetic medication used	NR	NR	NR	NR

Characteristic	Pioglitazone 30 mg once daily (N = 161)		Pioglitazone 30 mg + Vildagliptin 100 mg once daily (N = 148)	Vildagliptin 100 mg once daily (N = 154)
Nominal				
Blood pressure- lowering medication used	NR	NR	NR	NR
Statins/lipid- lowering medication used Nominal	NR	NR	NR	NR
Other treatment being received Nominal	NR	NR	NR	NR

92. Ross, 2015

Bibliographic Reference

Ross, S. A.; Caballero, A. E.; Del Prato, S.; Gallwitz, B.; Lewis-D'Agostino, D.; Bailes, Z.; Thiemann, S.; Patel, S.; Woerle, H. J.; von Eynatten, M.; Initial combination of linagliptin and metformin compared with linagliptin monotherapy in patients with newly diagnosed type 2 diabetes and marked hyperglycaemia: A randomized, double-blind, active-controlled, parallel group, multinational clinical trial; Diabetes Obes Metab; 2015; vol. 17 (no. 2); 136-144

Secondary publication of another included study- see primary study for details	No additional information.
Other publications associated with this study included in review	No additional information.
Trial name / registration number	NCT01512979.
Study location	Multicentre trial (Canada, India, Israel, Malaysia, Mexico, the Philippines, Russia, Sri Lanka, Thailand, Ukraine and the USA).
Study setting	Outpatient follow-up.
Study dates	24th January 2012 to 15th April 2013.
Sources of funding	Sponsored by Boehringer Ingelheim.
Inclusion criteria	Aged at least 18 years; newly diagnosed with type 2 diabetes (<12 months before the first visit); HbA1c 8.5-12.0%; no glucose-lowering drug in the previous 12 weeks; body mass index no more than 45kg/m2.
Exclusion criteria	Acute coronary syndrome, stroke or TIA within the previous 3 months; hepatic disease (serum level of ALT, AST and/or ALP >3 times the upper limit of normal); kidney disease (creatinine clearance of <60mL/min as calculated by the Cockcroft-Gault equation); contraindication to metformin or linagliptin; premenopausal women who were nursing, pregnant or not practicing birth control; bariatric surgery within the previous 2 years; history of cancer or pancreatitis; treatment with anti-obesity drugs or systemic steroids.

Recruitment / selection of participants	No additional information.
Intervention(s)	Linagliptin + Metformin N=159
	Linagliptin 5mg once daily and metformin 1500-2000mg daily (delivered split over two daily administrations). Metformin was initiated at 1000mg daily for the first week, then increased to 1500mg for the second week, then increased to 200mg daily over weeks 3-6 if tolerated and if fasting plasma glucose was >6.1mmol/L.
	Concomitant therapy: People received diet and exercise counselling before the open-label placebo run in phase (for 1 week) and were reminded at each subsequent visit to follow the diet and exercise plan. Rescue medication (with any glucose-lowering therapy apart from a DPP-4 inhibitor, metformin or a GLP-1 receptor agonist) was available if FPG >15mmol/L during weeks 1-6, >13.3mmol/L during weeks 7-12, >11.1mmol/L during weeks 13-24, confirmed by at least 2 measurements on 2 different days.
Strata 1: People with type 2 diabetes mellitus and heart failure	Not stated/unclear
Strata 2: People with atherosclerotic cardiovascular diseases	Mixed population
Strata 3: People with type 2 diabetes mellitus and chronic kidney disease	People without chronic kidney disease
Strata 4:	People at higher risk of developing cardiovascular disease
People with type 2 diabetes mellitus and high cardiovascular risk	Based on BMI, number of people with previous macro- or microvascular disease, number of people requiring antihypertensives and with a mild renal impairment and presence of diabetes.
Subgroup 1: People with frailty	Not stated/unclear

Subgroup 2: Onset of type 2 diabetes mellitus	Not stated/unclear
Subgroup 3: People with non-alcoholic fatty liver disease	People without non-alcoholic fatty liver disease
Subgroup 4: People with obesity	Not stated/unclear
Subgroup 5: eGFR category at baseline	eGFR ≥30mL/min/1.73m2
Subgroup 6: Albuminuria category at baseline	Not stated/unclear
Sensitivity analysis category: Enrichment trial status	6) No response criteria
Population subgroups	No additional information.
Comparator	Linagliptin N=157 Linagliptin 5mg once daily. Concomitant therapy: People received diet and exercise counselling before the open-label placebo run in phase (for 1 week) and were reminded at each subsequent visit to follow the diet and exercise plan. Rescue medication (with any glucose-lowering therapy apart from a DPP-4 inhibitor, metformin or a GLP-1 receptor agonist) was available if FPG >15mmol/L during weeks 1-6, >13.3mmol/L during weeks 7-12, >11.1mmol/L during weeks 13-24, confirmed by at least 2 measurements on 2 different days.
Number of participants	316
Duration of follow-up	24 weeks.
Indirectness	No additional information.
Method of analysis	Per protocol Per protocol completers cohort - all people who received at least 1 dose of the study drug

Additional	No additional information.
comments	

92.2.1. Linagliptin + Metformin (N = 159)

Linagliptin 5mg once daily and metformin 1500-2000mg daily (delivered split over two daily administrations). Metformin was initiated at 1000mg daily for the first week, then increased to 1500mg for the second week, then increased to 200mg daily over weeks 3-6 if tolerated and if fasting plasma glucose was >6.1mmol/L. Concomitant therapy: People received diet and exercise counselling before the open-label placebo run in phase (for 1 week) and were reminded at each subsequent visit to follow the diet and exercise plan. Rescue medication (with any glucose-lowering therapy apart from a DPP-4 inhibitor, metformin or a GLP-1 receptor agonist) was available if FPG >15mmol/L during weeks 1-6, >13.3mmol/L during weeks 7-12, >11.1mmol/L during weeks 13-24, confirmed by at least 2 measurements on 2 different days.

92.2.2. Linagliptin (N = 157)

Linagliptin 5mg once daily. Concomitant therapy: People received diet and exercise counselling before the open-label placebo run in phase (for 1 week) and were reminded at each subsequent visit to follow the diet and exercise plan. Rescue medication (with any glucose-lowering therapy apart from a DPP-4 inhibitor, metformin or a GLP-1 receptor agonist) was available if FPG >15mmol/L during weeks 1-6, >13.3mmol/L during weeks 7-12, >11.1mmol/L during weeks 13-24, confirmed by at least 2 measurements on 2 different days.

92.3. Characteristics

Characteristic	Linagliptin + Metformin (N = 159)	Linagliptin (N = 157)
% Male	n = 69; % = 43	n = 77 ; % = 49
Sample size		
Mean age (SD) (years)	49 (10.9)	48.6 (11.2)
Mean (SD)		
Ethnicity	n = NA ; % = NA	n = NA ; % = NA
Sample size		
White	n = 97; % = 61	n = 85 ; % = 54.1
Sample size		

Characteristic	Linagliptin + Metformin (N = 159)	Linagliptin (N = 157)
Asian	n = 57; % = 35.8	n = 64 ; % = 40.8
Sample size		
Black	n = 5; % = 3.1	n = 6; % = 3.8
Sample size		
Native American/Alaskan	n = 0; % = 0	n = 2; % = 1.3
Sample size		
Comorbidities	n = NA ; % = NA	n = NA ; % = NA
Sample size		
Retinopathy	n = 7; % = 4.4	n = 6; % = 3.8
Sample size		
Neuropathy	n = 14; % = 8.8	n = 13 ; % = 8.3
Sample size		
Nephropathy	n = 2; % = 1.3	n = 2; % = 1.3
Sample size		
Coronary artery disease	n = 10; % = 6.3	n = 13 ; % = 8.3
Sample size		ŕ
Peripheral artery disease	n = 7; % = 4.4	n = 1; % = 0.6
Sample size		., .,
Cerebrovascular disease	n = 7; % = 4.4	n = 9 ; % = 5.7
Sample size		
Hypertension	n = 65; % = 40.9	n = 69 ; % = 43.9
Sample size		
Presence of frailty	n = NR ; % = NR	n = NR ; % = NR
Sample size		11 1411, 70 - 1411
Time since type 2 diabetes diagnosis	n = NA ; % = NA	n = NA ; % = NA
Sample size		, ,,
<1 year	n = 159 ; % = 100	n = 155 ; % = 98.7
Sample size		
HbA1c (%)	9.79 (1.19)	9.88 (1.1)
		5.55 ()

Characteristic	Linagliptin + Metformin (N = 159)	Linagliptin (N = 157)
Mean (SD)		
Blood pressure	NR (NR)	NR (NR)
Mean (SD)		
Heart rate	NR (NR)	NR (NR)
Mean (SD)		
Smoking status	n = NR ; % = NR	n = NR ; % = NR
Sample size		
Alcohol consumption	n = NR ; % = NR	n = NR ; % = NR
Sample size Presence of severe mental illness	n = NR ; % = NR	
Sample size	II - INIX , /0 - INIX	n = NR ; % = NR
People with significant cognitive	n = NR ; % = NR	
impairment		n = NR ; % = NR
Sample size		
People with a learning disability Sample size	n = NR ; % = NR	n = NR ; % = NR
Weight	n = NR ; % = NR	
Sample size	11 - INIX , 70 - INIX	n = NR ; % = NR
BMI (kg/m2)	29.84 (5.82)	
Mean (SD)	20.0 . (6.62)	29.63 (5.43)
Number of people with obesity	n = NR ; % = NR	n = NR ; % = NR
Sample size		
Cholesterol and lipid levels	NR (NR)	NR (NR)
Mean (SD)		
Albumin creatinine ratio	NR (NR)	NR (NR)
Mean (SD)		
eGFR (mL/min/1.73m2) Sample size	n = NA ; % = NA	n = NA ; % = NA
•	n = 87 ; % = 54.7	
Normal (≥90 ml/min/1.73 m2)	11 - 01 , 70 - 04.1	n = 90 ; % = 57.3
Sample size		

Characteristic	Linagliptin + Metformin (N = 159)	Linagliptin (N = 157)
Mild impairment (60 to <90 ml/min/1.73 m2)	n = 69; % = 43.4	n = 64 ; % = 40.8
Sample size		
Moderate impairment (30 to <60 ml/min/1.73 m2)	n = 3; % = 1.9	n = 3; % = 1.9
Sample size		
Severe impairment (<30 ml/min/1.73 m2)	n = 0; % = 0	n = 0; % = 0
Sample size		
Other antidiabetic medication used	n = 0; % = 0	n = 0 ; % = 0
Sample size		
Blood pressure-lowering medication used	n = 65; % = 40.9	n = 65 ; % = 41.4
Sample size		
Statins/lipid-lowering medication used	n = 30 ; % = 18.9	n = 33 ; % = 21
Sample size		
Other treatment being received	n = NA ; % = NA	n = NA ; % = NA
Sample size	00.0/ 40.0	
Aspirin	n = 22 ; % = 13.8	n = 20 ; % = 12.7
Sample size		

93. Russell-Jones, 2012

Bibliographic Reference

Russell-Jones, D.; Cuddihy, R. M.; Hanefeld, M.; Kumar, A.; González, J. G.; Chan, M.; Wolka, A. M.; Boardman, M. K.; Efficacy and safety of exenatide once weekly versus metformin, pioglitazone, and sitagliptin used as monotherapy in drug-naive patients with type 2 diabetes (DURATION-4): a 26-week double-blind study; Diabetes Care; 2012; vol. 35 (no. 2); 252-8

Secondary publication of another included study- see primary study for details Other publications associated with this study included in review Trial name / registration number Study type Randomised controlled trial (RCT) Study location Multicentre trial (including Argentina, Belgium, Brazil, Canada, France, Germany, Hungary, India, Israel, Italy, South Korea, Mexico, Poland, Romania, the Russian Federation, Slovakia, South Africa, Spain, Taiwan, Turkey, the United Kingdom and the United States). Study dates November 2008 to June 2010. Sources of Funded by Amylin Pharmaceuticals and Eli Lilly. Authors received grants from various pharmaceutical companies.		,
publications associated with this study included in review Trial name / registration number Study type Randomised controlled trial (RCT) Study location Multicentre trial (including Argentina, Belgium, Brazil, Canada, France, Germany, Hungary, India, Israel, Italy, South Korea, Mexico, Poland, Romania, the Russian Federation, Slovakia, South Africa, Spain, Taiwan, Turkey, the United Kingdom and the United States). Study setting Outpatient follow-up. Study dates November 2008 to June 2010. Sources of funded by Amylin Pharmaceuticals and Eli Lilly. Authors received grants from various pharmaceutical companies. Inclusion Adults with type 2 diabetes; HbA1c 7.1-11.0%; BMI 23-45 kg/m2; history of stable weight. Exclusion Criteria Treated with any antihyperglycaemic drug for >7 days within 3 months of screening. No additional information.	publication of another included study- see primary study	No additional information.
registration number Study type Randomised controlled trial (RCT) Study location Multicentre trial (including Argentina, Belgium, Brazil, Canada, France, Germany, Hungary, India, Israel, Italy, South Korea, Mexico, Poland, Romania, the Russian Federation, Slovakia, South Africa, Spain, Taiwan, Turkey, the United Kingdom and the United States). Study setting Outpatient follow-up. Study dates November 2008 to June 2010. Sources of funded by Amylin Pharmaceuticals and Eli Lilly. Authors received grants from various pharmaceutical companies. Inclusion Adults with type 2 diabetes; HbA1c 7.1-11.0%; BMI 23-45 kg/m2; history of stable weight. Exclusion Criteria Treated with any antihyperglycaemic drug for >7 days within 3 months of screening. No additional information.	publications associated with this study included in	No additional information.
Study location Multicentre trial (including Argentina, Belgium, Brazil, Canada, France, Germany, Hungary, India, Israel, Italy, South Korea, Mexico, Poland, Romania, the Russian Federation, Slovakia, South Africa, Spain, Taiwan, Turkey, the United Kingdom and the United States). Study setting Outpatient follow-up. Study dates November 2008 to June 2010. Sources of funded by Amylin Pharmaceuticals and Eli Lilly. Authors received grants from various pharmaceutical companies. Inclusion Criteria Adults with type 2 diabetes; HbA1c 7.1-11.0%; BMI 23-45 kg/m2; history of stable weight. Exclusion Criteria Recruitment / selection of participants	registration	NCT00676338. DURATION-4 trial.
Germany, Hungary, India, Israel, Italy, South Korea, Mexico, Poland, Romania, the Russian Federation, Slovakia, South Africa, Spain, Taiwan, Turkey, the United Kingdom and the United States). Study setting Outpatient follow-up. Study dates November 2008 to June 2010. Sources of funded by Amylin Pharmaceuticals and Eli Lilly. Authors received grants from various pharmaceutical companies. Inclusion Criteria Adults with type 2 diabetes; HbA1c 7.1-11.0%; BMI 23-45 kg/m2; history of stable weight. Exclusion Criteria Treated with any antihyperglycaemic drug for >7 days within 3 months of screening. No additional information.	Study type	Randomised controlled trial (RCT)
Study dates November 2008 to June 2010. Sources of funding Funded by Amylin Pharmaceuticals and Eli Lilly. Authors received grants from various pharmaceutical companies. Inclusion Adults with type 2 diabetes; HbA1c 7.1-11.0%; BMI 23-45 kg/m2; history of stable weight. Exclusion Criteria Treated with any antihyperglycaemic drug for >7 days within 3 months of screening. Recruitment / selection of participants No additional information.	Study location	Germany, Hungary, India, Israel, Italy, South Korea, Mexico, Poland, Romania, the Russian Federation, Slovakia, South Africa, Spain, Taiwan,
Sources of funding Funded by Amylin Pharmaceuticals and Eli Lilly. Authors received grants from various pharmaceutical companies. Inclusion Adults with type 2 diabetes; HbA1c 7.1-11.0%; BMI 23-45 kg/m2; history of stable weight. Exclusion Treated with any antihyperglycaemic drug for >7 days within 3 months of screening. Recruitment / selection of participants Funded by Amylin Pharmaceuticals and Eli Lilly. Authors received grants from various pharmaceuticals and Eli Lilly. Authors received grants from various pharmaceuticals and Eli Lilly. Authors received grants from various pharmaceuticals and Eli Lilly. Authors received grants from various pharmaceuticals and Eli Lilly. Authors received grants from various pharmaceuticals and Eli Lilly. Authors received grants from various pharmaceuticals and Eli Lilly. Authors received grants from various pharmaceuticals and Eli Lilly. Authors received grants from various pharmaceuticals and Eli Lilly. Authors received grants from various pharmaceuticals and Eli Lilly. Authors received grants from various pharmaceuticals and Eli Lilly. Authors received grants from various pharmaceutical companies.	Study setting	Outpatient follow-up.
funding from various pharmaceutical companies. Inclusion Adults with type 2 diabetes; HbA1c 7.1-11.0%; BMI 23-45 kg/m2; history of stable weight. Exclusion Treated with any antihyperglycaemic drug for >7 days within 3 months of screening. Recruitment / selection of participants From various pharmaceutical companies. Adults with type 2 diabetes; HbA1c 7.1-11.0%; BMI 23-45 kg/m2; history of stable weight. Treated with any antihyperglycaemic drug for >7 days within 3 months of screening.	Study dates	November 2008 to June 2010.
criteria stable weight. Exclusion Treated with any antihyperglycaemic drug for >7 days within 3 months of screening. Recruitment / selection of participants Stable weight. Treated with any antihyperglycaemic drug for >7 days within 3 months of screening.		
criteria screening. Recruitment / No additional information. selection of participants		Adults with type 2 diabetes; HbA1c 7.1-11.0%; BMI 23-45 kg/m2; history of stable weight.
selection of participants		, ,, ,,
Intervention(s) Exenatide N=248	selection of	No additional information.
	Intervention(s)	Exenatide N=248

	Exenatide 2.0mg once a week subcutaneously plus daily placebo oral tablet for 26 weeks. Additional safety data was obtained 10 weeks after this. Concomitant therapy: Standard diet and exercise counselling was provided in each treatment group.
Strata 1: People with type 2 diabetes mellitus and heart failure	Not stated/unclear
Strata 2: People with atherosclerotic cardiovascular diseases	Not stated/unclear
Strata 3: People with type 2 diabetes mellitus and chronic kidney disease	Not stated/unclear
Strata 4: People with type 2 diabetes mellitus and high cardiovascular risk	People at higher risk of developing cardiovascular disease Based on BMI and presence of diabetes.
Subgroup 1: People with frailty	Not stated/unclear
Subgroup 2: Onset of type 2 diabetes mellitus	Not stated/unclear
Subgroup 3: People with non-alcoholic fatty liver disease	Not stated/unclear
Subgroup 4: People with obesity	Not stated/unclear

Subgroup 5: eGFR category at baseline	Not stated/unclear
Subgroup 6: Albuminuria category at baseline	Not stated/unclear
Sensitivity analysis category: Enrichment trial status	6) No response criteria
Population subgroups	No additional information.
Comparator	Metformin N=246
	Metformin increased in weekly increments up to target doses of 2000mg/day (or up to 2500mg/day based on glycaemic control) and subcutaneous placebo once a week for 26 weeks. Additional safety data was obtained 10 weeks after this.
	Concomitant therapy: Standard diet and exercise counselling was provided in each treatment group.
	Pioglitazone N=163
	Pioglitazone increased in weekly increments up to target doses of 45mg/day and subcutaneous placebo once a week for 26 weeks. Additional safety data was obtained 10 weeks after this.
	Concomitant therapy: Standard diet and exercise counselling was provided in each treatment group.
	Sitagliptin N=163
	Sitagliptin 100mg/day and subcutaneous placebo once a week for 26 weeks. Additional safety data was obtained 10 weeks after this.
	Concomitant therapy: Standard diet and exercise counselling was provided in each treatment group.
Number of participants	820
Duration of follow-up	26 weeks.
Indirectness	No additional information.

Method of analysis	ITT
Additional comments	No additional information.

93.2.1. Exenatide (N = 248)

Exenatide 2.0mg once a week subcutaneously plus daily placebo oral tablet for 26 weeks. Additional safety data was obtained 10 weeks after this. Concomitant therapy: Standard diet and exercise counselling was provided in each treatment group.

93.2.2. Metformin (N = 246)

Metformin increased in weekly increments up to target doses of 2000mg/day (or up to 2500mg/day based on glycaemic control) and subcutaneous placebo once a week for 26 weeks. Additional safety data was obtained 10 weeks after this. Concomitant therapy: Standard diet and exercise counselling was provided in each treatment group.

93.2.3. **Pioglitazone (N = 163)**

Pioglitazone increased in weekly increments up to target doses of 45mg/day and subcutaneous placebo once a week for 26 weeks. Additional safety data was obtained 10 weeks after this. Concomitant therapy: Standard diet and exercise counselling was provided in each treatment group.

93.2.4. Sitagliptin (N = 163)

Sitagliptin 100mg/day and subcutaneous placebo once a week for 26 weeks. Additional safety data was obtained 10 weeks after this. Concomitant therapy: Standard diet and exercise counselling was provided in each treatment group.

93.3. Characteristics

Characteristic	•	•	Pioglitazone (N = 163)	Sitagliptin (N = 163)
% Male	n = 139 ; % = 56	n = 154 ; % =	n = 97 ; % =	n = 94 ; % =
Sample size		62.6	59.5	57.7

Mean age (SD) (years) 54 (11) 54 (11) 55 (11) 52 (11)					
Mean (SD) 54 (11) 55 (11) 52 (11) Ethnicity n = NA; % = NA Sample size n = 169; % = 68.1 n = 160; % = 67.5 n = 110; % = n = 113; % = 69.3 n = 113; % = 69.3 East Asian n = 34; % = 12.6 n = 19; % = n = 18; % = 11.7 n = 18; % = 11.7 n = 18; % = 11.7 Sample size n = 21; % = 8.5 n = 20; % = 8.1 n = 15; % = 9.2 n = 13; % = 8 Sample size n = 16; % = 8.5 n = 21; % = 11; % = 11; % = 11 n = 15; % = 9.2 n = 13; % = 8 n = 11; % = 9.2 n = 13; % = 8 n = 15; % = 9.2 n = 13; % = 8 n = 11; % = 9.2 n = 13; % = 8 n = 15; % = 9.2 n = 13; % = 8 n = 15; % = 9.2 n = 13; % = 8 n = 11; % = 11 n = 11; % = 11 n = 13; % = 11 n = 11; % = 11 n = 13; % = 11 n = 11; % = 11 n = 13; % = 11 n = 13; % = 11 n = 11; % = 11 n = 13; % = 11 n = 11; % = 11 n = 11; % = 11 n = 13; % = 11 n = 11; % = 11 n = 13; % = 11 n = 11; % = 11 n = 13; % = 11 n = 13	Characteristic	•	•		
Mean (SD) Ethnicity n = NA; % = NA n = 113; % = NA n = 113; % = NA; % = NA n = 113; % = NA n	Mean age (SD) (years)	54 (11)	54 (11)	55 (11)	52 (11)
Sample size NA n = NA; % = n = NA; % = NA n = 110; % = n = 110; % = n = 110; % = n = 113; % = n = 112. n = 113; % = n = 112; % = n = 112. n = 19; % = n = n = 113; % = n = 112; % = n = 12. n = 15; % = 9.2 n = 13; % = n = 11; % = n =	Mean (SD)		` ,	,	, ,
Caucasian n = 169; % = 68.1 n = 160; % = 67.5 n = 110; % = 69.3 East Asian n = 34; % = 13.7 n = 31; % = 11.7 n = 19; % = 11.7 n = 18; % = 11.7 Sample size n = 21; % = 8.5 n = 20; % = 8.1 n = 15; % = 9.2 n = 15; % = 9.2 West Asian n = 16; % = 6.5 n = 21; % = 8.1 n = 15; % = 9.2 n = 15; % = 9.2 Sample size n = 16; % = 6.5 n = 21; % = 11	Ethnicity		•	n = NA ; % = NA	•
Sample size 68.1 n = 160; % = 67.5 n = 110; % = 69.3 n = 113; % = 69.3 East Asian n = 34; % = 13.7 n = 31; % = 11.7 n = 19; % = 11.7 n = 18; % = 11.7 West Asian n = 21; % = 8.5 n = 20; % = 8.1 n = 15; % = 9.2 n = 13; % = 18.8 n = 11; % = 9.2 n = 13; % = 18.8 n = 11; % = 9.2 n = 13; % = 18.8 n = 11; % = 9.2 n = 13; % = 18.8 n = 13; % = 18.8 n = 11; % = 9.2 n = 13; % = 18.8 n			INA		IVA
East Asian n = 34; % = 13.7 n = 31; % = 11.7 n = 19; % = 11.7 n = 18; % = 11.7 West Asian n = 21; % = 8.5 n = 20; % = 8.1 n = 15; % = 9.2 n = 13; % = 8.5 Sample size African n = 7; % = 2.8 n = 11; % = 1.8 n = 4; % = 2.5 n = 3; % = 1.8 n = 3; % = 1.8 n = 4; % = 2.5 n = 3; % = 1.8 n = 3; % = 1.8 n = 0; % = 0 n = 1; % = 0.6		,			· ·
Sample size 12.6 11.7 11 West Asian n = 21; % = 8.5 n = 20; % = 8.1 n = 15; % = 9.2 n = 15; % = 9.2 Sample size n = 16; % = 6.5 n = 21; % = n = 15; % = 9.2 n = 13; % = 8 Sample size n = 7; % = 2.8 n = 11; % = n = 4; % = 2.5 n = 3; % = 1.8 Sample size n = 1; % = 0.4 n = 3; % = 1.8 n = 0; % = 0 n = 1; % = 0.6 Comorbidities n = NR; % = NR Sample size n = NR; % = NR n = NR	East Asian	n = 34 ; % =			
Sample size 8.5 n = 20; % = 8.1 n = 15; % = 9.2 n = 13; % = 8 n = 21; % = n = 15; % = 9.2 n = 13; % = 8 n = 11; % = n = 4; % = 2.5 n = 13; % = 1.8 n = 3; % = 1.8 n = 0; % = 0 n = 1; % = 0.6	Sample size	· ·	•	•	·
Sample size 8.1 9.2 Hispanic n = 16; % = 6.5 n = 21; % = 8.5 n = 15; % = 9.2 n = 13; % = 8 Sample size n = 7; % = 2.8 n = 11; % = 1.8 n = 4; % = 2.5 n = 3; % = 1.8 Sample size n = 1; % = 0.4 n = 3; % = 1.8 n = 0; % = 0 n = 1; % = 0.6 Comorbidities n = NR; % = NR Sample size n = NR; % = NR Presence of frailty n = NR; % = NR Sample size n = NR; % = NR NR NR NR Heart size NR <	West Asian	· ·	n = 20 : % =	n = 15 : % = 9.2	n = 15 : % =
Sample size African n = 7; % = 2.8 n = 11; % = n = 4; % = 2.5 n = 3; % = 1.8 Sample size n = 1; % = 0.4 n = 11; % = 0.4 n = 4; % = 2.5 n = 3; % = 1.8 Other n = 1; % = 0.4 n = 3; % = n = 0; % = 0 n = 1; % = 0.6 n = 1; % = 0.6 Sample size n = NR; % = NR NR <td>Sample size</td> <td>0.0</td> <td>•</td> <td>, ,, ,, 0.12</td> <td></td>	Sample size	0.0	•	, ,, ,, 0.12	
African Sample size African Sample size Other Sample size Other Sample size Comorbidities Sample size Presence of frailty Sample size Time since type 2 diabetes diagnosis (years) Mean (SD) HbA1c (%) Blood pressure NR (NR)	Hispanic		•	n = 15 ; % = 9.2	n = 13 ; % = 8
Sample size 2.8 n = 11; % = 4.5 n = 4; % = 2.5 n = 3; % = 1.8 Other n = 1; % = 0.4 n = 3; % = 1.2 n = 0; % = 0 n = 1; % = 0.6 Sample size n = NR; % = NR Presence of frailty n = NR; % = NR Sample size 2.7 (3.2) 2.6 (3.6) 2.7 (3.7) 2.7 (3.7) Mean (SD) 8.5 (1.2) 8.6 (1.2) 8.5 (1.2) 8.5 (1.3) Mean (SD) NR (NR) NR (NR) NR (NR) NR (NR) NR (NR) Heart rate NR (NR) NR (NR) NR (NR) NR (NR) NR (NR)	Sample size		0.5		
Other n = 1; % = 0.4 n = 3; % = 1.2 n = 0; % = 0 n = 1; % = 0.6 Comorbidities n = NR; % = NR	African Sample size	·	·	n = 4; % = 2.5	·
Sample size 0.4 n = 3; % = n = 0; % = 0 n = 1; % = 0.6 Comorbidities n = NR; % = NR n = NR;		n - 1 · % -			
Comorbidities n = NR; % = NR n = NR;		·	· ·	n = 0; % = 0	· ·
Sample size NR n = NR; % = NR		n = NR · % =			
Presence of frailty n = NR; % = NR n = NR; NR n = NR				n = NR ; % = NR	
Sample size NR n = NR; % = NR n = NR; % = NR n = NR; % = NR Time since type 2 diabetes diagnosis (years) 2.7 (3.2) 2.6 (3.6) 2.7 (3.7) 2.7 (3.7) Mean (SD) 8.5 (1.2) 8.6 (1.2) 8.5 (1.2) 8.5 (1.3) Mean (SD) NR (NR) NR (NR) NR (NR) NR (NR) Mean (SD) NR (NR) NR (NR) NR (NR) NR (NR) Heart rate NR (NR) NR (NR) NR (NR) NR (NR)			IVIX		IVIX
Time since type 2 diabetes diagnosis (years) Mean (SD) HbA1c (%) Blood pressure NR (NR)	Presence of frailty		·	n = NR ; % = NR	·
diabetes diagnosis (years) 2.6 (3.6) 2.7 (3.7) 2.7 (3.7) Mean (SD) 8.5 (1.2) 8.6 (1.2) 8.5 (1.2) 8.5 (1.3) Mean (SD) NR (NR) NR (NR) NR (NR) NR (NR) Mean (SD) NR (NR) NR (NR) NR (NR) NR (NR) Heart rate NR (NR) NR (NR) NR (NR) NR (NR)	Sample size		NR		NR
HbA1c (%) 8.5 (1.2) 8.6 (1.2) 8.5 (1.2) 8.5 (1.3) Mean (SD) NR (NR) NR (NR) NR (NR) NR (NR) Mean (SD) NR (NR) NR (NR) NR (NR) NR (NR) Heart rate NR (NR) NR (NR) NR (NR)	Time since type 2 diabetes diagnosis (years)	2.7 (3.2)	2.6 (3.6)	2.7 (3.7)	2.7 (3.7)
NR (NR) NR (Mean (SD)				
Blood pressure NR (NR)	HbA1c (%)	8.5 (1.2)	8.6 (1.2)	8.5 (1.2)	8.5 (1.3)
Mean (SD) Heart rate NR (NR)	Mean (SD)				
Mean (SD) Heart rate NR (NR) NR (NR) NR (NR) NR (NR)	Blood pressure	NR (NR)	NR (NR)	NR (NR)	NR (NR)
NR (NR) NR (NR) NR (NR)	Mean (SD)				
	Heart rate	NR (NR)	NR (NR)	NR (NR)	NR (NR)
	Mean (SD)		(-)	()	(-/

Characteristic	Exenatide (N = 248)	Metformin (N = 246)	Pioglitazone (N = 163)	Sitagliptin (N = 163)
Smoking status Sample size	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
Alcohol consumption	n = NR ; % =			
Sample size	NR	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
Presence of severe mental illness	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
Sample size				
People with significant cognitive impairment	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
Sample size				
People with a learning disability	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
Sample size				
Weight (kg)	87.5 (18.9)	85.9 (19.6)	86.1 (17.8)	88.7 (18.7)
Mean (SD)				
BMI (kg/m2)	31.4 (5.3)	30.7 (5.5)	31.1 (5.3)	31.8 (5.4)
Mean (SD)				
Number of people with obesity	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
Sample size				
Cholesterol and lipid levels	NR (NR)	NR (NR)	NR (NR)	NR (NR)
Mean (SD)				
Albumin creatinine ratio	NR (NR)	NR (NR)	NR (NR)	NR (NR)
Mean (SD)				
eGFR (mL/min/1.73m2)	NR (NR)	NR (NR)	NR (NR)	NR (NR)
Mean (SD)				
Other antidiabetic medication used	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
Sample size				
Blood pressure-lowering medication used	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR

Characteristic	Exenatide (N = 248)	•	Pioglitazone (N = 163)	Sitagliptin (N = 163)
Sample size				
Statins/lipid-lowering medication used Sample size	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
Other treatment being	n = NR ; % =			
received	NR	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
Sample size				

94. Scherbaum, 2002

Bibliographic Reference

Scherbaum, W. A.; Göke, B.; Metabolic efficacy and safety of once-daily pioglitazone monotherapy in patients with type 2 diabetes: a double-blind, placebo-controlled study; Horm Metab Res; 2002; vol. 34 (no. 10); 589-95

94.1. S	tudy details
Secondary publication of another included study- see primary study for details	NA
Other publications associated with this study included in review	NA
Trial name / registration number	NR
Study type	Randomised controlled trial (RCT)
Study location	Germany
Study setting	Multicentre
Study dates	NR
Sources of funding	This study was supported by Takeda Pharmaceuticals, Europe
Inclusion criteria	- Female participants in the study had to be postmenopausal, surgically sterilised, or using appropriate contraceptive methods to avoid pregnancy
Exclusion criteria	 -Type 1 diabetes - Secondary failure to treatment with sulphonylureas - Requirement for other antidiabetic treatment - History of ketoacidosis, malabsorption, acute or chronic pancreatitis, liver disease, significant ventricular hypertrophy, complex cardiac arrhythmias, angina pectoris, heart failure, myocardial infarction, hypertension (diastolic pressure > 100 mmHg), stroke, or hypothyroidism - History of transient ischaemic attack or stroke

	- significant anaemia of any aetiology
	- clinically relevant haematological or malignant disease in the last 10 years
	- HIV infection
	- alcohol or drug abuse
	- participation in a clinical trial in the 3 months prior to study
Recruitment / selection of participants	Following enrolment in the study, patients were required to discontinue their previous oral antidiabetic therapy and to enter a 10-week placebo washout period. During this time, patients underwent a complete medical history and physical examination, including vital signs, laboratory tests, and an electrocardiogram to determine eligibility for the double-blind phase of the study. At the end of the washout period, HbA1C had to remain between 7.5% and 12 %, and FBG had to be between 140 mg/dl and 250 mg/ dl.
Intervention(s)	Eligible patients were randomised to one of three treatment regimens for 26 weeks: pioglitazone 15 mg once-daily + dietary controls, pioglitazone 30 mg once-daily + diet
Cointervention	Patients were required to follow a disease- and bodyweight-oriented diet throughout the study period. They were taught the nutritional recommendations of the Diabetes and Nutrition Study Group (DNSG) of the European Association for the Study of Diabetes (EASD). The recommended diet included a protein intake of $10\% - 20\%$ of total daily calories. The carbohydrate: fat ratio was individualised, based on patients' eating habits as well as on glucose and lipid goals. For patients with normal lipid profiles, 30% of the total daily caloric consumption was to be derived from fat, with an equal distribution among saturated, polyunsaturated, and monounsaturated fats. These recommendations were made prior to the start of the study. Diet was monitored at each visit to the clinic, and dietary advice was continuously available on request. However, compliance to diet was not monitored (such as in a diary). Also, bodyweight was recorded at the clinic at each visit, as well as each week by the patient at home.
Strata 1: People with type 2 diabetes mellitus and heart failure	People without heart failure
Strata 2: People with	Not stated/unclear
_	Myocardial infarction and stroke excluded, others not stated
Strata 3: People with type 2 diabetes	Not stated/unclear

mellitus and chronic kidney disease	
Strata 4: People with type 2 diabetes mellitus and high cardiovascular risk	Not stated/unclear
Subgroup 1: People with frailty	Not stated/unclear
Subgroup 2: Onset of type 2 diabetes mellitus	Not stated/unclear
Subgroup 3: People with non-alcoholic fatty liver disease	Not stated/unclear
Subgroup 4: People with obesity	Not stated/unclear
Subgroup 5: eGFR category at baseline	Not stated/unclear
Subgroup 6: Albuminuria category at baseline	Not stated/unclear
Sensitivity analysis category: Enrichment trial status	2) Excluding non-responders (but not exclusively including only responders) HbA1C had to remain between 7.5% and 12%, and FBG had to be between 140 mg/dl and 250 mg/ dl.
Population subgroups	
Comparator	diet + matching placebo (identical in colour, shape, taste, size, and odour to active treatment).
Number of participants	252
Duration of follow-up	26 weeks
Indirectness	Direct

Method of analysis	Per protocol
	ITT

94.3. Characteristics

Characteristic	Placebo + diet (N = 84)	Placebo 15mg / day + diet (N = 89)	Placebo 30mg / day + diet (N = 78)
Male (n (%))	n = 47; % = 55.9	n = 56 ; % = 62.9	n = 32 ; % = 41
Sample size			
Aged >= 65 years	n = 27 ; % = 32.1	n = 24 ; % = 26.9	n = 26 ; % = 33.3
Sample size			
Diabetes duration (yrs)	5.6	5.4	4.6
Nominal			

95. Scherbaum, 2008

Bibliographic Reference

Scherbaum, W. A.; Schweizer, A.; Mari, A.; Nilsson, P. M.; Lalanne, G.; Jauffret, S.; Foley, J.; Efficacy and tolerability of vildagliptin in drug-naive patients with type 2 diabetes and mild hyperglycaemia; Diabetes Obes Metab; 2008; vol. 10 (no. 8); 675-682

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Secondary	Yes, see:	
publication of another included study- see primary study for details	 Mari, A., Scherbaum, W. A., Nilsson, P. M., Lalanne, G., Schweizer, A., Dunning, B. E., & Foley, J. E. (2008). Characterization of the influence of vildagliptin on model-assessed β-cell function in patients with type 2 diabetes and mild hyperglycaemia. <i>The Journal of Clinical Endocrinology & Metabolism</i>, 93(1), 103-109. 	
Other	Original article reported in:	
publications associated	 Mari, A., Scherbaum, W. A., Nilsson, P. M., Lalanne, G., 	
with this study included in	Schweizer, A., Dunning, B. E., & Foley, J. E. (2008).	
review	Characterization of the influence of vildagliptin on model-assessed β-cell function in patients with type 2 diabetes and mild	
	hyperglycaemia. The Journal of Clinical Endocrinology & Metabolism, 93(1), 103-109.	
	Wetabolishi, 93(1), 103-109.	
Trial name /	NCT00101712	
registration number		
Study type	Randomised controlled trial (RCT)	
Study location	International (69 sites in 6 countries: in Finland (3 sites), France (4 sites), Germany (42 sites), Romania (5 sites), Spain (7 sites), Sweden (8 sites))	
Study setting	Community	
Study dates	10/2004 to 05/2006	
Sources of funding	Funded by Novartis Pharmaceuticals Corporation.	
Inclusion	Male or female (non-fertile or of childbearing potential using	
criteria	medically approved birth control method) • Aged≥18 years	
	Diagnosis of type 2 diabetes mellitus for at least 8 weeks proviously.	
	previouslyHbA1c level 6.2-7.5% inclusive at screening (upper limit of 7% in	
	Finland and Spain) • BMI 22-45 kg/m2 inclusive	
	• DIVIT ZZ-40 Ng/IIIZ IIIOUSIVE	

	 Drug-naive (no oral antidiabetic drug [OAD] for at least 12 weeks prior to screening and no OAD for more than 3 consecutive months at any time in past)
Exclusion criteria	 History of type 1 or secondary forms of diabetes Acute metabolic diabetic complications within the past 6 months or evidence of significant diabetic complications History of significant cardiac arrhythmia, congestive heart failure, or New York Heart Association Class III or IV Liver diseases (e.g. cirrhosis or chronic active hepatitis) Significant laboratory abnormalities
Recruitment / selection of participants	Participants recruited from 69 sites in 6 countries. Assessment at screening visit (week 2) for eligibility and randomisation at visit 2 to arms. All participants received individualized lifestyle counselling (weight management, diet, exercise) at each visit. Assessments at weeks 4, 12, 16, 24, 32, 40 and 52, followed by 4 week treatment-free period and final assessment at week 56. All lab assessments conducted by same laboratory according to standardised/validated procedures.
Intervention(s)	Vildagliptin 50 mg once daily
	Oral vildagliptin 50 mg once daily for 52 weeks.
Strata 1: People with type 2 diabetes mellitus and heart failure	People without heart failure Exclusion criteria: congestive heart failure, NYHA class 3 and 4
Strata 2: People with atherosclerotic cardiovascular diseases	Not stated/unclear
Strata 3: People with type 2 diabetes mellitus and chronic kidney disease	Not stated/unclear
Strata 4: People with type 2 diabetes mellitus and high cardiovascular risk	Not stated/unclear
Subgroup 1: People with frailty	Not stated/unclear

Subgroup 2: Onset of type 2 diabetes mellitus	Not stated/unclear	
Subgroup 3:	People without non-alcoholic fatty liver disease	
People with non-alcoholic fatty liver disease	Exclusion criteria: History of liver disease	
Subgroup 4: People with obesity	Not stated/unclear	
Subgroup 5: eGFR category at baseline	Not stated/unclear	
Subgroup 6: Albuminuria category at baseline	Not stated/unclear	
Sensitivity analysis	7) Mixed population	
category: Enrichment trial status	Inclusion criteria: no oral anti-diabetic drug (OAD) for at least 12 weeks prior to screening and no OAD for more than three consecutive months at any time in the past	
Population subgroups	Results also reported for these subgroups: Obesity Non-obesity	
Comparator	Placebo Oral matching placebo for 52 weeks.	
Number of participants	N=306	
Duration of follow-up	52 weeks + 4 week washout period	
Method of analysis	ACA	
anaiyəiə	Not reported but appears to be available case analysis (different number of participants for each reported outcome)	

95.2.1. Vildagliptin 50 mg once daily (N = 156)

Oral vildagliptin 50 mg once daily for 52 weeks.

95.2.2. Placebo (N = 150)

Matching placebo for 52 weeks.

95.3. Characteristics

95.3.1. Arm-level characteristics

Characteristic	Vildagliptin 50 mg once daily (N = 156)	Placebo (N = 150)
HbA1c (%) Significant difference between groups, p=0.0403	6.7 (0.4)	6.8 (0.4)
Mean (SD)		

See Mari 2008 for baseline characteristics

96. Schernthaner, 2004

Bibliographic Reference

Schernthaner, G.; Matthews, D. R.; Charbonnel, B.; Hanefeld, M.; Brunetti, P.; Efficacy and safety of pioglitazone versus metformin in patients with type 2 diabetes mellitus: a double-blind, randomized trial; J Clin Endocrinol Metab; 2004; vol. 89 (no. 12); 6068-76

Secondary publication of another included study- see primary study for details Other publications associated with this study included in review Trial name / registration number Study type Randomised controlled trial (RCT) Study location Multicentre trial (12 European countries).
publications associated with this study included in review Trial name / registration number Study type Randomised controlled trial (RCT) Study location Multicentre trial (12 European countries).
registration number Study type Randomised controlled trial (RCT) Study location Multicentre trial (12 European countries).
Study location Multicentre trial (12 European countries).
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Ctudy acting Outpatient follow up
Study setting Outpatient follow-up.
Study dates No additional information.
Sources of No additional information. funding
Inclusion People with HbA1c between 7.5% and 11% with stable or worsening glycaemic control for at least 3 months.
Prior use of glucose-lowering pharmacotherapy and specific contraindications to either drug (corticosteroids and beta blockers were permitted if commenced at least 4 weeks before screening, antihypertensives [except thiazides] were allowed dependent on clinical need, lipid lowering agents were allowed).
Recruitment / No additional information. selection of participants
Intervention(s) Pioglitazone N=597

	Initially 30mg pioglitazone, increased to up to 45mg of pioglitazone once a day with metformin placebo three times a day for a 12 week forced titration period followed by a 40 week maintenance period. 52 weeks in total.
	Concomitant therapy: Dietary advice was given at baseline and if body weight increased by more than 5% at any stage of HbA1c increased to greater than 9% after complete dose titrated, people were given additional intensive dietary counselling.
Strata 1: People with type 2 diabetes mellitus and heart failure	Not stated/unclear
Strata 2: People with atherosclerotic cardiovascular diseases	
Strata 3: People with type 2 diabetes mellitus and chronic kidney disease	Not stated/unclear
Strata 4: People with type 2 diabetes mellitus and high cardiovascular risk	People at higher risk of developing cardiovascular disease Based on BMI, age and presence of diabetes.
Subgroup 1: People with frailty	Not stated/unclear
Subgroup 2: Onset of type 2 diabetes mellitus	Not stated/unclear
Subgroup 3: People with non-alcoholic fatty liver disease	Not stated/unclear
Subgroup 4: People with obesity	Not stated/unclear

Not stated/unclear
NOT Stated/ulloleal
Not stated/unclear
5) All treatment naïve
No additional information.
Initially 850mg metformin once a day, increased up to a maximum of 850mg three times a day with pioglitazone placebo once a day for a 12 week forced titration period followed by a 40 week maintenance period. 52 weeks in total. Concomitant therapy: Dietary advice was given at baseline and if body weight increased by more than 5% at any stage of HbA1c increased to greater than 9% after complete dose titrated, people were given additional intensive dietary counselling.
1194
12 months.
No additional information.
ITT
No additional information.
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96.2.1. Pioglitazone (N = 597)

Initially 30mg pioglitazone, increased to up to 45mg of pioglitazone once a day with metformin placebo three times a day for a 12 week forced titration period followed by a 40 week maintenance period. 52 weeks in total. Concomitant therapy: Dietary advice was given at baseline and if body weight increased by more than 5% at any stage of HbA1c increased to greater than 9% after complete dose titrated, people were given additional intensive dietary counselling.

96.2.2. Metformin (N = 597)

Initially 850mg metformin once a day, increased up to a maximum of 850mg three times a day with pioglitazone placebo once a day for a 12 week forced titration period followed by a 40 week maintenance period. 52 weeks in total. Concomitant therapy: Dietary advice was given at baseline and if body weight increased by more than 5% at any stage of HbA1c increased to greater than 9% after complete dose titrated, people were given additional intensive dietary counselling.

96.3. Characteristics

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Characteristic	Pioglitazone (N = 597)	Metformin (N = 597)
% Male	n = 314 ; % = 53	n = 345 ; % = 58
Sample size		
Mean age (SD) (years)	57 (9.4)	56 (9.3)
Mean (SD)		
Ethnicity	n = NR ; % = NR	n = NR ; % = NR
Sample size		
Comorbidities Sample size	n = NR ; % = NR	n = NR ; % = NR
Presence of frailty	n = NR ; % = NR	
Presence of framty	II - INIX , /0 - INIX	n = NR; % = NR
Sample size		
Time since type 2 diabetes diagnosis (years)	3.4 (4.3)	3.1 (3.8)
Mean (SD)		
HbA1c (%)	8.7 (1)	8.7 (1)
Mean (SD)		
Blood pressure	NR (NR)	NR (NR)
Mean (SD)		
Heart rate	NR (NR)	NR (NR)
Mean (SD)		
Smoking status	n = NR ; % = NR	n = NR ; % = NR
Sample size		

Characteristic	Pioglitazone (N = 597)	Metformin (N = 597)
Alcohol consumption	n = NR ; % = NR	n = NR ; % = NR
Sample size		
Presence of severe mental illness Sample size	n = NR ; % = NR	n = NR ; % = NR
•	ND 0/ ND	
People with significant cognitive impairment	n = NR ; % = NR	n = NR ; % = NR
Sample size		
People with a learning disability	n = NR ; % = NR	n = NR ; % = NR
Sample size		
Weight (kg) Mean (SD)	88.2 (15.5)	89.7 (16.6)
	21 2 (4 0)	
BMI (kg/m2) Mean (SD)	31.2 (4.9)	31.4 (5.2)
Number of people with obesity	n = NR ; % = NR	n = NR ; % = NR
Sample size		
Cholesterol and lipid levels	NR (NR)	NR (NR)
Mean (SD)		
Albumin creatinine ratio Mean (SD)	NR (NR)	NR (NR)
eGFR (mL/min/1.73m2)	NR (NR)	
Mean (SD)	IVIX (IVIX)	NR (NR)
Other antidiabetic medication used	n = NR ; % = NR	
Sample size		n = NR ; % = NR
Blood pressure-lowering medication used	n = NR ; % = NR	n = NR ; % = NR
Sample size		
Statins/lipid-lowering medication used	n = NR ; % = NR	n = NR ; % = NR
Sample size		
Other treatment being received Sample size	n = NR ; % = NR	n = NR ; % = NR
Jampie Size		

97. Schwartz, 2006

Bibliographic Reference

Schwartz, Sherwyn; Fonseca, Vivian; Berner, Bret; Cramer, Marilou; Chiang, Yu-Kun; Lewin, Andrew; Efficacy, tolerability, and safety of a novel once-daily extended-release metformin in patients with type 2 diabetes.; Diabetes care; 2006; vol. 29 (no. 4); 759-64

97.1. 5	tudy details
Trial name / registration number	NR
Study type	Randomised controlled trial (RCT)
Study location	USA
Study setting	85 centres
Study dates	August 2001 - October 2003
Sources of funding	Financial support for this study was provided by Depomed
Inclusion criteria	 - 18-79 years of age - Type 2 diabetes - Patients were either drug naive (with newly diagnosed diabetes or treated with diet and exercise only) or had received prior drug therapy (monotherapy with oral hypoglycaemic agents other than metformin up to the maximum dose allowed, metformin monotherapy up to 2,000 mg/day, or metformin up to 1,500 mg/day with sulfonylurea up to one-half the maximum allowed dose) - HbA1c levels 7-12% (drug-naive patients) or 6.5 - 10% (prior drug therapy patients) - FPG levels 120-400mg/dl (drug-naive patients) or 120-250 mg/dl (prior drug therapy patients), C-peptide levels >1.0 ng/ml, BMI 22-50kg/m2 - Negative pregnancy test for female patients
Exclusion criteria	 -receiving insulin, systemic corticosteroids, nicotinic acid, or isoniazid -History of had a history of background retinopathy, symptomatic autonomic neuropathy, or unstable angina - Chronic gastroparesis or chronic severe gastrointestinal symptoms, a history of gastric or duodenal ulcers, abdominal surgery within 1 year, or active gastrointestinal disease within 2 years - Any uncontrolled or untreated cardiovascular, hepatic, pulmonary, renal, or neurological system conditions
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Recruitment / selection of participants	The trial enrolled male and female outpatients, 18–79 years of age, with type 2 diabetes. Patients were either drug naïve (with newly diagnosed diabetes or treated with diet and exercise only) or had received prior drug therapy (monotherapy with oral hypoglycaemic agents other than metformin up to the maximum dose allowed, metformin monotherapy up to 2,000 mg/day, or metformin up to 1,500 mg/day with sulfonylurea up to one-half the maximum allowed dose). Patients underwent a full physical examination. After a 6-week washout of current antihyperglycemic agents (for prior drug therapy patients), all patients began metformin dosing at 1,000 mg q.d., which was titrated to their assigned dose over 2–3 weeks and continued at that dose for a total treatment duration of 24 weeks. All study drugs and placebos were taken after a meal. Patients were evaluated every 1–2 weeks during the screening and washout periods, weekly for the first 4 weeks of treatment and then every 4 weeks until the end of study.
Intervention(s)	
Cointervention	appropriate placebo tablets to maintain the study blind
Strata 1: People with type 2 diabetes mellitus and heart failure	Not stated/unclear
Strata 2: People with atherosclerotic cardiovascular diseases	Mixed population Angina only mentioned
Strata 3: People with type 2 diabetes mellitus and chronic kidney disease	Not stated/unclear
Strata 4: People with type 2 diabetes mellitus and high cardiovascular risk	Not stated/unclear
Subgroup 1: People with frailty	Not stated/unclear
Subgroup 2: Onset of type 2 diabetes mellitus	Not stated/unclear

Subgroup 3: People with non-alcoholic fatty liver disease	Not stated/unclear
Subgroup 4: People with obesity	Not stated/unclear
Subgroup 5: eGFR category at baseline	Not stated/unclear
Subgroup 6: Albuminuria category at baseline	Not stated/unclear
Sensitivity analysis category: Enrichment trial status	7) Mixed population
Number of participants	750
Duration of follow-up	24 weeks
Indirectness	direct
Method of analysis	Per protocol ITT

97.2.1. ER metformin 1,500mg /day (N = 178) Extended-release metformin (Glumetza; Depomed, Menlo Park, CA),

97.2.2. ER metformin 1,500mg (AM/PM) (N = 182) 500 mg in the morning and 1,000 mg in the evening

97.2.3. ER metformin 2,000 mg / day (N = 172)

97.2.4. 1,500 mg immediate-release metformin (N = 174) (Glucophage; Bristol-Myers Squibb, Princeton, NJ) (500 mg in the morning and 1,000 mg in the evening),

97.3. Characteristics

97.3.1. Arm-level characteristics

97.3.1.	Arm-level characteristics					
Characteristic	ER metformin 1,500mg /day (N = 178)	ER metformin 1,500mg (AM/PM) (N = 182)	ER metformin 2,000 mg / day (N = 172)	1,500 mg immediate- release metformin (N = 174)		
Age (yrs)	54 (11.4)	54 (11.8)	55 (11.7)	54 (12.5)		
Mean (SD)						
≥65	n = 138 ; % = 77.5	n = 142 ; % = 78	n = 129 ; % = 75	n = 135 ; % = 77.6		
Sample size						
>= 65 Sample size	n = 40 ; % = 22.5	n = 40 ; % = 22	n = 43 ; % = 25	n = 39 ; % = 22.4		
Male	n = 83 ; % =					
Sample size	46.6	n = 111 ; % = 61	n = 91 ; % = 52.9	n = 95 ; % = 54.6		
Female	n = 95 ; % =					
Sample size	53.4	n = 71 ; % = 39	n = 81 ; % = 47.1	n = 79 ; % = 45.4		
Caucasian	n = 107 ; % =					
Sample size	60.1	n = 116 ; % = 63.7	n = 107 ; % = 62.2	n = 111; % = 63.8		
Black	n = 30 ; % =					
Sample size	16.9	n = 18; % = 9.9	n = 23 ; % = 13.4	n = 22 ; % = 12.6		
Asian	n = 5; % = 2.8	n = 5; % = 2.7	n = 3 ; % = 1.7	n = 3; % = 1.7		
Sample size						
Hispanic	n = 32 ; % = 18	n = 38 ; % = 20.9	n = 36 ; % = 20.9	n = 37 ; % = 21.3		
Sample size			20.0			
Native American	n = 1; % = 0.6	n = 3; % = 1.6	n = 1; % = 0.6	n = 0; % = 0		
Sample size						
Other Sample size	n = 3; % = 1.7	n = 2; % = 1.1	n = 2; % = 1.2	n = 1; % = 0.6		
Sample size	n = 60 · 0/ =					
≤30 less than 30	n = 60 ; % = 33.9	n = 67 ; % = 37	n = 53 ; % = 30.8	n = 58 ; % = 33.3		

Characteristic	ER metformin 1,500mg /day (N = 178)	ER metformin 1,500mg (AM/PM) (N = 182)	ER metformin 2,000 mg / day (N = 172)	1,500 mg immediate- release metformin (N = 174)
Sample size				
≤30 greater than or equal too eppi error	n = 117; % = 66.1	n = 114 ; % = 63	n = 119; % = 69.2	n = 116; % = 66.7
Sample size				
Duration of diabetes	3.9 (4.5)	4.5 (4.9)	3.9 (4.3)	4.4 (5.4)
Mean (SD)				
Drug naive	n = 81; % = 45.5	n = 86 ; % = 47.3	n = 84 ; % = 48.8	n = 87 ; % = 50
Sample size			10.0	
Metformin only Sample size	n = 43 ; % = 24.2	n = 44 ; % = 24.2	n = 45 ; % = 26.2	n = 43 ; % = 24.7
Sulfonylurea only	n = 29; % = 16.3	n = 30 ; % = 16.5	n = 22 ; % = 12.8	n = 30 ; % = 17.2
Sample size				
Metformin and sulfonylurea	n = 20 ; % = 11.2	n = 12 ; % = 6.6	n = 17 ; % = 9.9	n = 10; % = 5.7
Sample size				

98. Schweizer, 2007

Bibliographic Reference

Schweizer, A.; Couturier, A.; Foley, J. E.; Dejager, S.; Comparison between vildagliptin and metformin to sustain reductions in HbA1c over 1 year in drug-naive patients with Type 2 diabetes; Diabet Med; 2007; vol. 24 (no. 9); 955-61

Secondary publication of another included study- see primary study for details	No additional information.		
Other publications associated with this study included in review	No additional information.		
Trial name / registration number	NCT00099866.		
Study type	Randomised controlled trial (RCT)		
Study location	Multicentre trial (10 countries in the Americas and Europe).		
Study setting	Outpatient follow-up.		
Study dates	No additional information.		
Sources of funding	Funded by Novartis Pharmaceuticals Corporation.		
Inclusion criteria	People with type 2 diabetes mellitus who had an HbA1c of 7.5-11.0% while receiving no drug treatment; people who had previously taken no oral glucose lowering agents for more than three consecutive months at any time (considered drug naïve); male and female patients (non-fertile or of childbearing potential and using a medically approved birth control method); aged 18-78 years; fasting plasma glucose <15mmol/L.		
Exclusion criteria	History of type 1 diabetes or secondary forms of diabetes; acute metabolic diabetic complications within the past 6 months; congestive heart failure requiring pharmacological treatment, myocardial infarction, unstable angina or coronary artery bypass surgery within the previous 6 months; liver disease such as cirrhosis or chronic active hepatitis; renal disease or renal dysfunction suggested by elevated serum creatinine levels; ALT or AST greater than three times the upper limit of normal; direct bilirubin greater than 1.3 times the upper limit of normal; clinically significant abnormal TSH or fasting triglycerides >7.9 mmol/L.		

Recruitment / selection of participants	During the study, people were discontinued due to 'unsatisfactory therapeutic effect' if FPG >15mmol/L (or 13.3 mmol/L in Argentina) confirmed by a repeated measurement in the absence of intercurrent illness, or if they had symptoms of worsening hyperglycaemia in the absence of intercurrent illness or other incidental circumstances potentially causing deterioration of glucose control. People could also be withdrawn as a result of unsatisfactory therapeutic effect on the investigator's judgement alone.
Intervention(s)	Vildagliptin N=526
	Vildagliptin 100mg daily (given as equally divided doses).
	Concomitant therapy: No additional information.
Strata 1: People with type 2 diabetes mellitus and heart failure	People without heart failure
Strata 2: People with atherosclerotic cardiovascular diseases	
Strata 3: People with type 2 diabetes mellitus and chronic kidney disease	People without chronic kidney disease
Strata 4:	People at higher risk of developing cardiovascular disease
People with type 2 diabetes mellitus and high cardiovascular risk	Based on BMI, systolic blood pressure, triglycerides and presence of diabetes.
Subgroup 1: People with frailty	Not stated/unclear
Subgroup 2: Onset of type 2 diabetes mellitus	Not stated/unclear
Subgroup 3: People with non-alcoholic	People without non-alcoholic fatty liver disease

fatty liver disease	
Subgroup 4: People with obesity	Not stated/unclear
Subgroup 5: eGFR category at baseline	Not stated/unclear
Subgroup 6: Albuminuria category at baseline	Not stated/unclear
Sensitivity analysis category: Enrichment trial status	6) No response criteria
Population subgroups	No additional information.
Comparator	Metformin N=254 Metformin titrated to 2000mg daily (given as divided doses). Concomitant therapy: No additional information.
Number of participants	780
Duration of follow-up	52 weeks.
Indirectness	No additional information.
Method of analysis	ACA Other People who received at least one dose of the medication and had at least one post-baseline safety assessment
Additional comments	No additional information.

98.2.1. Vildagliptin (N = 526)

Vildagliptin 100mg daily (given as equally divided doses). Concomitant therapy: No additional information.

98.2.2. Metformin (N = 254)

Metformin titrated to 2000mg daily (given as divided doses). Concomitant therapy: No additional information.

98.3. Characteristics

98.3.1. Arm-level characteristics

30.3.1. Allii-level Characterist	.103	
Characteristic	Vildagliptin (N = 526)	Metformin (N = 254)
% Male	n = 278 ; % = 52.9	n = 146 ; % = 57.5
Sample size		
Mean age (SD) (years)	52.8 (11.7)	53.6 (10.2)
Mean (SD)		
Ethnicity	n = NA ; % = NA	n = NA ; % = NA
Sample size	0.57 0/ 07.0	
Caucasian Sample size	n = 357 ; % = 67.9	n = 177 ; % = 69.7
Hispanic or Latino	n = 104 ; % = 19.8	
Sample size	11 101, 70 10.0	n = 55 ; % = 21.7
Black	n = 42 ; % = 8	n = 13 ; % = 5.1
Sample size		
All Other	n = 23; % = 4.3	n = 9; % = 3.5
Sample size		
Comorbidities	n = NA ; % = NA	n = NA ; % = NA
Sample size	NIA - 0/ NIA	
Presence of frailty Sample size	n = NA ; % = NA	n = NA ; % = NA
Time since type 2 diabetes diagnosis	3.54	
(years)	J.J 4	3.28
IQR		
Time since type 2 diabetes diagnosis (years)	1.05 (NA to NA)	1.03 (NA to NA)
Median (IQR)		

Characteristic	Vildagliptin (N = 526)	Metformin (N = 254)
HbA1c (%)	8.7 (1.1)	8.7 (1.1)
Mean (SD)		
Blood pressure (mmHg)	NA (NA)	NA (NA)
Mean (SD)	400 (44)	
Systolic blood pressure Mean (SD)	133 (14)	133 (16)
Diastolic blood pressure	82 (8)	
·	<i>z</i> = (<i>z</i>)	82 (9)
Mean (SD)	>	
Heart rate	NR (NR)	NR (NR)
Mean (SD)		
Smoking status	n = NR ; % = NR	n = NR ; % = NR
Sample size		
Alcohol consumption Sample size	n = NR ; % = NR	n = NR ; % = NR
Presence of severe mental illness	n = NR ; % = NR	
Sample size	11 - IVIX , 70 - IVIX	n = NR ; % = NR
People with significant cognitive impairment	n = NR ; % = NR	n = NR ; % = NR
Sample size		
People with a learning disability	n = NR ; % = NR	n = NR ; % = NR
Sample size		
Weight	NR (NR)	NR (NR)
Mean (SD)		
BMI (kg/m2)	32.4 (5.7)	32.5 (5.7)
Mean (SD)		
Number of people with obesity	n = NA ; % = NA	n = NA ; % = NA
Sample size Cholostorol and linid lovels	NA (NA)	
Cholesterol and lipid levels	NA (NA)	NA (NA)
Mean (SD)		

Characteristic	Vilde alietie (N =	Mattausin (N -
Characteristic	Vildagliptin (N = 526)	Metformin (N = 254)
Total cholesterol	5.3 (1.1)	5.2 (1.1)
Mean (SD)		
HDL cholesterol	1.2 (0.2)	1.2 (0.3)
Mean (SD)		
LDL cholesterol Mean (SD)	3.1 (0.9)	3.1 (0.9)
,	2.4.(4.0)	
Triglycerides	2.4 (1.8)	2.4 (1.6)
Mean (SD)		
Albumin creatinine ratio	NR (NR)	NR (NR)
Mean (SD)		
eGFR (mL/min/1.73m2)	NR (NR)	NR (NR)
Mean (SD)		
Other antidiabetic medication used	NR (NR)	NR (NR)
Mean (SD)		
Blood pressure-lowering medication used	n = NR ; % = NR	n = NR ; % = NR
Sample size		
Statins/lipid-lowering medication used	n = NR ; % = NR	n = NR ; % = NR
Sample size		
Other treatment being received	n = NR ; % = NR	n = NR ; % = NR
Sample size		

99. Schweizer, 2009

Bibliographic Reference

Schweizer, A.; Dejager, S.; Bosi, E.; Comparison of vildagliptin and metformin monotherapy in elderly patients with type 2 diabetes: a 24-week, double-blind, randomized trial; Diabetes Obes Metab; 2009; vol. 11 (no. 8); 804-12

Secondary publication of another included study- see primary study for details	No additional information.
Other publications associated with this study included in review	No additional information.
Trial name / registration number	NCT00246619.
Study type	Randomised controlled trial (RCT)
Study location	Multicentre trial.
Study setting	Outpatient follow-up.
Study dates	14th September 2006 to 29th May 2008.
Sources of funding	Funded by Novartis Pharmaceutical Corporation.
Inclusion criteria	People with type 2 diabetes aged at least 65 years; HbA1c 7-9% at screening; people who were drug naïve (had taken no oral glucoselowering agents for at least 12 weeks prior to screening and no oral glucose-lowering agents for more than three consecutive months at any time in the past); FPG <15mmol/L; BMI 22-40kg/m2.
Exclusion criteria	History of type 1 or secondary forms of diabetes; acute metabolic diabetic complications within the past 6 months; congestive heart failure requiring pharmacological treatment or myocardial infarction, unstable angina or stroke or coronary artery bypass surgery within the past 6 months; liver disease such as cirrhosis or chronic active hepatitis; renal disease or renal dysfunction suggested by elevated serum creatinine levels.
Recruitment / selection of participants	No additional information.

Intervention(s)	Vildagliptin N=169
	Vildagliptin 100mg daily (as a once-daily dose) for 24 weeks of active treatment.
	Concomitant therapy: No additional information.
Strata 1: People with type 2 diabetes mellitus and heart failure	People without heart failure
Strata 2:	Mixed population
People with atherosclerotic cardiovascular diseases	33% of people had previous cardiac disorders
Strata 3:	Mixed population
People with type 2 diabetes mellitus and chronic kidney disease	Some with mild renal insufficiency defined by the study (60%)
Strata 4:	People at higher risk of developing cardiovascular disease
People with type 2 diabetes mellitus and high cardiovascular risk	Based on age, BMI, previous cardiac disease/CKD, dyslipidaemia (in 40%) and presence of diabetes.
Subgroup 1: People with frailty	Not stated/unclear
Subgroup 2: Onset of type 2 diabetes mellitus	Not stated/unclear
Subgroup 3: People with non-alcoholic fatty liver disease	People without non-alcoholic fatty liver disease
Subgroup 4: People with obesity	Not stated/unclear

Subgroup 5: eGFR category at baseline	Not stated/unclear
Subgroup 6: Albuminuria category at baseline	Not stated/unclear
Sensitivity analysis category: Enrichment trial status	6) No response criteria
Population subgroups	No additional information.
Comparator	Metformin N=166
	Metformin starting at 500mg/day titrated up, with weekly increases of 500mg to a maximum of 1500mg daily (as divided doses, 1000mg in the morning, 500mg in the evening) for 24 weeks of active treatment. Concomitant therapy: No additional information.
Number of participants	335
Duration of follow-up	24 weeks.
Indirectness	Outcome indirectness - the study reports acute coronary syndrome. This was included as a non-fatal myocardial infarction, but downgraded for indirectness.
Method of analysis	Other
anarysis	People who received at least one dose of the medication and had at least one post-baseline assessment
Additional comments	No additional information.

99.2.1. Vildagliptin (N = 169)

Vildagliptin 100mg daily (as a once-daily dose) for 24 weeks of active treatment. Concomitant therapy: No additional information.

99.2.2. Metformin (N = 166)

Metformin starting at 500mg/day titrated up, with weekly increases of 500mg to a maximum of 1500mg daily (as divided doses, 1000mg in the morning, 500mg in the

evening) for 24 weeks of active treatment. Concomitant therapy: No additional information.

99.3. Characteristics

99.3.1. Arm-level characteristics

Characteristic	Vildagliptin (N = 169)	Metformin (N = 166)
% Male	n = 75 ; % = 44.4	n = 88 ; % = 53
Sample size		
Mean age (SD) (years)	71.6 (5.2)	70.2 (5.1)
Mean (SD)		
Ethnicity	n = NA ; % = NA	n = NA ; % = NA
Sample size		
Caucasian	n = 123 ; % = 72.8	n = 117; % = 70.5
Sample size		
Asian	n = 32 ; % = 18.9	n = 36 ; % = 21.7
Sample size		
Hispanics Sample size	n = 13; % = 7.7	n = 10; % = 6
·	4 0/ 0.0	
All others Sample size	n = 1; % = 0.6	n = 3; % = 1.8
·	NIA O/ NIA	
Comorbidities Sample size	n = NA ; % = NA	n = NA ; % = NA
Presence of frailty	n = NA ; % = NA	
Sample size	, ,	n = NA ; % = NA
Time since type 2 diabetes diagnosis (years)	2.9 (4.2)	3 (4.7)
Mean (SD)		
HbA1c (%)	7.8 (0.6)	7.7 (0.6)
Mean (SD)		
Blood pressure	NR (NR)	NR (NR)

Characteristic	Vildagliptin (N = 169)	Metformin (N = 166)
Mean (SD)		
Heart rate	NR (NR)	NR (NR)
Mean (SD)		
Smoking status	n = NR ; % = NR	n = NR ; % = NR
Sample size	n = ND : 0/ = ND	
Alcohol consumption Sample size	n = NR ; % = NR	n = NR ; % = NR
Presence of severe mental illness	n = NR ; % = NR	
Sample size	11 - 141C, 70 - 141C	n = NR ; % = NR
People with significant cognitive impairment	n = NR ; % = NR	n = NR ; % = NR
Sample size		
People with a learning disability	n = NR ; % = NR	n = NR ; % = NR
Sample size	ND (ND)	
Weight Mean (SD)	NR (NR)	NR (NR)
BMI (kg/m2)	29.8 (4.4)	
Mean (SD)	29.0 (4.4)	29.4 (4.6)
Number of people with obesity	n = NR ; % = NR	
Sample size		n = NR ; % = NR
Cholesterol and lipid levels	NR (NR)	NR (NR)
Mean (SD)		
Albumin creatinine ratio	NR (NR)	NR (NR)
Mean (SD)	p = N(A + 0) = N(A)	
eGFR (mL/min/1.73m2) Sample size	n = NA ; % = NA	n = NA ; % = NA
>80 mL/min/1.73m2	n = 65; % = 38.5	
Sample size	11 - 00 , 70 - 00.0	n = 72; % = 43.4
50-80 mL/min/1.73m2	n = 102 ; % = 60.4	
Sample size		n = 90 ; % = 54.2

Characteristic	Vildagliptin (N = 169)	Metformin (N = 166)
30-49 mL/min/1.73m2	n = 2; % = 1.2	n = 4 ; % = 2.4
Sample size		
Other antidiabetic medication used	n = NA ; % = NA	n = NA ; % = NA
Sample size		
Blood pressure-lowering medication used	n = NA ; % = NA	n = NA ; % = NA
Sample size		
Statins/lipid-lowering medication used	n = NA ; % = NA	n = NA ; % = NA
Sample size		
Other treatment being received	n = NA ; % = NA	n = NA ; % = NA
Sample size		

100. Seino, 2018

Bibliographic Reference

Seino, Y.; Terauchi, Y.; Osonoi, T.; Yabe, D.; Abe, N.; Nishida, T.; Zacho, J.; Kaneko, S.; Safety and efficacy of semaglutide once weekly vs sitagliptin once daily, both as monotherapy in Japanese people with type 2 diabetes; Diabetes Obes Metab; 2018; vol. 20 (no. 2); 378-388

	-
Secondary publication of another included study- see primary study for details	NA
Other publications associated with this study included in review	NA
Trial name / registration number	NCT002254291
Study type	Randomised controlled trial (RCT)
Study location	Japan
Study setting	NR
Study dates	NR
Sources of funding	Novo Nordisk
Inclusion criteria	 Informed consent obtained before any trial-related activities. Trial-related activities are any procedures that are carried out as part of the trial, including activities to determine suitability for the trial Male or female, age ≥20 years at the time of signing informed consent HbA1c 6.5–9.5% (48–80 mmol/mol) (both inclusive) for subjects treated with oral antidiabetic drug (OAD) monotherapy and 7.0–10.5% (53–91 mmol/mol) (both inclusive) for subjects treated with diet and exercise at screening Japanese subjects with type 2 diabetes mellitus (diagnosed clinically) and on:

a) stable OAD monotherapy* in addition to diet and exercise therapy for at least 30 days prior to screening

OR

b) stable diet and exercise therapy for at least 30 days prior to screening

*Stable OAD monotherapy was defined as receiving half-maximum or below dose according to Japanese labelling for 30 days prior to screening. For metformin only, the maximum dose of 750 mg/day was allowed except for METGLUCO®. For METGLUCO®, the allowable half-maximum dose of 1125 mg/day was applied.

Exclusion criteria

- 1. Known or suspected hypersensitivity to trial product(s) or related products
- 2. Previous participation in this trial. Participation is defined as informed consent
- 3. Female who is pregnant, breast-feeding or intends to become pregnant or is of child-bearing potential and not using an adequate contraceptive method (e.g. abstinence [not having sex], diaphragm, condom [by the partner], intrauterine device, sponge, spermicide or oral contraceptives) throughout the trial including the 5-week follow-up period
- 4. Receipt of any investigational medicinal product within 90 days before screening
- 5. Treatment with glucose-lowering agent(s) other than stated in the inclusion criteria within 60 days before Visit 1 (week −2) and treatment with once-weekly glucagon-like peptide-1 (GLP-1) receptor agonists within 90 days before Visit 1 (week −2). An exception is short-term treatment (≤7 days in total) with insulin in connection with inter-current illness
- 6. Any disorder which, in the opinion of the investigator, might jeopardise subject's safety or compliance with the protocol
- 7. History of chronic or idiopathic acute pancreatitis
- 8. Screening calcitonin value ≥50 ng/L
- 9. Personal or family history of medullary thyroid carcinoma (MTC) or multiple endocrine neoplasia syndrome type 2 (MEN2)
- 10. Impaired renal function defined as estimated glomerular filtration rate (eGFR) <30 mL/min/1.73 m2 per Modification of Diet in Renal Disease (MDRD) formula (4 variable version)
- 11. Acute coronary or cerebrovascular event within 90 days before randomisation (Visit 2 [week 0])
- 12. Heart failure, New York Heart Association (NYHA) class IV

	13. Known proliferative retinopathy or maculopathy requiring acute treatment according to the opinion of the investigator
	14. Diagnosis of malignant neoplasm in the previous 5 years (except basal cell skin cancer or squamous cell skin cancer)
	15. Mental inability, unwillingness or language barrier precluding adequate understanding of or compliance with trial procedures
Recruitment / selection of participants	NR
Intervention(s)	S.C. Semaglutide 0.5 mg (n=103)
	Participants received 0.5 mg S.C. semaglutide once weekly for 30 weeks. Participants followed a fixed dose-escalation regimen of semaglutide 0.5 mg (maintenance dose reached after 4 weeks of 0.25 mg semaglutide once weekly)
	S.C. Semaglutide 1.0 mg (n=102)
	Participants received 1.0 mg S.C. semaglutide once weekly for 30 weeks. Participants followed a fixed dose-escalation regimen of semaglutide 1.0 mg (maintenance dose reached after 4 weeks of 0.25 mg semaglutide once weekly, followed by 4 weeks of 0.5mg semaglutide)
	, , , , , , , , , , , , , , , , , , , ,
Cointervention	
Cointervention Strata 1: People with type 2 diabetes mellitus and heart failure	
Strata 1: People with type 2 diabetes mellitus and	None People without heart failure Not stated/unclear
Strata 1: People with type 2 diabetes mellitus and heart failure Strata 2: People with atherosclerotic cardiovascular	None People without heart failure Not stated/unclear
Strata 1: People with type 2 diabetes mellitus and heart failure Strata 2: People with atherosclerotic cardiovascular diseases Strata 3: People with type 2 diabetes mellitus and chronic kidney	None People without heart failure Not stated/unclear

cardiovascular	
risk	
Subgroup 1: Not People with frailty	ot stated/unclear
Subgroup 2: Not Onset of type 2 diabetes mellitus	ot stated/unclear
Subgroup 3: Not People with non-alcoholic fatty liver disease	ot stated/unclear
Subgroup 4: Not People with obesity	ot stated/unclear
Subgroup 5: eG eGFR category at baseline	GFR ≥30mL/min/1.73m2
Subgroup 6: Not Albuminuria category at baseline	ot stated/unclear
Sensitivity 7) Nanalysis category: Enrichment trial status	Mixed population
Population NR subgroups	3
·	raglitpin 100mg (n=103) articipants received 100mg once daily for 30 weeks
Number of participants 308	8
Duration of 35 follow-up	weeks
Indirectness NA	A
Method of AC analysis	CA
Additional All comments	data analysed using all case analyses

100.2.1. Semaglutide 0.5 mg (N = 103)

Participants received 0.5mg S.C. semaglutide once weekly for 30 weeks

100.2.2. Semaglutide 1.0 mg (N = 102)

Participants received 1.0mg S.C. semaglutide once weekly for 30 weeks

100.2.3. Sitagliptin (N = 103)

Participants received once daily 100mg sitagliptin for 30 weeks

100.3. Characteristics

100.3.1. Arm-level characteristics

Characteristic	Semaglutide 0.5 mg (N = 103)	Semaglutide 1.0 mg (N = 102)	Sitagliptin (N = 103)
% Male	n = 79; % = 76.7	n = 75 ; % = 73.5	n = 81 ; % =
Sample size			78.6
Mean age (SD)	58.8 (10.4)	58.1 (11.6)	57.9 (10.1)
Mean (SD)			
Ethnicity	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
Sample size			INIX
Comorbidities	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
Sample size			1417
Presence of frailty	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
Sample size			INIX
Time since type 2 diabetes diagnosis (Years (mean, SD))	8 (5.2)	7.8 (6.9)	8.1 (6.7)
Mean (SD)			
Smoking status	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % =
Sample size			NR
Alcohol consumption	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % =
Sample size			NR

Characteristic	Semaglutide 0.5 mg (N = 103)	Semaglutide 1.0 mg (N = 102)	Sitagliptin (N = 103)
Presence of severe mental illness	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
Sample size			
People with significant congitive impairment	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
Sample size			
People with a learning disability	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
Sample size			
Number of people with obesity	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % =
Sample size			NR
Other treatment being received	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
Sample size			

101. Shihara, 2011

Bibliographic Reference

Shihara, Nobuyuki; Kitaoka, Masafumi; Inagaki, Nobuya; Kadowaki, Takashi; Koumoto, Seisuke; Satoh, Jo; Terauchi, Yasuo; Nunoi, Kiyohide; Yamada, Yuichiro; Sakamaki, Hiroyuki; Seino, Yutaka; Randomized controlled trial of single-agent glimepiride and pioglitazone in Japanese patients with type 2 diabetes: A comparative study.; Journal of diabetes investigation; 2011; vol. 2 (no. 5); 391-8

Secondary publication of another included study- see primary study for details	No additional information
Other publications associated with this study included in review	No additional information
Trial name / registration number	University Hospital Medical Information Network: UMIN000004582
Study type	Randomised controlled trial (RCT)
Study location	Japan
Study setting	No additional information
Study dates	August 2007 - February 2010
Sources of funding	Funded by Sanofi Aventis
Inclusion criteria	Outpatients of either sex with type 2 diabetes mellitus Aged 30–75 years Committed to a stable dietary and exercise regimen for >1 month before randomization HbA1c 6.9-10.4% 1 month before and at randomization, with a difference <1% between measurements
Exclusion criteria	Type 1 diabetes mellitus Use of insulin or any oral hypoglycaemic agent (including an alpha-glucosidase inhibitor) in the month before randomization

	Heart failure or history of heart failure
	Any serious intercurrent complication involving the heart, kidney, liver, pancreas or other organs, or hematological condition
Recruitment / selection of participants	No additional information
Intervention(s)	Participants allocated to the intervention initially received 0.5 mg glimepiride if their HbA1c was 6.9-7.4%, or 1.0 mg per day if their HbA1c was 7.4-10.4%. The dose could be increased to a maximum of 6 mg/day in order to achieve morning fasting blood glucose of<120 mg/dL. The dosage could be decreased according to the supervising physician's judgment if morning fasting blood glucose was <80 mg/dL. Drug doses were titrated according to morning fasting blood glucose measured at scheduled clinic visits.
Cointervention	
Strata 1: People with type 2 diabetes mellitus and heart failure	People without heart failure
Strata 2: People with atherosclerotic cardiovascular diseases	People without other cardiovascular diseases
Strata 3: People with type 2 diabetes mellitus and chronic kidney disease	People without chronic kidney disease
Strata 4: People with type 2 diabetes mellitus and high cardiovascular risk	Not stated/unclear
Subgroup 1: People with frailty	Not stated/unclear
Subgroup 2: Onset of type 2 diabetes mellitus	Not stated/unclear

Subgroup 3: People with non-alcoholic fatty liver disease Subgroup 4: People with obesity Subgroup 5: GefFR category at baseline Subgroup 6: Albuminuria category at baseline Sensitivity analysis category: Enrichment trial status Population subgroups Comparator Population subgroups Comparator No additional information subgroups Participants allocated to the comparator initially received 15 mg pioglitazone per day, which could be increased to a maximum of 45 and 30 mg/day in men and women, respectively, in order to achieve morning blood glucose of <120 mg/dL. The dosage could be decreased according to the supervising physician's judgment if morning fasting blood glucose was <80 mg/dL. Drug doses were titrated according to morning fasting blood glucose measured at scheduled clinic visits. Number of participants 191 randomised 95 received glimepiride, 86 completed 96 received pioglitazone, 91 completed Duration of follow-up Indirectness Method of analysis Additional comments		
People with obesity Subgroup 5: eGFR category at baseline Subgroup 6: Albuminuria category at baseline Sensitivity analysis category: Enrichment trial status Population subgroups Comparator Participants allocated to the comparator initially received 15 mg pioglitazone per day, which could be increased to a maximum of 45 and 30 mg/day in men and women, respectively, in order to achieve morning blood glucose of <120 mg/dL. The dosage could be decreased according to the supervising physician's judgment if morning fasting blood glucose was <80 mg/dL. Drug doses were titrated according to morning fasting blood glucose measured at scheduled clinic visits. Number of participants Number of participants Participants 191 randomised 95 received glimepiride, 86 completed 96 received pioglitazone, 91 completed 6 months follow-up Indirectness None Method of analysis Additional None	People with non-alcoholic fatty liver	People without non-alcoholic fatty liver disease
eGFR category at baseline Subgroup 6: Albuminuria category at baseline Sensitivity analysis category: Enrichment trial status Population subgroups Comparator Participants allocated to the comparator initially received 15 mg pioglitazone per day, which could be increased to a maximum of 45 and 30 mg/day in men and women, respectively, in order to achieve morning blood glucose of <120 mg/dL. The dosage could be decreased according to the supervising physician's judgment if morning fasting blood glucose was <80 mg/dL. Drug doses were titrated according to morning fasting blood glucose measured at scheduled clinic visits. Number of participants Participants 191 randomised 95 received glimepiride, 86 completed 96 received pioglitazone, 91 completed Duration of follow-up Indirectness None Method of analysis Additional None	People with	Not stated/unclear
Albuminuria category at baseline Sensitivity analysis category: Enrichment trial status Population subgroups Comparator Participants allocated to the comparator initially received 15 mg pioglitazone per day, which could be increased to a maximum of 45 and 30 mg/day in men and women, respectively, in order to achieve morning blood glucose of <120 mg/dL. The dosage could be decreased according to the supervising physician's judgment if morning fasting blood glucose was <80 mg/dL. Drug doses were titrated according to morning fasting blood glucose measured at scheduled clinic visits. Number of participants 191 randomised 95 received glimepiride, 86 completed 96 received pioglitazone, 91 completed Duration of follow-up Indirectness None Method of analysis Additional None	eGFR category	
analysis category: Enrichment trial status Population subgroups Comparator Participants allocated to the comparator initially received 15 mg pioglitazone per day, which could be increased to a maximum of 45 and 30 mg/day in men and women, respectively, in order to achieve morning blood glucose of <120 mg/dL. The dosage could be decreased according to the supervising physician's judgment if morning fasting blood glucose was <80 mg/dL. Drug doses were titrated according to morning fasting blood glucose measured at scheduled clinic visits. Number of participants 191 randomised 95 received glimepiride, 86 completed 96 received pioglitazone, 91 completed Duration of follow-up Indirectness None Method of analysis Additional None	Albuminuria category at	Not stated/unclear
Subgroups Comparator Participants allocated to the comparator initially received 15 mg pioglitazone per day, which could be increased to a maximum of 45 and 30 mg/day in men and women, respectively, in order to achieve morning blood glucose of <120 mg/dL. The dosage could be decreased according to the supervising physician's judgment if morning fasting blood glucose was <80 mg/dL. Drug doses were titrated according to morning fasting blood glucose measured at scheduled clinic visits. Number of participants 191 randomised 95 received glimepiride, 86 completed 96 received pioglitazone, 91 completed Duration of follow-up Indirectness None Method of analysis Additional None	analysis category: Enrichment	8) Not reported
pioglitazone per day, which could be increased to a maximum of 45 and 30 mg/day in men and women, respectively, in order to achieve morning blood glucose of <120 mg/dL. The dosage could be decreased according to the supervising physician's judgment if morning fasting blood glucose was <80 mg/dL. Drug doses were titrated according to morning fasting blood glucose measured at scheduled clinic visits. Number of participants 191 randomised 95 received glimepiride, 86 completed 96 received pioglitazone, 91 completed Duration of follow-up Indirectness None Method of analysis Additional None	•	No additional information
participants 95 received glimepiride, 86 completed 96 received pioglitazone, 91 completed Duration of follow-up Indirectness None Method of analysis Additional None	Comparator	pioglitazone per day, which could be increased to a maximum of 45 and 30 mg/day in men and women, respectively, in order to achieve morning blood glucose of <120 mg/dL. The dosage could be decreased according to the supervising physician's judgment if morning fasting blood glucose was <80 mg/dL. Drug doses were titrated according to morning fasting
follow-up Indirectness None Method of analysis Additional None		95 received glimepiride, 86 completed
Method of analysis Additional None		6 months
analysis Additional None	Indirectness	None
		ІТТ
		None

101.2.1. Glimepiride (N = 95)

Initially 0.5 (HbA1c 6.9-7.4%) or 1.0 mg (7.4-10.4%) per day up to a maximum of 6 mg per day

101.2.2. Pioglitazone (N = 96)

Initially 15 mg per day, up to a maximum of 45 (men) and 30 mg (women) per day

101.3. Characteristics

101.3.1. Arm-level characteristics

Characteristic Glimepiride (N = 95) Pioglitazone (N = 96) % Male n = 62; % = 65 n = 65; % = 68 Sample size 57.7 (10.4) 56.8 (10.3) Mean age (SD) (years) NR NR Nominal NR NR Name since type 2 diabetes diagnosis (years) 4.1 (4.3) Data for 41 and 52 participants 7.8 (0.9) Mean (SD) 7.8 (0.9)
Sample size Mean age (SD) (years) Mean (SD) Ethnicity NR Nominal Comorbidities NR Nominal Presence of frailty NR Nominal Time since type 2 diabetes diagnosis (years) Data for 41 and 52 participants Mean (SD) HbA1c (%) Data for 95 and 95 participants 57.7 (10.4) 56.8 (10.3) NR NR NR NR NR NR NR A.1 (4.3) 7.8 (0.9) 7.8 (0.9)
Mean age (SD) (years)57.7 (10.4)56.8 (10.3)Mean (SD)NRNRNominalNRNRNominalNRNRNominalNRNRNominalNRNRTime since type 2 diabetes diagnosis (years) Data for 41 and 52 participants6 (8.2)4.1 (4.3)Mean (SD)T.8 (0.9)7.8 (0.9)HbA1c (%) Data for 95 and 95 participants7.8 (0.9)
Mean (SD) Ethnicity NR Nominal Comorbidities NR Nominal Presence of frailty Nominal Time since type 2 diabetes diagnosis (years) Data for 41 and 52 participants Mean (SD) HbA1c (%) Data for 95 and 95 participants SUR NR NR NR NR A1 (4.3) 4.1 (4.3) 7.8 (0.9) 7.8 (0.9)
Ethnicity Nominal Comorbidities NR NR NR Nominal Presence of frailty NR Nominal Time since type 2 diabetes diagnosis (years) Data for 41 and 52 participants Mean (SD) HbA1c (%) Data for 95 and 95 participants NR NR NR NR NR NR NR NR NR N
Nominal Comorbidities NR Nominal Presence of frailty NR NR NR NR NR NR NR NR NR N
Comorbidities NR Nominal Presence of frailty NR Nominal Time since type 2 diabetes diagnosis (years) Data for 41 and 52 participants Mean (SD) HbA1c (%) Data for 95 and 95 participants NR NR NR NR NR NR NR NR NR N
NR Nominal Presence of frailty NR NR Nominal Time since type 2 diabetes diagnosis (years) Data for 41 and 52 participants Mean (SD) HbA1c (%) Data for 95 and 95 participants NR NR NR NR NR NR NR NR NR N
Presence of frailty NR Nominal Time since type 2 diabetes diagnosis (years) Data for 41 and 52 participants Mean (SD) HbA1c (%) Data for 95 and 95 participants NR 4.1 (4.3) 7.8 (0.9) 7.8 (0.9)
Nominal Time since type 2 diabetes diagnosis (years) Data for 41 and 52 participants Mean (SD) HbA1c (%) Data for 95 and 95 participants NR 4.1 (4.3) 7.8 (0.9)
Time since type 2 diabetes diagnosis (years) Data for 41 and 52 participants Mean (SD) HbA1c (%) Data for 95 and 95 participants 6 (8.2) 4.1 (4.3) 7.8 (0.9)
(years) Data for 41 and 52 participants Mean (SD) HbA1c (%) Data for 95 and 95 participants 7.8 (0.9) 7.8 (0.9)
HbA1c (%) Data for 95 and 95 participants 7.8 (0.9) 7.8 (0.9)
Data for 95 and 95 participants 7.8 (0.9)
Mean (SD)
Blood pressure NR NR Nominal
Heart rate NR NR Nominal
Smoking status NR Nominal
Alcohol consumption NR NR

Characteristic	Glimepiride (N = 95)	Pioglitazone (N = 96)
Nominal		
Presence of severe mental illness	NR	NR
Nominal		
People with significant cognitive impairment	NR	NR
Nominal		
People with a learning disability Nominal	NR	NR
Weight (kg)	65.6 (12.5)	
Data for 93 and 92 participants Mean (SD)	03.0 (12.0)	65.5 (14.6)
BMI (kg/m²)	24.6 (3.8)	
Data for 93 and 92 participants	24.0 (0.0)	24.5 (4.3)
Mean (SD)	ND	
Number of people with obesity	NR	NR
Nominal		
Cholesterol and lipid levels	NA (NA)	NA (NA)
Mean (SD)	007.5 (00.4)	
TC Mean (SD)	207.5 (39.1)	205.5 (38.2)
LDL	126.5 (36.5)	
	0.0 (00.0)	123.2 (32.6)
Mean (SD)		
HDL	59.3 (23)	52.8 (13.7)
Mean (SD)		, ,
TG	129.8 (68.4)	164 (112.4)
Mean (SD)		
Albumin creatinine ratio	NR	NR
Nominal		
eGFR (mL/min/1.73m2)	NR	NR
Nominal		

Characteristic	Glimepiride (N = 95)	Pioglitazone (N = 96)
Other antidiabetic medication used Nominal	NR	NR
Blood pressure-lowering medication used	NR	NR
Nominal		
Statins/lipid-lowering medication used	NR	NR
Nominal		
Other treatment being received	NR	NR
Nominal		

102. Sorli, 2017

Bibliographic Reference

Sorli, C.; Harashima, S. I.; Tsoukas, G. M.; Unger, J.; Karsbol, J. D.; Hansen, T.; Bain, S. C.; Efficacy and safety of once-weekly semaglutide monotherapy versus placebo in patients with type 2 diabetes (SUSTAIN 1): a double-blind, randomised, placebo-controlled, parallel-group, multinational, multicentre phase 3a trial; Lancet Diabetes Endocrinol; 2017; vol. 5 (no. 4); 251-260

Secondary publication of another included study- see primary study for details	NA	
Other publications associated with this study included in review	None	
Trial name / registration number	SUSTAIN 1/NCT02054897	
Study type	Randomised controlled trial (RCT)	
Study location	International (72 sites in Canada, Italy, Japan, Mexico, Russia, South Africa, UK and USA)	
Study setting	Community	
Study dates	02/2014 to 08/2014	
Sources of funding	Funded directly by Novo Nordisk A/S, Denmark.	
Inclusion criteria	 Aged ≥18 years (≥20 years for Japan) Diagnosis of type 2 diabetes Management with diet and exercise only ≥30 days before screening when enrolled HbA1c level 7·0-10·0% inclusive. 	
Exclusion criteria	 Female that is pregnant, breastfeeding, or intends to get pregnant, or is of child-bearing potential and not using adequate contraception as required by local practice, both during trial and 5-week FU period Use of glucose-lowering drugs ≤90 days before screening (except for ≤7 days treatment with insulin) History of chronic or idiopathic acute pancreatitis 	

	 Personal or family history of medullary thyroid carcinoma or multiple endocrine neoplasia syndrome type 2 Impaired renal function (eGFR <30 mL/min per 1·73 m²) Screening calcitonin values of at least 50 ng/L (pg/mL) Heart failure (New York Heart Association class IV) Any acute coronary or cerebrovascular events in the 90 days before randomisation
Recruitment / selection of participants	Participants recruited from 33 sites across 6 countries with some recruited using advertisements at some sites. Random assignment 2:2:1:1 to semaglutide 1mg, semaglutide 0.5 mg, placebo for semaglutide 1mg, or placebo for semaglutide 0.5 mg, using automated voice/web recognition system with no human involvement.
Intervention(s)	 Semaglutide 1.0 mg once weekly Semaglutide 0.5 mg once weekly Subcutaneous injection of semaglutide in prefilled 1.5 mL PDS290 peninjector. Participants administered own injections and encourage to administer them on same day each week in same body area (thigh, abdomen or upper arm). Time of day and proximity of meal times not specified. Participants in semaglutide 0.5 mg group reached maintenance dose after 4 weeks of 0.25 mg once weekly. Participants in semaglutide 1.0 mg group, maintenance dose reached after 4 weeks of 0.25 mg, followed by 4 weeks of 0.5 mg. Participants with unacceptable hyperglycaemia (assessed by FPG) could be offered rescue medication at discretion of investigator (ADA and EASD guidelines), either metformin as first choice or other antidiabetic drugs (except GLP-1RA and DPP-4 inhibitors) as add-ons to study treatment.
Strata 1: People with type 2 diabetes mellitus and heart failure	Not stated/unclear Exclusion criteria: heart failure (NYHA class IV). Trial may therefore include participants with NYHA class II and III.
Strata 2: People with atherosclerotic cardiovascular diseases	Not stated/unclear Exclusion criteria: Any acute coronary or cerebrovascular events in the 90 days before randomisation. Treial may therefore include participants with other CV diseases.
Strata 3: People with type 2 diabetes mellitus and chronic kidney disease	Not stated/unclear
Strata 4: People with type 2 diabetes mellitus and	Not stated/unclear

high cardiovascular risk	
Subgroup 1: People with frailty	Not stated/unclear
Subgroup 2: Onset of type 2 diabetes mellitus	Not stated/unclear
Subgroup 3: People with non-alcoholic fatty liver disease	Not stated/unclear
Subgroup 4: People with obesity	Not stated/unclear
Subgroup 5: eGFR category	eGFR ≥30mL/min/1.73m2
at baseline	Exclusion criteria: eGFR<30 mL/min per 1.73 m2
Subgroup 6: Albuminuria category at baseline	Not stated/unclear
Sensitivity analysis	7) Mixed population
category: Enrichment trial status	Inclusion criteria: Management with diet and exercise only ≥30 days before screening when enrolled
Population subgroups	
Comparator	 Placebo to semaglutide 1.0 mg once weekly Placebo to semaglutide 0.5 mg once weekly
	Volume-matched placebo injection provided in prefilled 1.5 mL PDS290 pen-injector identical in appearance, taste and smell to those used for semaglutide arms. See intervention details for dose-matching escalation. Participants with unacceptable hyperglycaemia could be offered rescue medication at discretion of investigator (ADA and EASD guidelines), either metformin as first choice or other antidiabetic drugs (except GLP-1RA and DPP-4 inhibitors) as add-ons to study treatment.
Number of participants	N=388
Duration of follow-up	30 weeks
Method of analysis	Modified ITT

Efficacy and safety analysis with all randomised participants who took at least one dose of study drug or placebo. Efficacy analysis used data before initiation of any rescue medication or before premature treatment discontinuation. Safety analysis used data before premature treatment discontinuation with window of 42 days to identify treatment-emergent AEs. Sensitivity analysis used all data regardless of whether data obtained while participants discontinued treatment or whether participant given rescue medication.

102.2. Study arms

102.2.1. Semaglutide 1 mg once weekly (N = 130)

Subcutaneous injection of semaglutide 1 mg once weekly for 30 weeks.

102.2.2. Semaglutide 0.5 mg once weekly (N = 129)

Subcutaneous injection of semaglutide 0.5 mg once weekly for 30 weeks.

102.2.3. Placebo once weekly (N = 129)

Subcutaneous volume-matched placebo injection (1 mg or 0.5 mg) once weekly for 30 weeks.

102.3. Characteristics

102.3.1. Arm-level characteristics

Characteristic	Semaglutide 1 mg once weekly (N = 130)	Semaglutide 0.5 mg once weekly (N = 129)	Placebo once weekly (N = 129)
% Male Sample size	n = 80 ; % = 62	n = 60 ; % = 47	n = 70 ; % = 54
Sample Size			
Mean age (SD) (years)	52.7 (11.9)	54.6 (11.1)	53.9 (11)
Mean (SD)			
Ethnicity	n = NA ; % = NA	n = NA ; % = NA	n = NA ; % = NA
Sample size			
Asian	n = 25 ; % = 19	n = 26 ; % = 20	n = 32 ; % = 25
Sample size			
Black/African American	n = 11; % = 8	n = 11 ; % = 9	n = 9; % = 7

Characteristic	Semaglutide 1 mg once weekly (N = 130)	Semaglutide 0.5 mg once weekly (N = 129)	Placebo once weekly (N = 129)
Sample size			
Hispanic or Latino	n = 45 ; % = 35	n = 34 ; % = 27	n = 36 ; % = 28
Sample size			
Not hispanic or latino	n = 85; % = 65	n = 94 ; % = 73	n = 93 ; % = 72
Sample size	00 · 0/ - 00		
White	n = 88 ; % = 68	n = 83; % = 65	n = 78 ; % = 60
Sample size			
Time since type 2 diabetes diagnosis (years)	3.62 (4.88)	4.81 (6.1)	4.06 (5.48)
Mean (SD)			
HbA1c (%)	8.12 (0.81)	8.09 (0.89)	7.95 (0.85)
Mean (SD)			
Weight (kg)	96.87 (25.59)	89.81 (22.96)	89.05 (22.16)
Mean (SD)			
BMI (kg/m2)	33.92 (8.43)	32.46 (7.62)	32.4 (6.86)
Mean (SD)			
eGFR (mL/min/1.73m2)	100.9 (27.74)	95.91 (26.23)	100.2 (24.97)
Mean (SD)			

103. Stenlöf, 2013

Bibliographic Reference

Stenlöf, K.; Cefalu, W. T.; Kim, K. A.; Alba, M.; Usiskin, K.; Tong, C.; Canovatchel, W.; Meininger, G.; Efficacy and safety of canagliflozin monotherapy in subjects with type 2 diabetes mellitus inadequately controlled with diet and exercise; Diabetes Obes Metab; 2013; vol. 15 (no. 4); 372-82

duy details		
No information available.		
No information available.		
NCT01081834		
Randomised controlled trial (RCT)		
17 countries including: US Sweden South Korea		
Hospital		
03/2010 to 03/2012		
Janssen Global Services, LLC. Canagliflozin is being developed by Janssen Research & Development, LLC, in collaboration with Mitsubishi Tanabe Pharma Corporation.		
 Men and women 18–80 years of age with T2DM who met one of the two following criteria: (i) not on an antihyperglycaemic agent at screening with HbA1c ≥7.0 and ≤10.0% or (ii) on 		

antihyperglycaemic agent monotherapy [except peroxisome proliferator-activated receptor-⟨PPAR⟩) agonist] or metformin plus sulfonylurea combination therapy (at \$50% of maximally or near-maximally effective doses) with HbA1c ≥6.5 and ≤9.5% at screening and HbA1c ≥7.0 and ≤10.0% and fasting plasma glucose <15.0 mmol/l at week −2. Exclusion criteria • Subjects were excluded if they had repeated Fasting Plasma Glucose measurements ≥15.0 mmol/l during the pre-treatment phase (or >19.4 mmol/l for the high glycaemic sub study) • A history of type 1 diabetes, hereditary glucose-galactose malabsorption, primary renal glucosuria or cardiovascular (CV) disease (including myocardia) infarction, unstable angina, revascularisation procedure or cerebrovascular accident) • Treatment with a PPARy agonist, insulin, another SGLT2 inhibitor or any other antihyperglycaemic agents except as specified in the inclusion criteria within 12 weeks before screening • Estimated glomerular filtration rate (eGFR) <50 ml/min/1.73 m2 at screening • Estimated glomerular filtration rate (eGFR) <50 ml/min/1.73 m2 at screening • Estimated glomerular filtration rate (eGFR) <50 ml/min/1.73 m2 at screening. • Estimated glomerular filtration rate (eGFR) <50 ml/min/1.73 m2 at screening. • Estimated glomerular filtration rate (eGFR) <50 ml/min/1.73 m2 at screening. • Estimated glomerular filtration rate (eGFR) <50 ml/min/1.73 m2 at screening. • Estimated glomerular filtration rate (eGFR) <50 ml/min/1.73 m2 at screening. • Estimated glomerular filtration rate (eGFR) <50 ml/min/1.73 m2 at screening. • Estimated glomerular filtration rate (eGFR) <50 ml/min/1.73 m2 at screening. • Estimated glomerular filtration rate (eGFR) <50 ml/min/1.73 m2 at screening. • Estimated glomerular filtration rate (eGFR) <50 ml/min/1.73 m2 at screening. • Estimated glomerular filtration rate (eGFR) <50 ml/min/1.73 m2 at screening. • Estimated glomerular filtration rate (eGFR) <50 ml/min/1.73 m2 at screening. • Estimated glomerular filtration rate (eGFR) <50 ml/min/1.73 m2 a		
Recruitment / selection of participants Men and women aged 18 - 80 years were recruited from 17 countries. The study included both subjects with inadequate control on diet and exercise and subjects on anantihyperglycaemic agent, who underwent a washout of the agent. Subjects not on an antihyperglycaemic directly entered a 2-week, single-blind, placebo run-in period (week -2 to day 1), while subjects on an antihyperglycaemic underwent an 8-week, antihyperglycaemic washout/diet and exercise period followed by the placebo run-in period. Intervention(s) Canagliflozin 300 mg daily, taken orally Canagliflozin 100 mg daily, taken orally During the double-blind treatment period, glycaemic rescue therapy with metformin was initiated if Fasting Plasma Glucose >15.0 mmol/l after day 1 to week 6, >13.3 mmol/l after week 6 to week 12 and >11.1 mmol/l after week 12 to week 26. Cointervention Strata 1: People with type 2 diabetes mellitus and heart failure Strata 2: People with attherosclerotic cardiovascular People with attherosclerotic cardiovascular		 proliferator-activated receptor-γ (PPARγ) agonist] or metformin plus sulfonylurea combination therapy (at ≤50% of maximally or near-maximally effective doses) with HbA1c ≥6.5 and ≤9.5% at screening and HbA1c ≥7.0 and ≤10.0% and fasting plasma glucose <15.0 mmol/l at week −2. Subjects were excluded if they had repeated Fasting Plasma Glucose measurements >15.0 mmol/l during the pre-treatment phase (or >19.4 mmol/l for the high glycaemic sub study) A history of type 1 diabetes, hereditary glucose-galactose malabsorption, primary renal glucosuria or cardiovascular (CV) disease (including myocardial infarction, unstable angina, revascularisation procedure or cerebrovascular accident) Treatment with a PPARγ agonist, insulin, another SGLT2 inhibitor or any other antihyperglycaemic agents except as specified in the inclusion criteria within 12 weeks before screening
selection of participants study included both subjects with inadequate control on diet and exercise and subjects on anantihyperglycaemic agent, who underwent a washout of the agent. Subjects not on an antihyperglycaemic directly entered a 2-week, single-blind, placebo run-in period (week -2 to day 1), while subjects on an antihyperglycaemic underwent an 8-week, antihyperglycaemic washout/diet and exercise period followed by the placebo run-in period. Intervention(s) Canagliflozin 300 mg daily, taken orally Canagliflozin 100 mg daily, taken orally During the double-blind treatment period, glycaemic rescue therapy with metformin was initiated if Fasting Plasma Glucose >15.0 mmol/l after day 1 to week 6, >13.3 mmol/l after week 6 to week 12 and >11.1 mmol/l after week 12 to week 26. Cointervention Strata 1: People with type 2 diabetes mellitus and heart failure Strata 2: People with atherosclerotic cardiovascular		· · · · · · · · · · · · · · · · · · ·
selection of participants study included both subjects with inadequate control on diet and exercise and subjects on anantihyperglycaemic agent, who underwent a washout of the agent. Subjects not on an antihyperglycaemic directly entered a 2-week, single-blind, placebo run-in period (week -2 to day 1), while subjects on an antihyperglycaemic underwent an 8-week, antihyperglycaemic washout/diet and exercise period followed by the placebo run-in period. Intervention(s) Canagliflozin 300 mg daily, taken orally Canagliflozin 100 mg daily, taken orally During the double-blind treatment period, glycaemic rescue therapy with metformin was initiated if Fasting Plasma Glucose >15.0 mmol/l after day 1 to week 6, >13.3 mmol/l after week 6 to week 12 and >11.1 mmol/l after week 12 to week 26. Cointervention Strata 1: People with type 2 diabetes mellitus and heart failure Strata 2: People with atherosclerotic cardiovascular		
Canagliflozin 100 mg daily, taken orally During the double-blind treatment period, glycaemic rescue therapy with metformin was initiated if Fasting Plasma Glucose >15.0 mmol/l after day 1 to week 6, >13.3 mmol/l after week 6 to week 12 and >11.1 mmol/l after week 12 to week 26. Cointervention None. Strata 1: People with type 2 diabetes mellitus and heart failure Strata 2: People with atherosclerotic cardiovascular	selection of	study included both subjects with inadequate control on diet and exercise and subjects on anantihyperglycaemic agent, who underwent a washout of the agent. Subjects not on an antihyperglycaemic directly entered a 2-week, single-blind, placebo run-in period (week -2 to day 1), while subjects on an antihyperglycaemic underwent an 8-week, antihyperglycaemic washout/diet and exercise period followed by the
Canagliflozin 100 mg daily, taken orally During the double-blind treatment period, glycaemic rescue therapy with metformin was initiated if Fasting Plasma Glucose >15.0 mmol/l after day 1 to week 6, >13.3 mmol/l after week 6 to week 12 and >11.1 mmol/l after week 12 to week 26. Cointervention None. Strata 1: People with type 2 diabetes mellitus and heart failure Strata 2: People with atherosclerotic cardiovascular	Intervention(s)	Canagliflozin 300 mg daily, taken orally
metformin was initiated if Fasting Plasma Glucose >15.0 mmol/l after day 1 to week 6, >13.3 mmol/l after week 6 to week 12 and >11.1 mmol/l after week 12 to week 26. Cointervention Strata 1: People with type 2 diabetes mellitus and heart failure Strata 2: People with atherosclerotic cardiovascular	. ,	
Strata 1: People with type 2 diabetes mellitus and heart failure Strata 2: People with atherosclerotic cardiovascular		metformin was initiated if Fasting Plasma Glucose >15.0 mmol/l after day 1 to week 6, >13.3 mmol/l after week 6 to week 12 and >11.1 mmol/l after
People with type 2 diabetes mellitus and heart failure Strata 2: People without other cardiovascular diseases People with atherosclerotic cardiovascular	Cointervention	None.
People with atherosclerotic cardiovascular	People with type 2 diabetes mellitus and	Not stated/unclear
	People with atherosclerotic cardiovascular	

Strata 3: People with type 2 diabetes mellitus and chronic kidney disease	Mixed population
Strata 4: People with type 2 diabetes mellitus and high cardiovascular risk	People at higher risk of developing cardiovascular disease
Subgroup 1: People with frailty	Not stated/unclear
Subgroup 2: Onset of type 2 diabetes mellitus	People with type 2 diabetes first diagnosed above 40 years of age
Subgroup 3: People with non-alcoholic fatty liver disease	Not stated/unclear
Subgroup 4: People with obesity	Not stated/unclear
Subgroup 5: eGFR category at baseline	eGFR ≥30mL/min/1.73m2
Subgroup 6: Albuminuria category at baseline	Not stated/unclear
Sensitivity analysis category: Enrichment trial status	7) Mixed population
Population subgroups	
Comparator	Placebo daily, taken orally
	During the double-blind treatment period, glycaemic rescue therapy with metformin was initiated if FPG >15.0 mmol/l after day 1 to week 6, >13.3

	mmol/l after week 6 to week 12 and >11.1 mmol/l after week 12 to week 26.
Number of participants	N=584
Duration of follow-up	26-week treatment period, no follow-up reported.
Indirectness	The study reports that 17 countries were included in this study, but the countries are not listed. The White, Black and Asian populations included in the study may provide some generalisability to a UK population but this could be limited due to the settings and countries.
Method of analysis	Modified ITT

103.2.1. Placebo once daily (N = 192)

103.2.2. Canagliflozin 100 mg (N = 195)

103.2.3. Canagliflozin 300 mg (N = 197)

103.3. Characteristics

103.3.1. Arm-level characteristics

Characteristic	Placebo once daily (N = 192)	Canagliflozin 100 mg (N = 195)	Canagliflozin 300 mg (N = 197)
% Male	n = 88 ; % = 45.8	n = 81 ; % = 41.5	n = 89 ; % = 45.2
No of events			
Mean age (SD)	55.7 (10.9)	55.1 (10.8)	55.3 (10.2)
Mean (SD)			
White	n = 20 ; % = 10.4	n = 26 ; % = 13.3	n = 17; % = 8.6
No of events			
Black or African American	n = 9; % = 4.7	n = 18; % = 9.2	n = 14 ; % = 7.1
No of events			

Characteristic	Placebo once daily (N = 192)	Canagliflozin 100 mg (N = 195)	Canagliflozin 300 mg (N = 197)
Asian	n = 29 ; % = 15.1	n = 27 ; % = 13.8	n = 29 ; % = 14.7
No of events			
Other	n = 20 ; % = 10.4	n = 26 ; % = 13.3	n = 17; % = 8.6
No of events			
Comorbidities	NR	NR	NR
Nominal			
Presence of frailty	NR	NR	NR
Nominal			
Time since type 2 diabetes diagnosis	4.2 (4.1)	4.5 (4.4)	4.3 (4.7)
Mean (SD)			
Smoking status	NR	NR	NR
Nominal			
Alcohol consumption	NR	NR	NR
Nominal			
Number of people with obesity	NR	NR	NR
Nominal			
Albumin creatinine ratio	NR	NR	NR
Nominal			
Other antidiabetic medication used	n = 92 ; % = 47.9	n = 94 ; % = 48.2	n = 95 ; % = 48.2
No of events			
Blood pressure-lowering medication used	NR	NR	NR
Nominal			
Statins/lipid-lowering medication used	NR	NR	NR
Nominal			
Other treatment being received	NR	NR	NR
Nominal			

104. Suzuki, 2017

Bibliographic Reference

Suzuki, Shuichi; Oura, Tomonori; Takeuchi, Masakazu; Boye, Kristina S; Evaluation of the impact of once weekly dulaglutide on patient-reported outcomes in Japanese patients with type 2 diabetes: comparisons with liraglutide, insulin glargine, and placebo in two randomized studies.; Health and quality of life outcomes; 2017; vol. 15 (no. 1); 123

104.1. 3	tudy details		
Secondary	26 week efficacy results, including placebo:		
publication of another included study- see primary study for details	 Miyagawa, J., Odawara, M., Takamura, T., Iwamoto, N., Takita, Y., & Imaoka, T. (2015). Once-weekly glucagon-like peptide-1 receptor agonist dulaglutide is non-inferior to once-daily liraglutide and superior to placebo in Japanese patients with type 2 diabetes: a 26-week randomized phase III study. <i>Diabetes, Obesity and Metabolism</i>, 17(10), 974-983. 		
Other	52 week efficacy results:		
publications associated with this study included in review	 Odawara, M., Miyagawa, J., Iwamoto, N., Takita, Y., Imaoka, T., & Takamura, T. (2016). Once-weekly glucagon-like peptide-1 receptor agonist dulaglutide significantly decreases glycated haemoglobin compared with once-daily liraglutide in J apanese patients with type 2 diabetes: 52 weeks of treatment in a randomized phase III study. <i>Diabetes, Obesity and Metabolism</i>, 18(3), 249-257. 		
Trial name / registration number	Secondary article reporting on NCT01558271 and NCT01584232		
Study type	Randomised controlled trial (RCT)		
Study location	Japan (33 sites in 14 cities)		
Study setting	Community		
Study dates	04/2012 to 10/2013		
Sources of funding	Eli Lilly Japan K.K., Kobe, Japan.		
Inclusion criteria	 Diagnosis of type 2 diabetes before screening Management with diet and exercise only or treatment with oral antidiabetic monotherapy (except for thiazolidinedione) and willing to discontinue medication with 8 week washout period before randomisation HbA1c level 7.0-10.0% inclusive at screening and randomisation for those on diet and exercise only; or HbA1c level 6.5-9.0% inclusive at screening and HbA1c level 7.0-10.0% inclusive at randomisation for those on oral antidiabetic monotherapy 		

	• BMI ≥18.5 to ≥35.0 kg/m2			
Exclusion criteria	 Diagnosis of type 1 diabetes. Previous treatment with any other GLP-1 analogue. Receiving more than half of maximum dose of sulfonylureas at screening Currently taking insulin or thiazolidinediones (TZD), or previous insulin or TZD treatment ≤3 months screening. Obvious clinical signs or symptoms of pancreatitis, history of chronic pancreatitis, or acute pancreatitis at screening, as determined by investigator Serum amylase concentration ≥3 times upper limit of reference range and/or a serum lipase concentration ≥2 times upper limit of reference range, as determined by central laboratory at screening History (personal or family) of medullary C-cell hyperplasia, focal hyperplasia, or medullary thyroid carcinoma 			
Recruitment / selection of participants	Participants recruited from 33 sites in Japan. Initial 2 week screening period, then 2 week lead-in period for treatment-naive participants and 8 week wash out period for participants on monotherapy. Eligible participants randomized to treatment in 4:2:1 ratio (dulaglutide; liraglutide; placebo) using computer-generated random sequence with interactive voice response system and stratified by pre-study oral antidiabetic medication status (yes/no), BMI group (<25; ≥25 kg/m2), and HbA1c (≤8.5; >8.5%). Participants and investigators masked to assignment to dulaglutide and placebo treatment but not masked to assignment to liraglutide treatment. At end of 26 weeks, participants in placebo group switched to dulaglutide 0.75 once weekly for remaining 26 weeks. Participants not tolerating study drugs discontinued them but remained in study to collect safety data.			
Intervention(s)	 Dulaglutide 0.75 mg once weekly Subcutaneous injection of dulaglutide 0.75 mg once weekly for 52 weeks, provided in non-identifiable solution in prefilled syringe and initiated at full dose. 			
Strata 1: People with type 2 diabetes mellitus and heart failure	Not stated/unclear			
Strata 2: People with atherosclerotic cardiovascular diseases	Not stated/unclear			
Strata 3: People with type 2	Not stated/unclear			

Not stated/unclear
Not stated/unclear
Not stated/unclear
Not stated/unclear
Not stated/unclear
5) All treatment naïve Participants were treatment-naive (diabetes managed with diet and exercise only) or if on oral antidiabetic monotherapy had 8-week washout period
 Liraglutide 0.9 mg once weekly Placebo Open label subcutaneous liraglutide uptitrated from 0.3 mg/day during week 1 to 0.6 mg/day during week 2 and 0.9 mg/day at start of week 3 for remaining 49 weeks. Participants in placebo arm received placebo for 26 weeks then were switched over to dulaglutide 0.75 mg once weekly for remaining 26 weeks.
N=492
26 weeks
All participants are Japanese so unlikely to be representative of UK population. Participants who were already on oral antidiabetic medication had 8-week washout period before randomisation.
Modified ITT mITT analysis for efficacy analysis - all randomised participants who took at least one dose of study drug.

Other

Safety analysis conducted on as-treated population according to actual treatments received.

104.2. Study arms

104.2.1. **Dulaglutide 0.75 mg once weekly (N = 281)**

Subcutaneous injection of dulaglutide 0.75 mg once weekly for 52 weeks.

104.2.2. Liraglutide 0.9 mg once daily (N = 141)

Subcutaneous injection of liraglutide 0.9 mg once daily for 52 weeks.

104.2.3. Placebo once weekly for 26 weeks then dulalgutide 0.75 mg once weekly for 26 weeks N=79 (N = 79)

Placebo injection once weekly using non-identifiable prefilled syringe (same as used for dulaglutide) for 26 weeks, then switched to dulaglutide 0.75 mg once weekly for 26 weeks.

105. Tan, 2005

Bibliographic Reference

Tan, M. H.; Baksi, A.; Krahulec, B.; Kubalski, P.; Stankiewicz, A.; Urquhart, R.; Edwards, G.; Johns, D.; Comparison of pioglitazone and gliclazide in sustaining glycemic control over 2 years in patients with type 2 diabetes; Diabetes Care; 2005; vol. 28 (no. 3); 544-50

Secondary publication of another included study- see				
primary study for details				
	Links to Charbonnel 2005 - different population numbers but states in the paper that the continuous outcomes were reported here previously.			
Trial name / Cregistration number	GLAL study.			
Study type F	Randomised controlled trial (RCT)			
Study location N	Multicentre trial.			
Study setting (Outpatient follow-up.			
Study dates N	No additional information.			
Sources of Funding	Funded by Takeda Europe Research and Development Centre.			
criteria d	People with type 2 diabetes inadequate controlled (HbA1c 7.5-11.0%) with diet alone; male and females aged 35-75 years; no prior use of any oral antidiabetic medication.			
Exclusion Notice of the Control of t	No additional information.			
Recruitment / N selection of participants	No additional information.			
4	Gliclazide N=297 Gliclazide starting at 80mg daily for 4 weeks, increased to 160mg daily for 4 weeks, increased to 240mg daily for 4 weeks then 320mg daily for 4 weeks. This was then continued for up to 2 years.			
	Concomitant therapy: No additional information.			

Strata 1: People with type 2 diabetes mellitus and heart failure	Not stated/unclear
Strata 2: People with atherosclerotic cardiovascular diseases	Not stated/unclear
Strata 3: People with type 2 diabetes mellitus and chronic kidney disease	Not stated/unclear
Strata 4: People with type 2 diabetes mellitus and high cardiovascular risk	Not stated/unclear
Subgroup 1: People with frailty	Not stated/unclear
Subgroup 2: Onset of type 2 diabetes mellitus	Not stated/unclear
Subgroup 3: People with non-alcoholic fatty liver disease	Not stated/unclear
Subgroup 4: People with obesity	Not stated/unclear
Subgroup 5: eGFR category at baseline	Not stated/unclear
Subgroup 6: Albuminuria category at baseline	Not stated/unclear
Sensitivity analysis	5) All treatment naïve

category: Enrichment trial status	
Population subgroups	No additional information.
Comparator	Pioglitazone N=270 Pioglitazone starting at 15mg daily for 4 weeks, increased to 30mg daily for 4 weeks, increased to 45mg daily for 8 weeks. This was then continued for up to 2 years.
Number of participants	567
Duration of follow-up	24 months, but reporting follow up for an additional 12 months, so will count as 12 months in the results spreadsheet.
Indirectness	No additional information.
Method of analysis	Not stated/unclear Appears to be completers only.
Additional comments	No additional information.

105.2.1. Gliclazide (N = 297)

Gliclazide starting at 80mg daily for 4 weeks, increased to 160mg daily for 4 weeks, increased to 240mg daily for 4 weeks then 320mg daily for 4 weeks. This was then continued for up to 2 years. Concomitant therapy: No additional information.

105.2.2. Pioglitazone (N = 270)

Pioglitazone starting at 15mg daily for 4 weeks, increased to 30mg daily for 4 weeks, increased to 45mg daily for 8 weeks. This was then continued for up to 2 years. Concomitant therapy: No additional information.

105.3. Characteristics

105.3.1. Arm-level characteristics

Characteristic	Gliclazide (N = 297)	Pioglitazone (N = 270)
% Male	n = 182; % = 61.3	n = 171 ; % = 63.3
Sample size		

Characteristic	Gliclazide (N = 297)	Pioglitazone (N = 270)
Mean age (SD) (years)	56 (9.9)	57 (9.8)
Mean (SD)		0.0)
Ethnicity	n = NA ; % = NA	n = NA ; % = NA
Sample size		11 - 1474 , 70 - 1474
White	n = 275; % = 92.6	n = 253 ; % = 93.7
Sample size		11 - 200 , 70 - 30.7
Others	n = 22 ; % = 7.4	n = 17; % = 6.3
Sample size		11 - 17 , 70 - 0.3
Comorbidities	n = NR ; % = NR	n = ND + 0/ = ND
Sample size		n = NR ; % = NR
Presence of frailty	n = NR ; % = NR	ND % ND
Sample size		n = NR ; % = NR
Time since type 2 diabetes diagnosis	2.9 (3.8)	
(years)	2.9 (3.0)	2.7 (3.5)
Mean (SD)		
Smoking status	n = NR ; % = NR	n = NR ; % = NR
Sample size		·
Alcohol consumption	n = NR ; % = NR	n = NR ; % = NR
Sample size		, ,,
Presence of severe mental illness	n = NR ; % = NR	n = NR ; % = NR
Sample size		11 - 141C, 70 - 141C
People with significant cognitive	n = NR ; % = NR	n = NR ; % = NR
impairment		11 - INIX , 70 - INIX
Sample size		
People with a learning disability	n = NR ; % = NR	n = NR ; % = NR
Sample size		
Number of people with obesity	n = NR; % = NR	n = NR ; % = NR
Sample size		,
Other antidiabetic medication used	n = NR ; % = NR	n = NR ; % = NR
Sample size		

Characteristic	Gliclazide (N = 297)	Pioglitazone (N = 270)
Blood pressure-lowering medication used	n = NR ; % = NR	n = NR ; % = NR
Sample size		
Statins/lipid-lowering medication used	n = NR ; % = NR	n = NR ; % = NR
Sample size		
Other treatment being received	n = NR ; % = NR	n = NR ; % = NR
Sample size		

106. Tao, 2018

Bibliographic Reference

Tao, T.; Wu, P.; Wang, Y.; Liu, W.; Comparison of glycemic control and beta-cell function in new onset T2DM patients with PCOS of metformin and saxagliptin monotherapy or combination treatment; BMC Endocrine Disorders; 2018; vol. 18 (no. 1); 14

Secondary publication of another included study- see primary study for details	No additional information.
Other publications associated with this study included in review	No additional information.
Trial name / registration number	ChiCTR-IPR-17011120.
Study type	Randomised controlled trial (RCT)
Study location	China.
Study setting	Outpatient follow-up.
Study dates	No additional information.
Sources of funding	Supported by the National Natural Science Foundation of China (grant number 81200628), the Chinese Medical Association Clinical Research and Special Funds - Squibb Endocrinology Diabetes Research projects [2012]; the Natural Science Foundation of Shanghai, China [grant number 12ZR1417800]l and the Shanghai Science and Technology Development Fund [grant number 08411953000].
Inclusion criteria	Newly diagnosed people with type 2 diabetes mellitus and polycystic ovarian syndrome.
Exclusion criteria	People with coronary atherosclerotic heart disease; abnormal liver and renal function; diabetic ketoacidosis; chronic inflammatory disease; severe gastrointestinal disease.
Recruitment / selection of participants	People were recruited from the Outpatient Department of Endocrinology and Metabolism at Shanghai Renji Hospital.
Intervention(s)	Saxagliptin + Metformin N=25

	Saxagliptin 5mg/day and metformin 2000mg/day for 24 weeks at a fixed dose.
	Concomitant therapy: All people were given a control diet for 2 weeks before the program. All people were given advice on diet and exercise and were asked to follow a behaviour modification program.
Strata 1: People with type 2 diabetes mellitus and heart failure	Not stated/unclear
Strata 2: People with atherosclerotic cardiovascular diseases	People without other cardiovascular diseases
Strata 3: People with type 2 diabetes mellitus and chronic kidney disease	People without chronic kidney disease
Strata 4: People with type 2 diabetes mellitus and high cardiovascular risk	People at higher risk of developing cardiovascular disease Based on BMI, triglycerides and presence of diabetes.
Subgroup 1: People with frailty	Not stated/unclear
Subgroup 2: Onset of type 2 diabetes mellitus	Not stated/unclear
Subgroup 3: People with non-alcoholic fatty liver disease	People without non-alcoholic fatty liver disease
Subgroup 4: People with obesity	Not stated/unclear

Subgroup 5: eGFR category at baseline Subgroup 6: Albuminuria category at baseline Sensitivity analysis category: Enrichment trial status Population subgroups Comparator Saxagliptin N=25 Saxagliptin 5mg/day for 24 weeks at a fixed dose. Concomitant therapy: All people were given a control diet for 2 weeks before the program. All people were given advice on diet and exercise and were asked to follow a behaviour modification program.
Albuminuria category at baseline Sensitivity analysis category: Enrichment trial status Population subgroups Comparator Saxagliptin N=25 Saxagliptin 5mg/day for 24 weeks at a fixed dose. Concomitant therapy: All people were given a control diet for 2 weeks before the program. All people were given advice on diet and exercise and
analysis category: Enrichment trial status Population subgroups Comparator Saxagliptin N=25 Saxagliptin 5mg/day for 24 weeks at a fixed dose. Concomitant therapy: All people were given a control diet for 2 weeks before the program. All people were given advice on diet and exercise and
Saxagliptin N=25 Saxagliptin 5mg/day for 24 weeks at a fixed dose. Concomitant therapy: All people were given a control diet for 2 weeks before the program. All people were given advice on diet and exercise and
Saxagliptin 5mg/day for 24 weeks at a fixed dose. Concomitant therapy: All people were given a control diet for 2 weeks before the program. All people were given advice on diet and exercise and
Concomitant therapy: All people were given a control diet for 2 weeks before the program. All people were given advice on diet and exercise and
before the program. All people were given advice on diet and exercise and
Metformin N=25
Metformin 2000mg/day for 24 weeks at a fixed dose.
Concomitant therapy: All people were given a control diet for 2 weeks before the program. All people were given advice on diet and exercise and were asked to follow a behaviour modification program.
Number of 75 participants
Duration of 24 weeks. follow-up
Indirectness Population indirectness - People with polycystic ovary syndrome and type 2 diabetes.
Method of analysis Per protocol
Additional No additional information. comments

106.2.1. Saxagliptin + Metformin (N = 25)

Saxagliptin 5mg/day and metformin 2000mg/day for 24 weeks at a fixed dose. Concomitant therapy: All people were given a control diet for 2 weeks before the program. All people were given advice on diet and exercise and were asked to follow a behaviour modification program.

106.2.2. Saxagliptin (N = 25)

Saxagliptin 5mg/day for 24 weeks at a fixed dose. Concomitant therapy: All people were given a control diet for 2 weeks before the program. All people were given advice on diet and exercise and were asked to follow a behaviour modification program.

106.2.3. Metformin (N = 25)

Metformin 2000mg/day for 24 weeks at a fixed dose. Concomitant therapy: All people were given a control diet for 2 weeks before the program. All people were given advice on diet and exercise and were asked to follow a behaviour modification program.

106.3. Characteristics

106.3.1. Arm-level characteristics

Characteristic	Saxagliptin + Metformin (N = 25)	Saxagliptin (N = 25)	Metformin (N = 25)
% Male	n = 0; % = 0	n = 0 ; % = 0	n = 0 ; % = 0
Sample size			
Mean age (SD) (years)	29 (5)	30 (5)	28 (3)
Mean (SD)			
Ethnicity	n = NR ; % = NR	n = NR ; % =	n = NR ; % =
Sample size		NR	NR
Comorbidities	n = NR ; % = NR	n = NR ; % =	n = NR ; % =
Sample size		NR	NR
Presence of frailty	n = NR ; % = NR	n = NR ; % =	n = NR ; % =
Sample size		NR	NR

Characteristic	Saxagliptin + Metformin (N = 25)	Saxagliptin (N = 25)	Metformin (N = 25)
Time since type 2 diabetes diagnosis	NR (NR)	NR (NR)	NR (NR)
Mean (SD)			
HbA1c (%)	7.4 (0.3)	7.4 (0.3)	7.3 (0.2)
Mean (SD)			
Blood pressure	NR (NR)	NR (NR)	NR (NR)
Mean (SD)			
Heart rate	NR (NR)	NR (NR)	NR (NR)
Mean (SD)	n = NR ; % = NR		
Smoking status Sample size	II - INK , 70 - INK	n = NR ; % = NR	n = NR ; % = NR
Alcohol consumption	n = NR ; % = NR	n = NR ; % =	n = NR ; % =
Sample size		NR	NR
Presence of severe mental illness	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
Sample size			
People with significant cognitive impairment	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
Sample size			
People with a learning disability	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
Sample size			
Weight (kg)	69.3 (64.6 to 74.1)	70.4 (63.7 to	67.9 (63.6 to
Mean (95% CI)		77.1)	72.2)
BMI (kg/m2)	26.38 (24.66 to 28.1)	27.2 (24.94 to	26.4 (24.63 to
Mean (95% CI)		29.46)	28.18)
Number of people with obesity Sample size	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
Cholesterol and lipid levels	n = NR ; % = NR		
Sample size		n = NR ; % = NR	n = NR ; % = NR
•			

Characteristic	Saxagliptin + Metformin (N = 25)	Saxagliptin (N = 25)	Metformin (N = 25)
Albumin creatinine ratio	NR (NR)	NR (NR)	NR (NR)
Mean (SD)			
eGFR (mL/min/1.73m2)	NR (NR)	NR (NR)	NR (NR)
Mean (SD)			
Other antidiabetic medication used	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
Sample size			
Blood pressure-lowering medication used	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
Sample size			
Statins/lipid-lowering medication used	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
Sample size			
Other treatment being received	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
Sample size			

107. Wainstein, 2012

Bibliographic Reference

Wainstein, J; Katz, L; Engel, S S; Xu, L; Golm, G T; Hussain, S; O'Neill, E A; Kaufman, K D; Goldstein, B J; Initial therapy with the fixed-dose combination of sitagliptin and metformin results in greater improvement in glycaemic control compared with pioglitazone monotherapy in patients with type 2 diabetes.; Diabetes, obesity & metabolism; 2012; vol. 14 (no. 5); 409-18

I T
No additional information.
No additional information.
NCT00532935.
Randomised controlled trial (RCT)
Multicentre trial.
Outpatient follow-up.
No additional information.
All authors except one were employees of Merck Sharp and Dohme Corporation, with the other author receiving honoraria for lecturing with the organisation.
At least 18 to no more than 78 years of age with a diagnosis of type 2 diabetes and inadequate glycaemic control (defined as HbA1c no less than 7.5% and no more than 12.0% while on a diet/exercise regimen); not on an antihyperglycaemic agent in the 3 months prior to the screening visit and were to have had less than 4 weeks of cumulative duration of treatment and an antihyperglycaemic over the 3 years prior to the screening visit.
History of type 1 diabetes; contraindication to biguanide or thiazolidinedione medications; previous treatment with any DPP-4 inhibitor or incretin mimetic; required treatment with CYP2C8 inhibitors or inducers; had impaired renal function (creatinine clearance <60mL/min); alanine aminotransferase or aspartate aminotransferase levels more than twofold the upper limit of normal, or a fasting glucose value <130 mg/dL or >320 mg/dL prior to randomisation.

Recruitment / selection of participants	No additional information.
Intervention(s)	Sitagliptin + Metformin N=261
	Sitagliptin 50mg + Metformin 500mg twice daily in a fixed dose combination. The dose was increased to 50/500mg in the morning and 50/1000mg in the evening at week 2, then to 50/1000mg twice daily at week 4. Given for a total of 32 weeks.
Cointervention	Concomitant therapy: No additional information.
Strata 1: People with type 2 diabetes mellitus and heart failure	Not stated/unclear
Strata 2: People with atherosclerotic cardiovascular diseases	Not stated/unclear
Strata 3: People with type 2 diabetes mellitus and chronic kidney disease	Not stated/unclear
Strata 4: People with type 2 diabetes mellitus and high cardiovascular risk	Not stated/unclear
Subgroup 1: People with frailty	Not stated/unclear
Subgroup 2: Onset of type 2 diabetes mellitus	Not stated/unclear
Subgroup 3: People with non-alcoholic fatty liver disease	Not stated/unclear

Subgroup 5: eGFR category at baseline Not stated/unclear Subgroup 5: eGFR category at baseline Not stated/unclear Subgroup 6: Albuminuria category at baseline Not response criteria Sensitivity analysis category: Enrichment trial status 6) No response criteria Population subgroups No additional information. Comparator Pioglitazone N=256 Pioglitazone 30mg per day, increased to 45mg per day at week 4. Given for a total of 32 weeks. Number of participants 521 (includes 2 people who were randomised twice at two different sites but who were not included in any analyses). Duration of follow-up 32 weeks. Indirectness No additional information. Method of analysis ACA Efficacy outcomes = People who took at least one dose of the medication and had both a baseline measurement and at least one post- randomisation measurement of the respective endpoint. Safety = People who took at least one dose of the medication. Additional comments No additional information.		
eGFR category at baseline Subgroup 6: Albuminuria category at baseline Sensitivity analysis category: Enrichment trial status Population subgroups Comparator Pioglitazone N=256 Pioglitazone 30mg per day, increased to 45mg per day at week 4. Given for a total of 32 weeks. Number of participants but who were not included in any analyses). Duration of follow-up Indirectness Method of analysis Efficacy outcomes = People who took at least one dose of the medication and had both a baseline measurement and at least one post-randomisation measurement of the respective endpoint. Safety = People who took at least one dose of the medication. Additional No additional information.	People with	Not stated/unclear
Albuminuria category at baseline Sensitivity analysis category: Enrichment trial status Population subgroups Comparator Pioglitazone N=256 Pioglitazone 30mg per day, increased to 45mg per day at week 4. Given for a total of 32 weeks. Number of participants Duration of follow-up Indirectness No additional information. Method of analysis Efficacy outcomes = People who took at least one dose of the medication and had both a baseline measurement and at least one post-randomisation measurement of the respective endpoint. Safety = People who took at least one dose of the medication. Additional No additional information.	eGFR category	Not stated/unclear
analysis category: Enrichment trial status Population subgroups Comparator Pioglitazone N=256 Pioglitazone 30mg per day, increased to 45mg per day at week 4. Given for a total of 32 weeks. Number of participants Duration of follow-up Indirectness No additional information. ACA Efficacy outcomes = People who took at least one dose of the medication and had both a baseline measurement and at least one post- randomisation measurement of the respective endpoint. Safety = People who took at least one dose of the medication. Additional No additional information.	Albuminuria category at	Not stated/unclear
Comparator Pioglitazone N=256 Pioglitazone 30mg per day, increased to 45mg per day at week 4. Given for a total of 32 weeks. Number of participants Duration of follow-up Indirectness No additional information. ACA Efficacy outcomes = People who took at least one dose of the medication and had both a baseline measurement and at least one post-randomisation measurement of the respective endpoint. Safety = People who took at least one dose of the medication. Additional No additional information.	analysis category: Enrichment	6) No response criteria
Pioglitazone 30mg per day, increased to 45mg per day at week 4. Given for a total of 32 weeks. Number of participants 521 (includes 2 people who were randomised twice at two different sites but who were not included in any analyses). Duration of follow-up Indirectness No additional information. Method of analysis Efficacy outcomes = People who took at least one dose of the medication and had both a baseline measurement and at least one post-randomisation measurement of the respective endpoint. Safety = People who took at least one dose of the medication. Additional No additional information.	-	No additional information.
participants but who were not included in any analyses). Duration of follow-up Indirectness No additional information. Method of analysis Efficacy outcomes = People who took at least one dose of the medication and had both a baseline measurement and at least one post-randomisation measurement of the respective endpoint. Safety = People who took at least one dose of the medication. Additional No additional information.	Comparator	Pioglitazone 30mg per day, increased to 45mg per day at week 4. Given
Indirectness No additional information. Method of analysis Efficacy outcomes = People who took at least one dose of the medication and had both a baseline measurement and at least one post-randomisation measurement of the respective endpoint. Safety = People who took at least one dose of the medication. Additional No additional information.		
Method of analysis Efficacy outcomes = People who took at least one dose of the medication and had both a baseline measurement and at least one post-randomisation measurement of the respective endpoint. Safety = People who took at least one dose of the medication. Additional No additional information.		32 weeks.
analysis Efficacy outcomes = People who took at least one dose of the medication and had both a baseline measurement and at least one post-randomisation measurement of the respective endpoint. Safety = People who took at least one dose of the medication. Additional No additional information.	Indirectness	No additional information.
		Efficacy outcomes = People who took at least one dose of the medication and had both a baseline measurement and at least one post-randomisation measurement of the respective endpoint. Safety = People
		No additional information.

107.2.1. Sitagliptin + Metformin (N = 261)

Sitagliptin 50mg + Metformin 500mg twice daily in a fixed dose combination. The dose was increased to 50/500mg in the morning and 50/1000mg in the evening at week 2, then to 50/1000mg twice daily at week 4. Given for a total of 32 weeks. Concomitant therapy: No additional information.

107.2.2. Pioglitazone (N = 256)

Pioglitazone 30mg per day, increased to 45mg per day at week 4. Given for a total of 32 weeks. Concomitant therapy: No additional information.

107.3. Characteristics

107.3.1. Arm-level characteristics

7		
Characteristic	Sitagliptin + Metformin (N = 261)	Pioglitazone (N = 256)
% Male	n = 143 ; % = 54.8	n = 134 ; % = 52.3
Sample size		
Mean age (SD) (years)	52.4 (10.7)	52.2 (11)
Mean (SD)		
Ethnicity	n = NA ; % = NA	n = NA ; % = NA
Sample size		
White	n = 168; % = 64.4	n = 167; % = 65.2
Sample size		
Asian	n = 58 ; % = 22.2	n = 55 ; % = 21.5
Sample size		
Multiracial	n = 27 ; % = 10.3	n = 29 ; % = 11.3
Sample size		
Black or African Sample size	n = 6; % = 2.3	n = 5; % = 2
·	0 : 0/ 0 0	
American Indian Sample size	n = 2; % = 0.8	n = 0; % = 0
Comorbidities	n = NR ; % = NR	n = ND : 0/ = ND
Sample size		n = NR ; % = NR
Presence of frailty	n = NR	n = NR ; % = NR
Sample size		
Time since type 2 diabetes diagnosis (years)	3.2 (4)	3.3 (3.5)
Mean (SD)		

Characteristic	Sitagliptin + Metformin (N = 261)	Pioglitazone (N = 256)
Smoking status	n = NA ; % = NA	n = NA ; % = NA
Sample size		
Alcohol consumption	n = NA ; % = NA	n = NA ; % = NA
Sample size		
Presence of severe mental illness	n = NA ; % = NA	n = NA ; % = NA
Sample size		,
People with significant cognitive impairment	n = NA ; % = NA	n = NA
Sample size		
People with a learning disability	n = NA ; % = NA	n = NA ; % = NA
Sample size		
Number of people with obesity	n = NA ; % = NA	n = NA ; % = NA
Sample size		
Other antidiabetic medication used	n = NA ; % = NA	n = NA ; % = NA
Sample size		
Blood pressure-lowering medication used	n = NA ; % = NA	n = NA ; % = NA
Sample size		
Statins/lipid-lowering medication used	n = NA ; % = NA	n = NA ; % = NA
Sample size		
Other treatment being received	n = NA ; % = NA	n = NA ; % = NA
Sample size		

108. Wang, 2013

Bibliographic Reference

Wang, H.; Ni, Y.; Yang, S.; Li, H.; Li, X.; Feng, B.; The effects of gliclazide, metformin, and acarbose on body composition in patients with newly diagnosed type 2 diabetes mellitus; Curr Ther Res Clin Exp; 2013; vol. 75; 88-92

Secondary publication of another included study- see primary study for details	No additional information.
Other publications associated with this study included in review	No additional information.
Trial name / registration number	No additional information.
Study type	Randomised controlled trial (RCT)
Study location	China.
Study setting	Outpatient follow-up.
Study dates	No additional information.
Sources of funding	Supported by a Young Medical Talents Training Program Grant of Pudong Health Bureau of Shanghai (No. PWRq2012-08) and the Key Disciplines Group Construction Project of Pudong Health Bureau of Shanghai (No. PWZxkq2010-04).
Inclusion criteria	People newly diagnosed with type 2 diabetes according to the results of oral glucose tolerance test; hyperglycaemia (HbA1c 7-10%); drug naïve.
Exclusion criteria	Severe congestive heart failure (NYHA III-IV); liver dysfunction (AST/ALT >1.5 x upper limit of normal); renal dysfunction (creatinine clearance <90mL/min; creatinine clearance calculated using the Cockcroft-Gault formula); extraordinary body weight (body mass index <18.5 or >35kg/m2); dyslipidaemia (total cholesterol >6.21mmol/L; triglycerides >2.25mmol/L, LDL cholesterol >4.13mmol/L); people receiving antidiabetes treatment before the study; taking pharmacologic agents known to affect carbohydrate homeostasis or influence lipid levels; type 1 diabetes mellitus.
Recruitment / selection of participants	No additional information.

Intervention(s)	Gliclazide N=30
	Gliclazide 120mg/day for 6 months.
	Concomitant therapy: All people received diet therapy.
Strata 1: People with type 2 diabetes mellitus and heart failure	People without heart failure
Strata 2: People with atherosclerotic cardiovascular diseases	Not stated/unclear
Strata 3: People with type 2 diabetes mellitus and chronic kidney disease	People without chronic kidney disease
Strata 4: People with type 2 diabetes mellitus and high cardiovascular risk	People at higher risk of developing cardiovascular disease Based on age, triglycerides and presence of diabetes.
Subgroup 1: People with frailty	Not stated/unclear
Subgroup 2: Onset of type 2 diabetes mellitus	Not stated/unclear
Subgroup 3: People with non-alcoholic fatty liver disease	People without non-alcoholic fatty liver disease
Subgroup 4: People with obesity	Not stated/unclear

Subgroup 5: eGFR category at baseline	Not stated/unclear
Subgroup 6: Albuminuria category at baseline	Not stated/unclear
Sensitivity analysis category: Enrichment trial status	5) All treatment naïve
Population subgroups	No additional information.
Comparator	Metformin N=30
	Metformin 1700mg/day for 6 months.
	Concomitant therapy: All people received diet therapy.
	A third arm (n=30) was reported where people received acarbose. This arm is not extracted as it is not relevant to the protocol.
Number of participants	90 (including the acarbose arm, 60 without).
Duration of follow-up	6 months.
Indirectness	Outcome indirectness - only withdrawal due to adverse events was reported in a manner where it could be attributed to study arms and so this was extracted instead of withdrawal due to any reason.
Method of analysis	Not stated/unclear
Additional comments	No additional information.

108.2.1. Gliclazide (N = 30)

Gliclazide 120mg/day for 6 months. Concomitant therapy: All people received diet therapy.

108.2.2. Metformin (N = 30)

Metformin 1700mg/day for 6 months. Concomitant therapy: All people received diet therapy.

108.3. Characteristics

108.3.1. Arm-level characteristics

100.0.1. Alli-level characteristic		
Characteristic	Gliclazide (N = 30)	Metformin (N = 30)
% Male	n = 21 ; % = 70	n = 18 ; % = 60
Sample size		
Mean age (SD) (years)	55.89 (10.5)	54 (10.3)
Mean (SD)		
Ethnicity	n = NR ; % = NR	n = NR ; % = NR
Sample size		
Comorbidities Sample size	n = NR ; % = NR	n = NR ; % = NR
Sample size	ND - 0/ ND	
Presence of frailty	n = NR ; % = NR	n = NR ; % = NR
Sample size		
Time since type 2 diabetes diagnosis	NR (NR)	NR (NR)
Mean (SD)		
HbA1c (%)	8.4 (0.93)	8.07 (0.77)
Mean (SD)		
Blood pressure (mmHg)	NA (NA)	NA (NA)
Mean (SD)		
Systolic blood pressure	125.1 (9.4)	126.4 (9)
Mean (SD)		
Diastolic blood pressure	76.3 (7.4)	74.9 (7)
Mean (SD)		
Heart rate	NR (NR)	NR (NR)
Mean (SD)		
Smoking status	n = NR ; % = NR	n = NR ; % = NR
Sample size		

Characteristic		Metformin (N = 30)
Alcohol consumption	n = NR ; % = NR	n = NR ; % = NR
Sample size		
Presence of severe mental illness	n = NR ; % = NR	n = NR ; % = NR
Sample size		
People with significant cognitive impairment	n = NR ; % = NR	n = NR ; % = NR
Sample size		
People with a learning disability	n = NR ; % = NR	n = NR ; % = NR
Sample size		,
Weight (kg)	70.4 (11.7)	71.6 (12.7)
Mean (SD)		,
ВМІ	NR (NR)	NR (NR)
Mean (SD)		()
Number of people with obesity	n = NR ; % = NR	n = NR ; % = NR
Sample size		11111, 70 1111
Cholesterol and lipid levels (mmol/L)	NA (NA)	NA (NA)
Mean (SD)		
Total cholesterol	5.23 (1.02)	5.02 (0.85)
Mean (SD)		3.02 (0.03)
HDL cholesterol	1.05 (0.33)	1.09 (0.32)
Mean (SD)		1.03 (0.32)
LDL cholesterol	2.6 (0.68)	2.72 (0.67)
Mean (SD)		2.12 (0.01)
Triglycerides	2.12 (1.25)	1.93 (0.53)
Mean (SD)		1.93 (0.33)
Albumin creatinine ratio	NR (NR)	NR (NR)
Mean (SD)		IVIX (IVIX)
eGFR (mL/min/1.73m2)	NR (NR)	ND (ND)
Mean (SD)		NR (NR)
Other antidiabetic medication used	n = NR ; % = NR	n = ND : 0/ = ND
Sample size		n = NR ; % = NR
- Sample 0120		

Characteristic	Gliclazide (N = 30)	Metformin (N = 30)
Blood pressure-lowering medication used	n = NR ; % = NR	n = NR ; % = NR
Sample size		
Statins/lipid-lowering medication used	n = NR ; % = NR	n = NR ; % = NR
Sample size		
Other treatment being received	n = NR ; % = NR	n = NR ; % = NR
Sample size		

109. Wang, 2022

Bibliographic Reference

Wang, X.; Zhao, B.; Sun, H.; You, H.; Qu, S.; Effects of sitagliptin on intrahepatic lipid content in patients with non-alcoholic fatty liver disease;

Frontiers in endocrinology; 2022; vol. 13; 866189

100.11.	ludy details
Secondary publication of another included study- see primary study for details	No additional information.
Other publications associated with this study included in review	No additional information.
Trial name / registration number	NCT05480007.
Study type	Randomised controlled trial (RCT)
Study location	China.
Study setting	Outpatient follow-up.
Study dates	No additional information.
Sources of funding	Supported by the Climbing Talent Program of Shanghai Tenth People's Hospital (2021SYPDRC047) and National Nature Science Foundation of China (NO.81900781).
Inclusion criteria	Age 30-70 years; fulfilment of the diagnostic criteria for T2DM by WHO in 1999 (HbA1c ranged from 7% to 10%, FPG <11 mol/L, 2-hour blood glucose postprandial <20mmol/L); fulfilment of the diagnostic criteria for NAFLD according to the guidelines of the Chinese Medical Association in 2010.
Exclusion criteria	T2DM complicated with ketoacidosis hyperosmolarity, acute and chronic infection; serious heart, liver, kidney, lung disease and liver damage; alcoholic fatty liver; drug use that influences glucose metabolism such as thiazide diuretics and hormones within three months; hypertension of more than or equal to 180/110 mmHg; gastrointestinal disease or absorption dysfunction; recent trauma, surgery or other conditions resulting in an increased stress response within the past three months.
Recruitment / selection of participants	Recruited from the outpatient department of Shanghai Tenth People's Hospital.

Intervention(s)	Sitagliptin + Metformin N=20
	Sitagliptin 100mg per day and metformin 500mg three times a day for 24 weeks.
	Concomitant therapy: Health education regarding eating a diabetic diet and performing routine exercise during the treatment period.
Strata 1: People with type 2 diabetes mellitus and heart failure	People without heart failure
Strata 2: People with atherosclerotic cardiovascular diseases	People without other cardiovascular diseases
Strata 3: People with type 2 diabetes mellitus and chronic kidney disease	People without chronic kidney disease
Strata 4: People with type 2 diabetes mellitus and high cardiovascular risk	People at higher risk of developing cardiovascular disease NAFLD and T2DM
Subgroup 1: People with frailty	Not stated/unclear
Subgroup 2: Onset of type 2 diabetes mellitus	Not stated/unclear
Subgroup 3: People with non-alcoholic fatty liver disease	People with non-alcoholic fatty liver disease
Subgroup 4: People with obesity	Not stated/unclear

Subgroup 5: eGFR category at baseline	Not stated/unclear
Subgroup 6: Albuminuria category at baseline	Not stated/unclear
Sensitivity analysis category: Enrichment trial status	8) Not reported
Population subgroups	No additional information.
Comparator	Sitagliptin N=17
	Sitagliptin 100mg per day for 24 weeks.
	Concomitant therapy: Health education regarding eating a diabetic diet and performing routine exercise during the treatment period.
	Metformin N=17
	Metformin 500mg three times a day for 24 weeks.
	Concomitant therapy: Health education regarding eating a diabetic diet and performing routine exercise during the treatment period.
Number of participants	54
Duration of follow-up	24 weeks.
Indirectness	No additional information.
Method of analysis	Not stated/unclear
Additional comments	No additional information.

109.2.1. Sitagliptin + Metformin (N = 20)

Sitagliptin 100mg per day and metformin 500mg three times a day for 24 weeks. Concomitant therapy: Health education regarding eating a diabetic diet and performing routine exercise during the treatment period.

109.2.2. Sitagliptin (N = 17)

Sitagliptin 100mg per day for 24 weeks. Concomitant therapy: Health education regarding eating a diabetic diet and performing routine exercise during the treatment period.

109.2.3. **Metformin (N = 17)**

Metformin 500mg three times a day for 24 weeks. Concomitant therapy: Health education regarding eating a diabetic diet and performing routine exercise during the treatment period.

109.3. Characteristics

109.3.1. Arm-level characteristics

7			
Characteristic	Sitagliptin + Metformin (N = 20)	Sitagliptin (N = 17)	Metformin (N = 17)
% Male	n = 9; % = 45	n = 10 ; % = 59	n = 9; % = 53
Sample size			
Mean age (SD) (years)	54.55 (6.6)	54.4 (9)	55.63 (10.9)
Mean (SD)			
Ethnicity	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
Sample size			
Comorbidities Sample size	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
•			
Presence of frailty Sample size	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
·	ND 0/ ND		
Time since type 2 diabetes diagnosis	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
Sample size			
HbA1c (%)	7.83 (0.58)	7.93 (0.91)	8.6 (1.17)
Mean (SD)			
Blood pressure	NR (NR)	NR (NR)	NR (NR)
Mean (SD)			
Heart rate	NR (NR)	NR (NR)	NR (NR)
Mean (SD)			

Characteristic	Sitagliptin + Metformin (N = 20)	Sitagliptin (N = 17)	Metformin (N = 17)
Smoking status	n = NR ; % = NR	n = NR ; % =	n = NR ; % =
Sample size		NR	NR
Alcohol consumption	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
Sample size	ND 0/ ND	·	
Presence of severe mental illness	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
Sample size			
People with significant cognitive impairment	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
Sample size	ND 0/ ND		
People with a learning disability Sample size	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
Weight	NR (NR)		
	,	NR (NR)	NR (NR)
Mean (SD)	22.27 (2.24)		
BMI (kg/m2)	26.07 (3.24)	25.41 (3.45)	26.46 (2.86)
Mean (SD)			
Number of people with obesity	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
Sample size		Turk	TVIX
Cholesterol and lipid levels (mmol/L)	NA (NA)	NA (NA)	NA (NA)
Mean (SD)			
HDL cholesterol	1.34 (0.27)	1.31 (0.38)	1.18 (0.31)
Mean (SD)	0.54 (4.04)		
LDL cholesterol	2.54 (1.01)	3.02 (0.97)	2.83 (0.65)
Mean (SD)			
Triglycerides	1.26 (0.62)	1.7 (0.82)	1.94 (0.97)
Mean (SD)			
Total cholesterol	4.46 (1.03)	5.22 (0.92)	4.88 (0.64)
Mean (SD)			
Albumin creatinine ratio	NR (NR)	NR (NR)	NR (NR)

Characteristic	Sitagliptin + Metformin (N = 20)	Sitagliptin (N = 17)	Metformin (N = 17)
Mean (SD)			
eGFR (mL/min/1.73m2)	NR (NR)	NR (NR)	NR (NR)
Mean (SD)			
Other antidiabetic medication used	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
Sample size			
Blood pressure-lowering medication used	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
Sample size			
Statins/lipid-lowering medication used	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
Sample size			
Other treatment being received	n = NR ; % = NR	n = NR ; % =	· · · · · · · · · · · · · · · · · · ·
Sample size		NR	NR

110. Wang, 2016

Bibliographic Reference

Wang, Y.; Xu, L.; Yuan, L.; Li, D.; Zhang, Y.; Zheng, R.; Liu, C.; Feng, X.; Li, Q.; Ma, J.; Sodium-glucose co-transporter-2 inhibitors suppress atrial natriuretic peptide secretion in patients with newly diagnosed Type 2 diabetes; Diabetic Med; 2016; vol. 33 (no. 12); 1732-1736

	tudy details
Secondary publication of another included study- see primary study for details	No information available.
Other publications associated with this study included in review	Li FF, Gao G, Li Q, Zhu HH, Su XF, Wu JD, Ye L, Ma JH. Influence of Dapagliflozin on Glycemic Variations in Patients with Newly Diagnosed Type 2 Diabetes Mellitus. J Diabetes Res. 2016;2016:5347262.
Trial name / registration number	No information available.
Study type	Randomised controlled trial (RCT)
Study location	China
Study setting	Hospital
Study dates	07/2010 to 03/2012
Sources of funding	Grants from the National Natural Science Foundation of China, Jiangsu Planned Projects of Postdoctoral Research Funds, the Peak of Six Personnel in Jiangsu, and the Nanjing Medical Science and Technique Development Foundation.
Inclusion criteria	Newly diagnosed or drug-naive Type 2 diabetes
Exclusion criteria	 History of diabetes Severe uncontrolled hypertension (systolic blood pressure ≥ 180 mmHg and/or diastolic blood pressure ≥ 110 mmHg) and use of any renin angiotensin system blocker. Replacement of chronic system corticosteroid treatment History or current diagnosis of significant comorbid diseases, such as cardiovascular, hepatic and renal diseases and/or Positive test for islet cell autoantibodies (such as glutamic acid decarboxylase autoantibodies, islet-cell antibodies or insulinomalike antigen 2 indicating the possibility of type 2 diabetes

Recruitment / selection of participants	After receiving 8 weeks of lifestyle management counselling, those who continued to experience inadequate glycaemic control, as defined by HbA1c levels of 58 - 80 mmol/mol (7.5-10.5%) were recruited.
Intervention(s)	Dapagliflozin After 4 weeks of treatment, patients lacking glycaemic control (fasting blood glucose >11.1 mmol/l) were eligible to receive another antihyperglycaemic drug such as metformin.
Cointervention	
Strata 1: People with type 2 diabetes mellitus and heart failure	Not stated/unclear
Strata 2: People with atherosclerotic cardiovascular diseases	
Strata 3: People with type 2 diabetes mellitus and chronic kidney disease	Not stated/unclear
Strata 4: People with type 2 diabetes mellitus and high cardiovascular risk	Not stated/unclear
Subgroup 1: People with frailty	Not stated/unclear
Subgroup 2: Onset of type 2 diabetes mellitus	People with type 2 diabetes first diagnosed above 40 years of age
Subgroup 3: People with non-alcoholic fatty liver disease	Not stated/unclear

Not stated/unclear
NOL Stated/uniclear
Not stated/unclear
Not stated/unclear
5) All treatment naïve
No information reported.
Placebo
N=28
24-week treatment period + no follow-up.
Study was conducted in a hospital in China, the results may not be as generalisable to a UK population.
Not stated/unclear
Unpaired Student's t-tests were used to compare all the variables measured for each treatment group and to determine significant differences between groups. Analyses of continuous outcomes were based on a ANCOVA model, with treatment as a fixed effect and baseline as the covariate, and were also used to estimate differences between blacebo and treatment groups.

110.2.1. Dapagliflozin (N = 18)

110.2.2. Placebo (N = 10)

110.3. Characteristics

110.3.1. Arm-level characteristics

110.5.1. Allii-level characteristics			
Characteristic	Dapagliflozin (N = 18)	Placebo (N = 10)	
% Male	n = 11; % = 61	n = 2; % = 20	
No of events		,	
Mean age (SD)	60.7 (10.1)	59.3 (9)	
Mean (SD)		00.0 (0)	
Ethnicity	NR	NR	
Nominal		INIX	
Comorbidities	NR	ND	
Nominal		NR	
Presence of frailty	NR		
Nominal		NR	
Smoking status	NR		
•		NR	
Nominal Alabal consumption	ND		
Alcohol consumption	NR	NR	
Nominal			
Presence of severe mental illness	NR	NR	
Nominal			
People with significant cognitive impairment	NR	NR	
Nominal			
People with a learning disability	NR	NR	
Nominal			
ВМІ	NR	NR	
Nominal		T T T	
Number of people with obesity	NR	NR	
Nominal		INIX	
Albumin creatinine ratio	NR	ND	
Nominal		NR	
eGFR (mL/min/1.73m2)	NR		
()		NR	

Characteristic	Dapagliflozin (N = 18)	Placebo (N = 10)
Nominal		
Other antidiabetic medication used	NR	NR
Nominal		
Blood pressure-lowering medication used	NR	NR
Nominal		
Statins/lipid-lowering medication used	NR	NR
Nominal		
Other treatment being received	NR	NR
Nominal		

111. Wolever, 2000

Bibliographic Reference

Wolever, T. M. S.; Assiff, L.; Basu, T.; Chiasson, J. L.; Boctor, M.; Gerstein, H. C.; Hunt, J. A.; Josse, R. G.; Lau, D.; Leiter, L. A.; Maheux, P.; Murphy, L.; Rodger, N. W.; Ross, S. A.; Ryan, E.; Tildesley, H. D.; Yale, J. F.; Miglitol, an alpha-glucosidase inhibitor, prevents the metformin-induced fall in serum folate and vitamin B12 in subjects with type 2 diabetes; Nutr Res; 2000; vol. 20 (no. 10); 1447-56

Secondary publication of another included study- see primary study for details	No additional information.
Other publications associated with this study included in review	No additional information.
Trial name / registration number	No additional information.
Study type	Randomised controlled trial (RCT)
Study location	Canada.
Study setting	Outpatient follow-up.
Study dates	No additional information.
Sources of funding	Supported by Bayer Canada Inc which also provided the study medications.
Inclusion criteria	Euthyroid male and non-pregnant females >40 years of age; BMI no more than 40 kg/m2; oral glucose lowering drugs had to be withdrawn and people had to undergo an 8 week placebo baseline period.
Exclusion criteria	Type 1 diabetes; insulin treatment; unwillingness to perform the self-blood glucose monitoring specified in the protocol; major debilitating diseases; recent cardiovascular events or surgery; presence of gastrointestinal diseases or use of drugs associated with abnormal intestinal motility or altered absorption; raised serum creatinine or aspartate transaminase; use of steroids; the presence of emotional disorders or substance abuse.
Recruitment / selection of participants	No additional information.

Intervention(s)	Metformin N=83
	Metformin 500mg three times daily plus miglitol placebo with meals for 36 weeks.
	Concomitant therapy: No additional information.
Strata 1: People with type 2 diabetes mellitus and heart failure	Not stated/unclear
Strata 2: People with atherosclerotic cardiovascular diseases	Not stated/unclear
Strata 3: People with type 2 diabetes mellitus and chronic kidney disease	Not stated/unclear
Strata 4: People with type 2 diabetes mellitus and high cardiovascular risk	People at higher risk of developing cardiovascular disease Based on BMI and presence of diabetes
Subgroup 1: People with frailty	Not stated/unclear
Subgroup 2: Onset of type 2 diabetes mellitus	Not stated/unclear
Subgroup 3: People with non-alcoholic fatty liver disease	Not stated/unclear
Subgroup 4: People with obesity	Not stated/unclear

Subgroup 5: eGFR category at baseline	Not stated/unclear
Subgroup 6: Albuminuria category at baseline	Not stated/unclear
Sensitivity analysis category: Enrichment trial status	6) No response criteria
Population subgroups	No additional information.
Comparator	Placebo N=83
	Placebo metformin and placebo miglitol tablets three times a day for 36 weeks.
	Concomitant therapy: No additional information.
	Two additional arms were reported, one received a combination of miglitol and metformin (n=76) while the other received miglitol only (n=82). Neither of these arms were extracted as they were not relevant to the protocol.
Number of participants	324
Duration of follow-up	9 months
Indirectness	No additional information.
Method of analysis	Other Appears to be completers only included in the analysis
Additional	No additional information.
comments	TO GEGINOLIGITICATION OF THE SECOND OF THE S

111.2.1. Metformin (N = 62)

Metformin 500mg three times daily plus miglitol placebo with meals for 36 weeks. Concomitant therapy: No additional information.

111.2.2. Placebo (N = 45)

Placebo metformin and placebo miglitol tablets three times a day for 36 weeks. Concomitant therapy: No additional information.

111.3. Characteristics

111.3.1. Arm-level characteristics

Characteristic	Metformin (N = 62)	Placebo (N = 45)
% Male	n = 45 ; % = 73	n = 27 ; % = 60
Sample size		
Mean age (SD) (years)	58.7 (1.1)	58.5 (1.6)
Mean (SE)		
Ethnicity	n = NR ; % = NR	n = NR ; % = NR
Sample size		
Comorbidities Sample size	n = NR ; % = NR	n = NR ; % = NR
•	n - ND · 0/ - ND	
Presence of frailty	n = NR ; % = NR	n = NR; % = NR
Sample size		
Time since type 2 diabetes diagnosis (years)	7.3 (0.9)	4.5 (0.6)
Mean (SD)		
HbA1c (%)	8.2 (0.1)	7.8 (0.1)
Mean (SE)		
Blood pressure	NR (NR)	NR (NR)
Mean (SD)		
Heart rate	NR (NR)	NR (NR)
Mean (SD)		
Smoking status	n = NR ; % = NR	n = NR ; % = NR
Sample size		
Alcohol consumption	n = NR ; % = NR	n = NR ; % = NR
Sample size		
Presence of severe mental illness	n = NR ; % = NR	n = NR ; % = NR
Sample size		

Characteristic	Metformin (N = 62)	Placebo (N = 45)
People with significant cognitive impairment	n = NR ; % = NR	n = NR ; % = NR
Sample size		
People with a learning disability	n = NR ; % = NR	n = NR ; % = NR
Sample size		
Weight	NR (NR)	NR (NR)
Mean (SD)		
BMI (kg/m2)	30.5 (0.6)	30.8 (0.6)
Mean (SE)		
Number of people with obesity	n = NR ; % = NR	n = NR ; % = NR
Sample size	()	
Cholesterol and lipid levels	NR (NR)	NR (NR)
Mean (SD)	ND (ND)	
Albumin creatinine ratio Mean (SD)	NR (NR)	NR (NR)
eGFR (mL/min/1.73m2)	NR (NR)	
Mean (SD)	NIX (INIX)	NR (NR)
Other antidiabetic medication used	n = NR ; % = NR	n = NR ; % = NR
Sample size		
Blood pressure-lowering medication used	n = NR ; % = NR	n = NR ; % = NR
Sample size		
Statins/lipid-lowering medication used	n = NR ; % = NR	n = NR ; % = NR
Sample size		
Other treatment being received	n = NR ; % = NR	n = NR ; % = NR
Sample size		

112. Wu, 2015

Bibliographic Reference

Wu, W.; Li, Y.; Chen, X.; Lin, D.; Xiang, S.; Shen, F.; Gu, X.; Effect of linagliptin on glycemic control in Chinese patients with newly-diagnosed, drug-naive type 2 diabetes mellitus: A randomized controlled trial; Med Sci Monit; 2015; vol. 21; 2678-2684

112.1. Study details		
Secondary publication of another included study- see primary study for details	NA	
Other publications associated with this study included in review	None	
Trial name / registration number	Not reported	
Study type	Randomised controlled trial (RCT)	
Study location	Zhejiang, China	
Study setting	Community	
Study dates	09/2013 to 01/2014	
Sources of funding	Supported by the National Science Foundation for Young Scholars of China (Grant No.81000356)	
Inclusion criteria	 Aged 18-80 years, inclusive Diagnosis of type 2 diabetes mellitus (WHO classification) Fasting plasma glucose≤13.3 mmol/L HbA1c level 7-10% inclusive at baseline BMI 20-35 kg/m2 inclusive Drug-naive (diet and exercise only) 	
Exclusion criteria	 Type 1 diabetes or secondary diabetes Acute complications of diabetes Myocardial infarction, stroke, unstable angina, or coronary artery bypass graft (CABG) in the past 6 months Congestive heart failure Impaired hepatic function (serum alanine aminotransferase, aspartate aminotransferase or alkaline phosphatase level exceeding twice the upper limit of normal) Thyroid disorders 	

	 Chronic intestinal tract disorders History of acute pancreatitis or pancreatic tumor Fertile women not using contraceptives
Recruitment / selection of participants	Participants recruited from outpatient department of First Affiliated Hospital of Wenzhou Medical University, Zhejiang, China. Randomisation using computer-generated random sequence and sealed envelopes. Double blinded to treatment with assigned medication number enter in case report form and corresponding drug kit given to participant. Kit prepared by pharmacist who had not contact with participants All participants received health education before trial. Visits at weeks 6, 12, 18 and 24 with lab assessments conducted by same lab. Medication adherence was also assessed.
Intervention(s)	 Linagliptin 5 mg once daily Oral linagliptin 5 mg once daily for 24 weeks.
Strata 1:	People without heart failure
People with type 2 diabetes mellitus and heart failure	Exclusion criteria: congestive heart failure
Strata 2: People with atherosclerotic cardiovascular	People without other cardiovascular diseases Exclusion criteria: Myocardial infarction, stroke, unstable angina, or coronary artery bypass graft (CABG) in the past 6 months
diseases	coronary artory bypass gran (c), ibc) in the past of monaris
Strata 3: People with type 2 diabetes mellitus and chronic kidney disease	Not stated/unclear
Strata 4: People with type 2 diabetes mellitus and high cardiovascular risk	Not stated/unclear
Subgroup 1: People with frailty	Not stated/unclear
Subgroup 2: Onset of type 2 diabetes mellitus	Not stated/unclear

Not stated/unclear
Not stated/unclear
Not stated/unclear
Not stated/unclear
5) All treatment naïve Inclusion criteria: drug naive (diabetes treated only with diet and exercise)
 Placebo Matching placebo once daily for 24 weeks.
N=57
24 weeks
Other Not explicitly reported by appears to be completer analysis for efficacy and safety analyses

112.2.1. Linagliptin 5 mg once daily (N = 34)

Oral linagliptin tablet 5 mg once daily for 24 weeks.

112.2.2. Placebo (N = 23)

Matching placebo for 24 weeks.

112.3. Characteristics

112.3.1. Arm-level characteristics

% Male daily (N = 34) 23) Reported % of males is not consistent with reported number of participants n = 17; % = 50 n = 15; % = 65.7 Sample size Mean age (SD) (years) 51.2 (7.5) 52.5 (11) Mean (SD) NR NR Nominal NA (NA) NA (NA) Mean (SD) NA (NA) NA (NA) Systolic blood pressure 133.65 (11.91) 136.6 (13.55) Mean (SD) 81.39 (6.96) 81 (5.69) Mean (SD) NR NR Mean (SD) NR <	7 10.10.10.10.10.10.10.10.10.10.10.10.10.1		
Reported % of males is not consistent with reported number of participants Sample size Mean age (SD) (years) Mean (SD) Ethnicity NR NR NR NR NR NR NR NR NR N	Characteristic		Placebo (N = 23)
Mean age (SD) (years) 51.2 (7.5) 52.5 (11) Mean (SD) NR NR Nominal NA (NA) NA (NA) Mean (SD) NA (NA) NA (NA) Mean (SD) 33.65 (11.91) 136.6 (13.55) Mean (SD) 31.39 (6.96) 31 (5.69) Mean (SD) NR NR NR NR NR NR NR NR NR NR NR	% Male Reported % of males is not consistent with reported number of participants	n = 17; % = 50	·
Section Sect	Sample size		
Mean (SD) Ethnicity NR NR Nominal NR NR Comorbidities NR NR Nominal NR NR Presence of frailty NR NR Nominal NR NR Nominal NR NR Nominal NA NA HbA1c (%) 8 (0.69) 7.97 (0.68) Mean (SD) NA (NA) NA (NA) Mean (SD) NA (NA) NA (NA) Mean (SD) 33.65 (11.91) 136.6 (13.55) Mean (SD) 81.39 (6.96) 81 (5.69) Mean (SD) NR NR Neart rate NR NR Nominal NR NR Nominal NR NR Alcohol consumption NR NR	Mean age (SD) (years)	51.2 (7.5)	52.5 (11)
Nominal Comorbidities Nominal Presence of frailty Nominal Time since type 2 diabetes diagnosis NR Nominal HbA1c (%) Blood pressure (mmHg) NA (NA) Mean (SD) Systolic blood pressure Mean (SD) Diastolic blood pressure NR Nominal NA (NA) NA (Mean (SD)		()
ComorbiditiesNRNRNominalNRNRPresence of frailtyNRNRNominalNRNRNominalNRNRHbA1c (%)8 (0.69)7.97 (0.68)Mean (SD)NA (NA)NA (NA)Blood pressure (mmHg)NA (NA)NA (NA)Mean (SD)133.65 (11.91)136.6 (13.55)Mean (SD)81.39 (6.96)81 (5.69)Mean (SD)NRNRMean (SD)NRNRNominalNRNRNominalNRNRNominalNRNRNominalNRNR	Ethnicity	NR	NR
Nominal NR Presence of frailty NR NR Nominal NR NR Nominal NR NR Nominal NR NR HbA1c (%) 8 (0.69) 7.97 (0.68) Mean (SD) NA (NA) NA (NA) Blood pressure (mmHg) NA (NA) NA (NA) Mean (SD) 133.65 (11.91) 136.6 (13.55) Mean (SD) 81.39 (6.96) 81 (5.69) Mean (SD) NR NR Nominal NR NR Nominal NR NR Nominal NR NR Alcohol consumption NR NR		ND	
Presence of frailty Nominal Time since type 2 diabetes diagnosis NR Nominal HbA1c (%) Mean (SD) Blood pressure (mmHg) NA (NA) Mean (SD) Systolic blood pressure Mean (SD) Diastolic blood pressure Mean (SD) Heart rate NR Nominal Smoking status NR	Comorbidities	INIX	NR
NR Nominal Time since type 2 diabetes diagnosis NR Nominal HbA1c (%) Mean (SD) Blood pressure (mmHg) NA (NA) Mean (SD) Systolic blood pressure Mean (SD) Diastolic blood pressure Mean (SD) Heart rate NR Nominal Smoking status NR	Nominal		
Time since type 2 diabetes diagnosis NR Nominal HbA1c (%) Mean (SD) Blood pressure (mmHg) NA (NA) Mean (SD) Systolic blood pressure 133.65 (11.91) 136.6 (13.55) Mean (SD) Diastolic blood pressure 81.39 (6.96) Mean (SD) Heart rate NR NR NR NR NR NR NR NR NR N	Presence of frailty	NR	NR
NR Nominal NR NR	Nominal		
HbA1c (%) 8 (0.69) 7.97 (0.68) Mean (SD) NA (NA) NA (NA) Mean (SD) 133.65 (11.91) 136.6 (13.55) Mean (SD) 81.39 (6.96) 81 (5.69) Mean (SD) NR NR Mean (SD) NR NR Nominal NR NR Nominal NR NR Nominal NR NR Nominal NR NR	Time since type 2 diabetes diagnosis	NR	NR
Mean (SD)		8 (0.69)	
Blood pressure (mmHg) NA (NA) NA (NA) Mean (SD) 133.65 (11.91) 136.6 (13.55) Mean (SD) 81.39 (6.96) 81 (5.69) Mean (SD) NR NR Nominal NR NR Nominal NR NR Nominal NR NR Nominal NR NR		(6.66)	7.97 (0.68)
NA (NA)	` '		
Systolic blood pressure 133.65 (11.91) 136.6 (13.55) Mean (SD) 81.39 (6.96) 81 (5.69) Mean (SD) NR NR Nominal NR NR Nominal NR NR Nominal NR NR Alcohol consumption NR NR	Blood pressure (mmHg)	NA (NA)	NA (NA)
136.6 (13.55) Mean (SD)	Mean (SD)		
Mean (SD) Diastolic blood pressure Mean (SD) Mean (SD) Heart rate NR Nominal Smoking status NR Nominal Alcohol consumption 81.39 (6.96) 81 (5.69) 81 (5.69) NR NR NR	Systolic blood pressure	133.65 (11.91)	136 6 (13 55)
Mean (SD) Heart rate NR Nominal Smoking status NR Nominal Alcohol consumption NR NR NR NR NR	Mean (SD)		100.0 (10.00)
Mean (SD) Heart rate NR Nominal Smoking status Nominal Alcohol consumption NR NR NR NR NR NR	Diastolic blood pressure	81.39 (6.96)	91 (5 60)
Heart rate NR Nominal Smoking status NR Nominal Alcohol consumption NR NR NR NR	Mean (SD)		01 (3.09)
Nominal Smoking status NR Nominal Alcohol consumption NR NR	Heart rate	NR	ND
Smoking status NR Nominal Alcohol consumption NR NR	Nominal		NK
NR Nominal Alcohol consumption NR NR		NR	
Alcohol consumption NR NR	•		NR
NR NR		ND	
Nominal	Alconol consumption	NK	NR
	Nominal		

Characteristic	Linagliptin 5 mg once	Placebo (N =
Durant of a community of the community o	daily (N = 34)	23)
Presence of severe mental illness	NR	NR
Nominal		
People with significant cognitive impairment	NR	NR
Nominal		
People with a learning disability	NR	NR
Nominal		
Weight (kg)	65.24 (8.45)	67.05 (8.12)
Mean (SD)		,
BMI (kg/m2)	24.11 (2.28)	24.37 (2.09)
Mean (SD)		21.07 (2.00)
Number of people with obesity	NR	NR
Nominal		IVIX
Cholesterol and lipid levels (mmol/L)	NA (NA)	NIA (NIA)
Mean (SD)		NA (NA)
Total cholesterol	4.41 (0.72)	4.07.(0.70)
Mean (SD)		4.67 (0.79)
HDL-cholesterol	1.07 (0.12)	4.00 (0.45)
Mean (SD)		1.09 (0.15)
LDL-cholesterol	2.46 (0.56)	
Mean (SD)		2.63 (0.6)
Triglycerides	1.71 (0.59)	
	,	1.75 (0.43)
Mean (SD) Albumin creatinine ratio	NR	
		NR
Nominal eGFR (mL/min/1.73m2)	NR	
·	IVIX	NR
Nominal Other antidiabetic medication used	ND	
	NR	NR
Nominal	ND	
Blood pressure-lowering medication used	NR	NR

Characteristic	Linagliptin 5 mg once daily (N = 34)	Placebo (N = 23)
Nominal		
Statins/lipid-lowering medication used	NR	NR
Nominal		
Other treatment being received	NR	NR
Nominal		

113. Xu, 2015

Bibliographic Reference

Xu, W.; Bi, Y.; Sun, Z.; Li, J.; Guo, L.; Yang, T.; Wu, G.; Shi, L.; Feng, Z.; Qiu, L.; Li, Q.; Guo, X.; Luo, Z.; Lu, J.; Shan, Z.; Yang, W.; Ji, Q.; Yan, L.; Li, H.; Yu, X.; Li, S.; Zhou, Z.; Lv, X.; Liang, Z.; Lin, S.; Zeng, L.; Yan, J.; Ji, L.; Weng, J.; Comparison of the effects on glycaemic control and betacell function in newly diagnosed type 2 diabetes patients of treatment with exenatide, insulin or pioglitazone: a multicentre randomized parallel-group trial (the CONFIDENCE study); J Intern Med; 2015; vol. 277 (no. 1); 137-50

	tudy details
Secondary publication of another included study- see primary study for details	NA
Other publications associated with this study included in review	None
Trial name / registration number	CONFIDENCE/NCT01147627
Study type	Randomised controlled trial (RCT)
Study location	China (25 university-affiliated hospitals in 13 provinces)
Study setting	Community
Study dates	08/2010 to 08/2012
Sources of funding	Funded by Key Projects of Clinical Disciplines of Hospitals Affiliated to Ministry of Health from Ministry of Health of the People's Republic of China, the National Science Fund for Distinguished Young Scholars (81025005), investigator-initiated trial research funds from Eli Lilly and Co. and Amylin Pharmaceuticals, Inc., and the 5010 Project of Sun Yat-sen University.
Inclusion criteria	 Aged ≥30 to ≤70 years Newly-diagnosed type 2 diabetic patients Treatment-naive to glucose-lowering drugs HbA1c ≥7.0 to ≤10.0% BMI ≥20 to ≤35 kg/m2 Stable body weight (≤10% variation)≥3 months prior to screening Female patients of reproductive age should practice a reliable method of birth control throughout study

Exclusion criteria

- Pregnancy
- Acute or severe chronic diabetic complications
- Congestive heart failure (NYHA grade 3-4)
- Severe gastrointestinal disease
- Severe osteoporosis or history of pathological fracture, or use of bisphosphonates preparation
- Other severe intercurrent illness
- Serum aminotransferase (ALT and AST) level higher than 2 times of the upper normal limits and/or serum creatinine≥133µmol/L (1.5mg/dL)
- Tested positive for glutamic acid decarboxylase antibody
- Use of weight loss drugs, corticosteroids, drugs known to affect gastrointestinal motility, transplantation medications, or any investigational drug
- History of pancreatitis
- Serum triglyceride ≥ 5.0 mmol/L

Recruitment / selection of participants

Participants recruited from 25 university-affiliated hospitals in 13 Chinese provinces. Participants randomised 1:1:1 to groups using randomisation list generated by SAS software with allocation by secure Oracle-based interactive web-based response system in accordance with list. Participants followed up every 4 weeks for 12 weeks, then every 12 weeks until week 48. At each visit, anthropomorphic data, adverse events and hypoglycaemia episodes recorded and HbA1c, FPG and 2-h postprandialglucose (PPG) after mixed-meal test measured. Telephone calls scheduled at weeks 16, 20, 28, 32, 40 and 44 to collect information and provide guidance. All participants received diabetes information and lifestyle counselling at enrolment, reinforced during study. Baseline assessments repeated at week 48, with participants instructed to stop all antihyperglycaemic therapy 2 days beforehand to avoid acute drug effects on collected data. HbA1c assessed centrally at the Diabetes Centre of the Third Affiliated Hospital of Sun Yat-sen University, and insulin and proinsulin levels were measured centrally at the Beijing North Institute of Biological Technology. Remaining biochemical variables assessed locally at participating centres.

Intervention(s)

Exenatide 10 mcg twice daily

Subcutaneous injection of exenatide 5 mcg twice daily for 4 weeks then exenatide 10 mcg twice daily for remaining 44 weeks. Participants instructed to reduce dose to 5 mcg twice daily if experiencing frequent hypoglycaemia or could not tolerate adverse events.

Strata 1: People with type 2 diabetes mellitus and heart failure

Not stated/unclear

Exclusion criteria: NYHA grade III and IV. Trial may include participants with NYHA grade 2.

Strata 2: People with atherosclerotic cardiovascular diseases

Not stated/unclear

Strata 3: People with type 2 diabetes mellitus and chronic kidney disease	Not stated/unclear
Strata 4: People with type 2 diabetes mellitus and high cardiovascular risk	Not stated/unclear
Subgroup 1: People with frailty	Not stated/unclear
Subgroup 2: Onset of type 2 diabetes mellitus	Not stated/unclear
Subgroup 3: People with non-alcoholic fatty liver disease	Not stated/unclear
Subgroup 4: People with obesity	Not stated/unclear
Subgroup 5: eGFR category at baseline	Not stated/unclear
Subgroup 6: Albuminuria category at baseline	Not stated/unclear
Sensitivity analysis category: Enrichment trial status	5) All treatment naïve Inclusion criteria: treatment-naive to glucose lowering drugs and newly diagnosed type 2 diabetes
Population subgroups	
Comparator	 Premixed biphasic insulin lispro 25%/insulin lispro protamine 75% twice daily Pioglitazone 45 mg once daily Insulin injected twice daily at initial dose of 0.4 IU/kg, 50% 15 min before breakfast and 50% 15 min before dinner. Doses self-titrated after this
	Dieakiasi and 50% 15 min defore dinner. Doses self-titrated after this

	based on self-monitored blood glucose levels. Pioglitazone administered at 30 mg once daily for 4 weeks, increasing to 45 mg once daily for remaining 44 weeks.
Number of participants	N=416
Duration of follow-up	48 weeks
Method of analysis	Per protocol Per protocol population conducted for efficacy analysis. Reports conducting analysis on ITT population but data not reported because results were similar to analysis on PP population. Modified ITT Safety analysis conducted on ITT population - all participants randomised who received at least one dose of study drug.

113.2.1. Exenatide 10 mcg twice daily (N = 142)

Subcutaneous injection of exenatide 5 mcg twice daily for 4 weeks, increased to 10 mcg twice daily for remaining 44 weeks.

113.2.2. Biphasic insulin lispro 25%/insulin lispro protamine 75% twice daily (N = 138)

Subcutaneous injection of premixed biphasic insulin lispro 25%/insulin lispro protamine 75% twice daily titrated based on self-monitored blood glucose levels for 48 weeks.

113.2.3. Pioglitazone 45 mg once daily (N = 136)

Oral pioglitazone tablets 30 mg once daily for 4 weeks, increased to 45 mg once daily for remaining 44 weeks.

113.3. Characteristics

113.3.1. Arm-level characteristics

113.3.1.	Attil-level characteristics			
Characteristic	Exenatide 10 mcg twice daily (N = 142)	Biphasic insulin lispro 25%/insulin lispro protamine 75% twice daily (N = 138)	Pioglitazone 45 mg once daily (N = 136)	
% Male Sample size	n = 98 ; % = 69	n = 85; % = 61.6	n = 83; % = 61	
Mean age (SD) (years)	50 (0.8)	51 (0.8)	50 (0.8)	
Mean (SD)				
HbA1c (%)	8 (0.1)	8.1 (0.1)	8 (0.1)	
Mean (SE)				
Blood pressure Mean (SE)	NA (NA)	NA (NA)	NA (NA)	
	400 (4)			
Systolic blood pressure	126 (1)	124 (1)	125 (1)	
Mean (SE)				
Diastolic blood pressure	80 (1)	79 (1)	80 (1)	
Mean (SE)				
Weight (kg)	72.6 (1)	70.3 (1)	71.2 (1)	
Mean (SE)				
BMI (kg/m2)	26.1 (0.3)	25.6 (0.3)	25.8 (0.3)	
Mean (SE)	NIA (NIA)			
Cholesterol and lipid levels (mmol/L)	NA (NA)	NA (NA)	NA (NA)	
Mean (SE)				
Total cholesterol	5.1 (0.1)	5.2 (0.1)	5.3 (0.1)	
Mean (SE)				
HDL-cholesterol	1.14 (0.02)	1.21 (0.03)	1.18 (0.02)	
Mean (SE)				

Characteristic	Exenatide 10 mcg twice daily (N = 142)	Biphasic insulin lispro 25%/insulin lispro protamine 75% twice daily (N = 138)	Pioglitazone 45 mg once daily (N = 136)
LDL-cholesterol Mean (SE)	3.2 (0.1)	3.2 (0.1)	3.2 (0.1)
, ,			
Triglycerides	1.9 (0.1)	1.9 (0.1)	2 (0.1)
Mean (SE)			

114. Yamada, 2020

Bibliographic Reference

Yamada, Y.; Katagiri, H.; Deenadayalan, S.; Navarria, A.; Nishijima, K.; Seino, Y.; Fukushima, Y.; Hamamoto, Y.; Hisatomi, A.; Ide, Y.; Inoue, S.; Kawada, T.; Kim, H.; Kiyosue, A.; Matoba, K.; Matsuoka, O.; Nishimura, H.; Noguchi, M.; Osonoi, T.; Sawada, S.; Shibasaki, Y.; Shin, K.; Doseresponse, efficacy, and safety of oral semaglutide monotherapy in Japanese patients with type 2 diabetes (PIONEER 9): a 52-week, phase 2/3a, randomised, controlled trial; Lancet Diabetes Endocrinol; 2020; vol. 8 (no. 5); 377-391

117.1. 0	tudy details		
Secondary publication of another included study- see primary study for details	NA		
Other publications associated with this study included in review	None		
Trial name / registration number	PIONEER 9/NCT03018028		
Study type	Randomised controlled trial (RCT)		
Study location	Japan (16 sites)		
Study setting	Community		
Study dates	01/2017 to 07/2017		
Sources of funding	Funded by Novo Nordisk.		
Inclusion criteria	 Japanese adults aged ≥20 years or older Diagnosis of type 2 diabetes ≥30 days before screening Managed with diet and exercise only or received oral glucose-lowering drug monotherapy (at stable daily dose less than or equal to a half of the maximum approved dose in Japan) ≥30 days before screening HbA1c level 7·0-10·0% inclusive for diet and exercise only or 6·5-9·5% inclusive for those on oral glucose-lowering drug monotherapy 		
Exclusion criteria	 Female who was pregnant, breast-feeding or had intention to become pregnant, or not using adequate contraception 		

- Any disorder that might affect safety of participant or compliance with trial protocol
- Treatment ≤90 before screening with once-weekly GLP-1 receptor agonist, once-weekly DPP-4 inhibitor, or thiazolidinedione
- Severe renal impairment (eGFR <30 mL/min per 1·73 m²)
- Proliferative retinopathy or maculopathy requiring acute treatment
- Family or personal history of multiple endocrine neoplasia type 2 or medullary thyroid carcinoma
- History of pancreatitis (acute or chronic)
- History of major surgical procedures involving stomach potentially affecting absorption of trial product (e.g. subtotal and total gastrectomy, sleeve gastrectomy, gastric bypass surgery).
- Any of: myocardial infarction, stroke or hospitalisation for unstable angina, or transient ischaemic attack within the past 180 days prior to the day of screening and randomisation
- Subjects classified as being in New York Heart Association Class
 IV
- Planned coronary, carotid or peripheral artery revascularisation known on the day of screening
- Alanine aminotransferase >2.5 x upper limit of normal
- Renal impairment (eGFR<30 mL/min/1·73 m2) as per Chronic Kidney Disease Epidemiology collaboration
- Treatment with once-weekly glucagon-like peptide-1 receptor agonist, once-weekly DPP-4 inhibitor or thiazolidinedione in a period of 90 days before the day of screening
- Treatment with any medication for the indication of diabetes or obesity other than stated in inclusion criteria in a period of 60 days before the day of screening (with exception of short-term insulin treatment for acute illness for a total of ≤14 days)
- Proliferative retinopathy or maculopathy requiring acute treatment.
 Verified by fundus photography or dilated fundoscopy performed within 90 days prior to randomisation
- History or presence of malignant neoplasms within the last 5 years (except basal and squamous cell skin cancer and in situ carcinomas)
- Initiation of glucose-lowering medication between the day of screening and the day of randomisation

Recruitment / selection of participants

Participants recruited from 16 sites (clinics and university hospitals) in Japan. They were randomly assigned 1:1:1:1:1 to one of 5 arms using trial-specific interactive web response system that assigned treatment codes, stratified by pre-trial treatment at screening (without oral glucose lowering drug monotherapy; with monotherapy). Participants receiving diet and exercise only screened over 2 week period; those receiving monotherapy had 8 week screening/washout period.

Intervention(s)

- Semaglutide 14 mg once daily
- Semaglutide 7 mg once daily
- Semaglutide 3 mg once daily

	Participants received treatment for 52 weeks with a 5 week post-treatment follow up visit. Oral semaglutide initiated with 3 mg once daily tablet dose, escalated after 4 weeks to 7 mg for 7 mg and 14 mg arms, and an additional 7 mg 4 weeks after this for 14 mg arm. Tablets taken with ≤120 ml water in morning, in fasting state, and minimum of 30 min before first meal of day or other oral medication. Rescue medication could be offered at week 8+ for participants with persistent hypoglycaemia (>240 mg/dL from weeks 8-13, or >200 mg/dL week 14+, or if HbA1c>8.5% at week 26+). Rescue medication (excluding use of GLP-1RA, DPP-4 inhibitors, and amylin analogues) offered at investigator's discretion and participants could continue in trial.
Strata 1: People with	Not stated/unclear
type 2 diabetes mellitus and heart failure	Exclusion criteria: NYHA Class IV
Strata 2:	People without other cardiovascular diseases
People with atherosclerotic cardiovascular diseases	Exclusion criteria: myocardial infarction, stroke or hospitalisation for unstable angina, or transient ischaemic attack within the past 180 days prior to the day of screening and randomisation;
Strata 3: People with type 2 diabetes mellitus and chronic kidney disease	Not stated/unclear
Strata 4: People with type 2 diabetes mellitus and high cardiovascular risk	Not stated/unclear
Subgroup 1: People with frailty	Not stated/unclear
Subgroup 2: Onset of type 2 diabetes mellitus	Not stated/unclear
Subgroup 3: People with non-alcoholic fatty liver disease	Not stated/unclear

Not stated/unclear
Not stated/uniclear
eGFR ≥30mL/min/1.73m2
Exclusion criteria: eGFR<30 mL/min per 1.73 m2
Not stated/unclear
7) Mixed population
Participants were drug-naive (diet and exercise only) or if on oral glucose-lowering monotherapy (for more than 30 days before screening) then had 8 week screening and washout period.
Placebo for semaglutideLiraglutide 0.9 mg once daily
Placebo for semaglutide was visually identical tablets and packaging as used for semaglutide arms. Placebo tablets taken with ≤120 ml water in morning, in fasting state, and minimum of 30 min before first meal of day or other oral medication. Liraglutide initiated with 0.3 mg daily subcutaneous injection, escalated 0.3 mg after 1 and 2 weeks until 0.9 mg once daily dose achieved, and administered at fixed time in morning or evening irrespective of meal times. Rescue medication strategy same as for interventions.
N=243
52 weeks
ITT
All randomised participants regardless of discontinuation or use of rescue medication with imputation of missing data for 6 groups (one group of those who discontinued treatment or initiated rescue medication; 5 groups (1 for each randomised treatment arm including all those in each group who had not initiated rescue medication). Missing data addressed using multiple imputation using assumption that participants at week 26 or week 52 were similar in terms of treatment arm and adherence, or rescue medication status.
Trial specifies 2 estimands: trial product and treatment policy. Trial product estimand is evaluation of treatment effect for all randomly assigned participants under assumption that all remained on trial product for entire length of trial and didn't use rescue medication. Model includes all post-baseline measurements collected at scheduled visits to week 26 or week 52 excluding those who took rescue medication. Treatment policy estimand is evaluation of treatment effect for all randomly assigned participants regardless of whether they discontinued or used rescue medication (that is, it is an ITT analysis).

114.2.1. Semaglutide 14 mg once daily (N = 48)

Oral semaglutide tablet 14 mg once daily for 52 weeks.

114.2.2. Semaglutide 7 mg once daily (N = 49)

Oral semaglutide tablet 7 mg once daily for 52 weeks.

114.2.3. Semaglutide 3 mg once daily (N = 49)

Oral semaglutide tablet 3 mg once daily for 52 weeks.

114.2.4. Liraglutide 0.9 mg once daily (N = 48)

Subcutaneous injection of liraglutide 0.9 mg once daily for 52 weeks.

114.2.5. Placebo (N = 49)

Placebo tablet once daily for 52 weeks, using visually identical tablet and packaging as used for oral semaglutide.

114.3. Characteristics

114.3.1. Arm-level characteristics

Characteristic	Semaglutide 14 mg once daily (N = 48)	Semaglutide 7 mg once daily (N = 49)	Semaglutide 3 mg once daily (N = 49)	Liraglutide 0.9 mg once daily (N = 48)	Placebo (N = 49)
% Male Sample size	n = 40 ; % = 83	n = 36 ; % = 73	n = 36 ; % = 73	n = 39 ; % = 81	n = 40 ; % = 82
Mean age (SD) (years) Mean (SD)	61 (9)	60 (10)	58 (9)	59 (10)	59 (9)
Ethnicity Nominal	NR	NR	NR	NR	NR
Comorbidities Nominal	NR	NR	NR	NR	NR

Characteristic	Semaglutide 14 mg once daily (N = 48)	Semaglutide 7 mg once daily (N = 49)	Semaglutide 3 mg once daily (N = 49)	Liraglutide 0.9 mg once daily (N = 48)	Placebo (N = 49)
Presence of frailty	NR	NR	NR	NR	NR
Nominal					
Time since type 2 diabetes diagnosis (years) Mean (SD)	7.9 (5.9)	7.4 (5.6)	7.4 (5.5)	6.7 (5.2)	8.4 (6)
	9 (0 0)				
HbA1c (%) Mean (SD)	8 (0.9)	8.3 (1)	8.1 (0.8)	8.3 (0.8)	8.3 (1.1)
Blood pressure (mmHg)	NA (NA)	NA (NA)	NA (NA)	NA (NA)	NA (NA)
Mean (SD)					
Systolic blood pressure	127 (13)	129 (12)	127 (14)	128 (13)	128 (13)
Mean (SD)					
Diastolic blood pressure	76 (9)	80 (10)	76 (8)	81 (11)	78 (12)
Mean (SD)					
Heart rate	NR	NR	NR	NR	NR
Nominal					
Smoking status	NR	NR	NR	NR	NR
Nominal	ND				
Alcohol consumption	NR	NR	NR	NR	NR
Nominal					
Presence of severe mental illness	NR	NR	NR	NR	NR
Nominal					
People with significant cognitive impairment	NR	NR	NR	NR	NR
Nominal					

Characteristic	Semaglutide 14 mg once daily (N = 48)	Semaglutide 7 mg once daily (N = 49)	Semaglutide 3 mg once daily (N = 49)	Liraglutide 0.9 mg once daily (N = 48)	Placebo (N = 49)
People with a learning disability	NR	NR	NR	NR	NR
Nominal					
Weight (kg)	68 (13)	71.3 (10.8)	71.4 (14.3)	74.7 (15.4)	70.3 (12.4)
Mean (SD)	24.7 (4.1)				()
BMI (kg/m2) Mean (SD)	24.7 (4.1)	26.3 (3.5)	26.5 (4.6)	26.9 (4.8)	25.1 (3.9)
Number of people with obesity	NR	NR	NR	NR	NR
Nominal					
Cholesterol and lipid levels	NR	NR	NR	NR	NR
Nominal					
Albumin creatinine ratio	NR	NR	NR	NR	NR
Nominal					
eGFR (mL/min/1.73m2)	94 (13)	96 (14)	99 (12)	99 (9)	96 (12)
Mean (SD)					
Other antidiabetic medication used Pre-trial glucose- lowering therapy	n = NA ; % = NA	n = NA ; % = NA	n = NA ; % = NA	n = NA ; % = NA	n = NA ; % = NA
Sample size					
Metformin	n = 8 ; % = 17	n = 4; % = 8	n = 7; % = 14	n = 8 ; % =	n = 9; % = 18
Sample size	0.0/.40				10
DPP-4 inhibitor Sample size	n = 6; % = 13	n = 10 ; % = 20	n = 5; % = 10	n = 2; % = 4	n = 7; % = 14
SGLT2 inhibitor	n = 1; % = 2	m = 0 · 0/ 0	- F.O/ 40	n = 4 · 0/ 0	0 · 0/
Sample size		11 = 3; % = 6	n = 5; % = 10	11 = 4 ; % = 8	n = 2; % = 4

Characteristic	Semaglutide 14 mg once daily (N = 48)	Semaglutide 7 mg once daily (N = 49)	Semaglutide 3 mg once daily (N = 49)	Liraglutide 0.9 mg once daily (N = 48)	Placebo (N = 49)
Alpha- glucosidase inhibitor Sample size	n = 3; % = 6	n = 2; % = 4	n = 1; % = 2	n = 3; % = 6	n = 1; % = 2
·					
Sulfonylurea	n = 0; % = 0	n = 0; % = 0	n = 1; % = 2	n = 1; % = 2	n = 0; % = 0
Sample size					·
Blood pressure- lowering medication used	NR	NR	NR	NR	NR
Nominal					
Statins/lipid- lowering medication used	NR	NR	NR	NR	NR
Nominal					
Other treatment being received	NR	NR	NR	NR	NR
Nominal					

115. Yamanouchi, 2005

Bibliographic Reference

Yamanouchi, T.; Sakai, T.; Igarashi, K.; Ichiyanagi, K.; Watanabe, H.; Kawasaki, T.; Comparison of metabolic effects of pioglitazone, metformin, and glimepiride over 1 year in Japanese patients with newly diagnosed Type 2 diabetes; Diabet Med; 2005; vol. 22 (no. 8); 980-5

110.11. 0	tudy details
Secondary publication of another included study- see primary study for details	No additional information.
Other publications associated with this study included in review	No additional information.
Trial name / registration number	No additional information.
Study location	Japan.
Study setting	Outpatient follow-up.
Study dates	No additional information.
Sources of funding	No additional information.
Inclusion criteria	People with a short duration of Type 2 diabetes; had not received an oral hypoglycaemic agent or a lipid drug; treated with diet and exercise alone for at least 3 months, including the 1 month for baseline measurements before the study (observation period) after which they had an HbA1c at least 7.0% and a fasting plasma glucose at least 7.78 mmol/L; BMI 22-35kg/m2.
Exclusion criteria	People who had unstable or rapidly progressive diabetic retinopathy, nephropathy or neuropathy; people with liver dysfunction (AST and ALT >1.5 x the upper limit of normal); impaired kidney function (serum creatinine >133 micromol/L); anaemia; myocardial infarction, angina, congestive heart failure or a documented cerebrovascular accident.
Recruitment / selection of participants	No additional information.
Intervention(s)	Pioglitazone N=38

	Pioglitazone 30-45mg/day. Concomitant therapy: Antihypertensive drugs or other concurrent
	treatments, including dietary regimens, remained unchanged throughout the study.
Strata 1: People with type 2 diabetes mellitus and heart failure	People without heart failure
Strata 2: People with atherosclerotic cardiovascular diseases	People without other cardiovascular diseases
Strata 3: People with type 2 diabetes mellitus and chronic kidney disease	People without chronic kidney disease
Strata 4: People with type 2 diabetes mellitus and high cardiovascular risk	People at higher risk of developing cardiovascular disease Based on systolic blood pressure, triglycerides and presence of diabetes.
Subgroup 1: People with frailty	Not stated/unclear
Subgroup 2: Onset of type 2 diabetes mellitus	Not stated/unclear
Subgroup 3: People with non-alcoholic fatty liver disease	People without non-alcoholic fatty liver disease
Subgroup 4: People with obesity	Not stated/unclear

Subgroup 5: eGFR category at baseline	Not stated/unclear
Subgroup 6: Albuminuria category at baseline	Not stated/unclear
Sensitivity analysis category: Enrichment trial status	5) All treatment naïve
Population subgroups	No additional information.
Comparator	Metformin N=39
	Metformin 750mg/day.
	Concomitant therapy: Antihypertensive drugs or other concurrent treatments, including dietary regimens, remained unchanged throughout the study.
	Glimepiride N=37
	Glimepiride 1.0-2.0mg/day.
	Concomitant therapy: Antihypertensive drugs or other concurrent treatments, including dietary regimens, remained unchanged throughout the study.
Number of participants	114
Duration of follow-up	12 months.
Indirectness	No additional information.
Method of analysis	Not stated/unclear
Additional comments	No additional information.

115.2.1. Pioglitazone (N = 38)

Pioglitazone 30-45mg/day. Concomitant therapy: Antihypertensive drugs or other concurrent treatments, including dietary regimens, remained unchanged throughout the study.

115.2.2. Metformin (N = 39)

Metformin 750mg/day. Concomitant therapy: Antihypertensive drugs or other concurrent treatments, including dietary regimens, remained unchanged throughout the study.

115.2.3. Glimepiride (N = 37)

Glimepiride 1.0-2.0mg/day. Concomitant therapy: Antihypertensive drugs or other concurrent treatments, including dietary regimens, remained unchanged throughout the study.

115.3. Characteristics

115.3.1. Arm-level characteristics

Characteristic	Pioglitazone (N = 38)	Metformin (N = 39)	Glimepiride (N = 37)
% Male	n = 20 ; % = 53	n = 19 ; % = 49	n = 18 ; % = 49
Sample size			
Mean age (SD) (years)	55.2 (9.2)	54.7 (9.8)	55.6 (9.3)
Mean (SD)			
Ethnicity	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
Sample size		IVIX	
Comorbidities	n = NR ; % = NR	·	n = NR ; % = NR
Sample size		NR	
Presence of frailty	n = NR ; % = NR	n = NR ; % = NR	n = NR ; % = NR
Sample size		IVIX	
Time since type 2 diabetes diagnosis (Months)	3.2 (2.1)	3 (2.5)	3.3 (2.6)
Mean (SD)			

Characteristic	Pioglitazone (N = 38)	Metformin (N = 39)	Glimepiride (N = 37)
HbA1c (%)	10.2 (0.8)	9.9 (0.7)	9.8 (0.7)
Mean (SD)		,	,
Blood pressure (mmHg)	NA (NA)	NA (NA)	NA (NA)
Mean (SD)		,	,
Systolic blood pressure	142.8 (17.1)	143.3 (18.8)	141.3 (21.3)
Mean (SD)		110.0 (10.0)	111.0 (21.0)
Diastolic blood pressure	85.3 (9.8)	86.3 (10.1)	84.9 (7.7)
Mean (SD)		00.0 (10.1)	04.5 (1.1)
Heart rate	NR (NR)	NR (NR)	NR (NR)
Mean (SD)		TVIX (IVIX)	TVIC (IVIC)
Smoking status	n = NR ; % = NR	n = NR ; % =	n = NR ; % = NR
Sample size		NR	11 - INIX , 70 - INIX
Alcohol consumption	n = NA ; % = NA	m - NIA : 0/ -	- NIA : 0/ - NIA
Sample size		n = NA ; % = NA	n = NA ; % = NA
Presence of severe mental illness	n = NA ; % = NA	NIA 0/	NIA O/ NIA
Sample size		n = NA ; % = NA	n = NA ; % = NA
People with significant cognitive	n = NA ; % = NA		
impairment		n = NA ; % = NA	n = NA ; % = NA
Sample size			
People with a learning disability	n = NA ; % = NA	n = NA ; % =	n = NA ; % = NA
Sample size		NA	,
Weight	NA (NA)	NA (NA)	NA (NA)
Mean (SD)		,	,
BMI (kg/m2)	10.2 (0.8)	9.9 (0.7)	9.8 (0.7)
Mean (SD)		(6)	(011)
Number of people with obesity	n = NR ; % = NR	n = NR ; % =	n = NR ; % = NR
Sample size		NR	, 70
Cholesterol and lipid levels	NA (NA)	NA (NA)	NA (NA)
Mean (SD)		(1.0.1)	()

Characteristic	Pioglitazone (N = 38)	Metformin (N = 39)	Glimepiride (N = 37)
Total cholesterol	5.77 (0.57)	5.7 (0.36)	5.89 (0.49)
Mean (SD)			
HDL cholesterol	1.38 (0.12)	1.33 (0.09)	1.35 (0.11)
Mean (SD)			
Triglycerides	2.47 (1.26)	2.31 (1.14)	2.63 (1.37)
Mean (SD)			
Albumin creatinine ratio	NA (NA)	NA (NA)	NA (NA)
Mean (SD)			
eGFR (mL/min/1.73m2)	NA (NA)	NA (NA)	NA (NA)
Mean (SD)			
Other antidiabetic medication used	n = NA ; % = NA	n = NA ; % = NA	n = NA ; % = NA
Sample size			
Blood pressure-lowering medication used	n = NA ; % = NA	n = NA ; % = NA	n = NA ; % = NA
Sample size			
Statins/lipid-lowering medication used	n = NA ; % = NA	n = NA ; % = NA	n = NA ; % = NA
Sample size			
Other treatment being received	n = NA ; % = NA	n = NA ; % = NA	n = NA ; % = NA
Sample size		1 1// 1	

116. Yoon, 2012

Bibliographic Reference

Yoon, K H; Steinberg, H; Teng, R; Golm, G T; Lee, M; O'Neill, E A; Kaufman, K D; Goldstein, B J; Efficacy and safety of initial combination therapy with sitagliptin and pioglitazone in patients with type 2 diabetes: a 54-week study.; Diabetes, obesity & metabolism; 2012; vol. 14 (no. 8); 745-52

Secondary publication of another included study- see primary study for details Other publications associated with this study included in review Trial name / registration number Study type Randomised controlled trial (RCT) Double-blind single-dummy placebo RCT with open-label pioglitazone. International (sites in 14 countries: Brazil, Columbia, Costa Rica, Czech Republic, India, Lithuania, Malaysia, Peru, Philippines, Poland, Puerto Rico, Russian Federation, South Korea, USA) Study setting Study dates Sources of funding Inclusion criteria NOT00397631 NCT00397631 PCOUNTIENT OF THE NOT O
publications associated with this study included in review Trial name / registration number Study type Randomised controlled trial (RCT) Double-blind single-dummy placebo RCT with open-label pioglitazone. Study location International (sites in 14 countries: Brazil, Columbia, Costa Rica, Czech Republic, India, Lithuania, Malaysia, Peru, Philippines, Poland, Puerto Rico, Russian Federation, South Korea, USA) Study setting Community Study dates 09/2007 to 01/2009 Sources of funding Inclusion • Aged≥18 years
Tegistration number Study type Randomised controlled trial (RCT) Double-blind single-dummy placebo RCT with open-label pioglitazone. Study location International (sites in 14 countries: Brazil, Columbia, Costa Rica, Czech Republic, India, Lithuania, Malaysia, Peru, Philippines, Poland, Puerto Rico, Russian Federation, South Korea, USA) Study setting Study dates O9/2007 to 01/2009 Sources of funding Inclusion Aged≥18 years
Double-blind single-dummy placebo RCT with open-label pioglitazone. Study location International (sites in 14 countries: Brazil, Columbia, Costa Rica, Czech Republic, India, Lithuania, Malaysia, Peru, Philippines, Poland, Puerto Rico, Russian Federation, South Korea, USA) Study setting Community Study dates 09/2007 to 01/2009 Sources of funding Inclusion • Aged≥18 years
Republic, India, Lithuania, Malaysia, Peru, Philippines, Poland, Puerto Rico, Russian Federation, South Korea, USA) Study setting Study dates O9/2007 to 01/2009 Sources of funding Inclusion Aged≥18 years
Study dates 09/2007 to 01/2009 Sources of funding Inclusion
Sources of funding Inclusion Funded by Merck Sharp & Dohme LLC • Aged≥18 years
funding Inclusion • Aged≥18 years
5 7
 Not receiving oral antihyperglycaeic agents prior to screening Not received anti-hyperglycaemic agent for ≥4 cumulative weeks in prior 2 years and none within 4-mo screening HbA1c level 8-12% inclusive
 Fasting fingerstick glucose level <7.2 mmol/l or >17.8 mmol./l at randomisation Diagnosis of type 2 diabetes Unstable cardiac disease

	 Significant renal impairment (estimated creatinine clearance <60 ml/min) or elevated ALT or AST levels (>2 times upper limit of normal)
Recruitment / selection of participants	Recruited from sites in 14 countries. Participants received diet and exercise counselling and then started 2 week single-blind placebo run-in period. Participants with adequate compliance (≥85% via tablet counts) during run-in period randomised 1:1 ratio to arms. Lab assessments conducted at central laboratory by technicians blinded to treatment. Thirty-week Extension study to original 24-week study (Yoon KH, Shockey GR, Teng R, Golm GT, Thakkar PR, Meehan AG, Williams-Herman DE, Kaufman KD, Amatruda JM, Steinberg H. Effect of initial combination therapy with sitagliptin, a dipeptidyl peptidase-4 inhibitor, and pioglitazone on glycemic control and measures of β-cell function in patients with type 2 diabetes. International journal of clinical practice. 2011 Feb;65(2):154-64.).
Intervention(s)	 Sitagliptin 100 mg once daily + Pioglitazone 30 mg once daily Participants in this arm took one oral sitagliptin tablet, double blinded, and one open-label pioglitazone tablet, both once daily for 24 weeks.
Cointervention	Placebo for sitagliptin in comparator arm only
Strata 1: People with type 2 diabetes mellitus and heart failure	Not stated/unclear
Strata 2: People with atherosclerotic cardiovascular diseases	Not stated/unclear
Strata 3: People with type 2 diabetes mellitus and chronic kidney disease	Not stated/unclear
Strata 4: People with type 2 diabetes mellitus and high cardiovascular risk	Not stated/unclear
Subgroup 1: People with frailty	Not stated/unclear

Subgroup 2: Nonset of type 2 diabetes mellitus	Not stated/unclear
Subgroup 3: Neople with non-alcoholic fatty liver disease	Not stated/unclear
Subgroup 4: Neople with obesity	Not stated/unclear
Subgroup 5: NegFR category at baseline	Not stated/unclear
Subgroup 6: N Albuminuria category at baseline	Not stated/unclear
analysis category: Enrichment	7) Mixed population Inclusion criteria: no oral anti-hyperglycaemic agent prior to screening, with less than 4 weeks cumulative prior treatment in prior 2 years and none within 4-mo of screening visit.
	Pioglitazone 30 mg once daily Oral pioglitazone tablet 30 mg once daily with matching placebo (for sitagliptin) for 24 weeks. Pioglitazone tablet was open-label.
Number of participants	N=520 in original trial; N=317 in extension trial
Duration of 5 follow-up	54 weeks
Indirectness	None
analysis E	Modified ITT Efficacy analyses conducted on all randomised participants who received at least one study drug dose, and had both baseline and at least one post-baseline assessment. Safety analysis conducted on all randomised participants. Missing data not imputed.
Additional comments	

116.2.1. Pioglitazone 30 mg + Sitagliptin 100 mg once daily (N = 261) Oral sitagliptin 100 mg + Oral pioglitazone 30 mg once daily for 24 weeks.

116.2.2. Pioglitazone 30 mg once daily (N = 259)

Oral pioglitazone 30 mg once daily with matching placebo for 24 weeks.

116.3. Characteristics

116.3.1. Arm-level characteristics

110.3.1. Affil-level characteristics			
Characteristic	Pioglitazone 30 mg + Sitagliptin 100 mg once daily (N = 261)	Pioglitazone 30 mg once daily (N = 259)	
% Male	n = 86 ; % = 52.4	n = 90 ; % = 58.8	
Sample size			
Mean age (SD) (years)	51.4 (10)	52.3 (11.5)	
Mean (SD)			
Ethnicity	n = NA ; % = NA	n = NA ; % = NA	
Sample size			
Asian	n = 42 ; % = 25.6	n = 41; % = 26.8	
Sample size	7 0/ 40		
Black	n = 7; % = 4.3	n = 3; % = 2	
Sample size			
Hispanic or Latino	n = 64 ; % = 39	n = 58 ; % = 37.9	
Sample size			
Not hispanic or latino	n = 100 ; % = 61	n = 95; % = 62.1	
Sample size	00 0/ 10 1		
Other	n = 22 ; % = 13.4	n = 24 ; % = 15.7	
Sample size			
White	n = 93 ; % = 56.7	n = 85; % = 55.6	
Sample size			
Comorbidities	NR	NR	
Nominal			
Presence of frailty	NR	NR	
Nominal			

Characteristic	Pioglitazone 30 mg + Sitagliptin 100 mg once daily (N = 261)	Pioglitazone 30 mg once daily (N = 259)
Time since type 2 diabetes diagnosis (years)	2.6 (4)	1.6 (3.7)
Mean (SD)		
Smoking status	NR	NR
Nominal		
Alcohol consumption Nominal	NR	NR
	ND	
Presence of severe mental illness	NR	NR
Nominal		
People with significant cognitive impairment	NR	NR
Nominal		
Number of people with obesity	NR	NR
Nominal		
Albumin creatinine ratio Nominal	NR	NR
Other antidiabetic	NR	
medication used	INK	NR
Nominal		
Blood pressure-lowering medication used	NR	NR
Nominal		
Statins/lipid-lowering medication used	NR	NR
Nominal		
Other treatment being received	NR	NR
Nominal		

117. Yoon, 2011

Bibliographic Reference

Yoon, K. H.; Shin, J. A.; Kwon, H. S.; Lee, S. H.; Min, K. W.; Ahn, Y. B.; Yoo, S. J.; Ahn, K. J.; Park, S. W.; Lee, K. W.; Sung, Y. A.; Park, T. S.; Kim, M. S.; Kim, Y. K.; Nam, M. S.; Kim, H. S.; Park Ie, B.; Park, J. S.; Woo, J. T.; Son, H. Y.; Comparison of the efficacy of glimepiride, metformin, and rosiglitazone monotherapy in korean drug-naïve type 2 diabetic patients: the practical evidence of antidiabetic monotherapy study; Diabetes Metab J; 2011; vol. 35 (no. 1); 26-33

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Secondary publication of another included study- see primary study for details	No additional information.
Other publications associated with this study included in review	No additional information.
Trial name / registration number	No additional information.
Study type	Randomised controlled trial (RCT)
Study location	South Korea.
Study setting	Outpatient follow-up.
Study dates	February 2007 to December 2008.
Sources of funding	Grants from the Korean Diabetes Association. The investigators received drugs from Handok Pharmaceuticals Co. Ltd., GlaxoSmithKline Ltd. Korea, and Merck Ltd. Korea.
Inclusion criteria	Ages 30-65; HbA1c 6.5-9.5%; never taken an oral hypoglycaemic agent.
Exclusion criteria	Glucocorticoid users; pregnant women; people who had clinically significant liver disease (AST, ALT >2.5 x upper limit of normal); significant renal disease (serum creatinine >1.5 mg/dL in men, >1.4 mg/dL in women); a history of lactic acidosis; a history of unstable angina or severe angina pectoris; a history or treatment for congestive heart failure; contraindications to metformin or sulfonylurea treatment.
Recruitment / selection of participants	No additional information.

Intervention(s)	Glimepiride N=118
	Glimepiride initially 2mg once a day, up to 4mg twice a day for 48 weeks.
	Concomitant therapy: All people received a lifestyle intervention including individualised education according to current, recommended guidelines for medical nutritional treatment. People were recommended to perform at least 150 minutes per week of moderate-intensity aerobic physical activity, provided exercise was not contraindicated.
Strata 1: People with type 2 diabetes mellitus and heart failure	Not stated/unclear
Strata 2: People with atherosclerotic cardiovascular diseases	Not stated/unclear
Strata 3: People with type 2 diabetes mellitus and chronic kidney disease	People without chronic kidney disease
Strata 4: People with type 2 diabetes mellitus and high cardiovascular risk	Not stated/unclear
Subgroup 1: People with frailty	Not stated/unclear
Subgroup 2: Onset of type 2 diabetes mellitus	Not stated/unclear
Subgroup 3: People with non-alcoholic fatty liver disease	People without non-alcoholic fatty liver disease

Subgroup 4: People with obesity	Not stated/unclear
Subgroup 5: eGFR category at baseline	Not stated/unclear
Subgroup 6: Albuminuria category at baseline	Not stated/unclear
Sensitivity analysis category: Enrichment trial status	5) All treatment naïve
Population subgroups	No additional information.
Comparator	Metformin N=114
	Metformin initially 500mg once a day, up to 1000mg twice a day for 48 weeks. Concomitant therapy: All people received a lifestyle intervention including individualised education according to current, recommended guidelines for medical nutritional treatment. People were recommended to perform at least 150 minutes per week of moderate-intensity aerobic physical activity, provided exercise was not contraindicated. A third arm (n=117) received rosiglitazone. This arm was not included in this data extraction as it was not relevant to the review protocol.
Number of participants	349 (including rosiglitazone, 232 if excluding rosiglitazone).
Duration of follow-up	48 weeks.
Indirectness	No additional information.
Method of analysis	ITT
Additional comments	No additional information.

117.2.1. Glimepiride (N = 118)

Glimepiride initially 2mg once a day, up to 4mg twice a day for 48 weeks. Concomitant therapy: All people received a lifestyle intervention including

individualised education according to current, recommended guidelines for medical nutritional treatment. People were recommended to perform at least 150 minutes per week of moderate-intensity aerobic physical activity, provided exercise was not contraindicated.

117.2.2. Metformin (N = 114)

Metformin initially 500mg once a day, up to 1000mg twice a day for 48 weeks. Concomitant therapy: All people received a lifestyle intervention including individualised education according to current, recommended guidelines for medical nutritional treatment. People were recommended to perform at least 150 minutes per week of moderate-intensity aerobic physical activity, provided exercise was not contraindicated.

117.3. Characteristics

117.3.1. Arm-level characteristics

Sample size Mean age (SD) (years) 50.8 (8.9) 51.8 (8.9)	6; % = 57.89 8.5)
Mean age (SD) (years) 50.8 (8.9)	8.5)
51.8 (8.5)
Marana (OD)	/
Mean (SD)	
Ethnicity $n = NR$; % = NR $n = NR$	R ; % = NR
Sample size	
	R ; % = NR
Sample size	
Presence of frailty NR (NR) NR (NR)	IR)
Mean (SD)	
Time since type 2 diabetes diagnosis NR (NR) NR (NR)	IR)
Mean (SD)	
HbA1c (%) 7.8 (0.8) 7.9 (0.8)	.8)
Mean (SD)	
Blood pressure (mmHg) NA (NA) NA (N	A)
Mean (SD)	

Characteristic	Glimepiride (N = 118)	Metformin (N = 114)
Systolic blood pressure	126.3 (12.8)	128.2 (12.4)
Mean (SD)		(,
Diastolic blood pressure	78.4 (8.7)	79.8 (8.6)
Mean (SD)		,
Heart rate	NR (NR)	NR (NR)
Mean (SD)		,
Smoking status	n = NR ; % = NR	n = NR ; % = NR
Sample size		11111, 70 1111
Alcohol consumption	n = NR ; % = NR	n = NR ; % = NR
Sample size		11 - 141X , 70 - 141X
Presence of severe mental illness	n = NR ; % = NR	n = NR ; % = NR
Sample size		11 - INIX , 70 - INIX
People with significant cognitive impairment	n = NR ; % = NR	n = NR ; % = NR
Sample size		
People with a learning disability	n = NR ; % = NR	n = NR ; % = NR
Sample size	07.0 (40.0)	
Weight (kg)	67.9 (10.9)	68.9 (11.1)
Mean (SD)		
BMI (kg/m2)	25.5 (3.1)	25.7 (3.2)
Mean (SD)		
Number of people with obesity	n = NR ; % = NR	n = NR ; % = NR
Sample size		
Cholesterol and lipid levels (mg/dL)	NA (NA)	NA (NA)
Mean (SD)		,
Total cholesterol	190.2 (41.3)	186.8 (34.3)
Mean (SD)		(5.1.5)
HDL cholesterol	45 (12)	43 (13)
Mean (SD)		(- /

Characteristic	Glimepiride (N = 118)	Metformin (N = 114)
LDL cholesterol	110.9 (42)	106 (33.7)
Mean (SD)		
Triglyceride	127 (114)	144 (99)
Mean (SD)		
Albumin creatinine ratio	NR (NR)	NR (NR)
Mean (SD)		
eGFR (mL/min/1.73m2)	NR (NR)	NR (NR)
Mean (SD)		
Other antidiabetic medication used	n = NR	n = NR ; % = NR
Sample size		
Blood pressure-lowering medication used	n = 34 ; % = 28.81	n = 33 ; % = 28.95
Sample size		
Statins/lipid-lowering medication used	n = 35 ; % = 29.66	n = 33 ; % = 28.95
Sample size		
Other treatment being received	n = NR ; % = NR	n = NR ; % = NR
Sample size		

118. Yuan, 2012

Bibliographic Reference

Yuan, G. H.; Song, W. L.; Huang, Y. Y.; Guo, X. H.; Gao, Y.; Efficacy and tolerability of exenatide monotherapy in obese patients with newly diagnosed type 2 diabetes: a randomized, 26 weeks metformin-controlled, parallel-group study; Chin Med J (Engl); 2012; vol. 125 (no. 15); 2677-81

Secondary publication of another included study- see primary study for details	No additional information.
Other publications associated with this study included in review	No additional information.
Trial name / registration number	No additional information.
Study type	Randomised controlled trial (RCT)
Study location	China.
Study setting	Outpatient follow-up.
Study dates	No additional information.
Sources of funding	No additional information.
Inclusion criteria	Greater than 18 years of age; new type 2 diabetes (<1 month); HbA1c 7-10%; BMI 28-40kg/m2 or waist circumference >90cm (male) or 85cm (female); managing type 2 diabetes with diet and exercise prior to the study.
Exclusion criteria	Ever been treated with antidiabetic or lipid lowering agents; had blood pressure >150/100mmHg; had a history or presence of clinically significant cardiac disease within the year prior to inclusion in the study; had renal or hepatic dysfunction; had a history or clinically suspected hyper- or hypothyroid disease; Cushing syndrome.
Recruitment / selection of participants	No additional information.
Intervention(s)	Exenatide N=33

	Exenatide 5 micrograms injected subcutaneously twice daily before morning and evening meals for 4 weeks, then increased to 10 micrograms twice daily thereafter. Given for a total of 26 weeks. Concomitant therapy: On confirmed hypoglycaemic event (documented blood glucose <3.3 mmol/L) and 2 unconfirmed events allowed the dose of
	either drug to be decreased by 50% (additional episodes allowed further decrease or discontinuation).
Strata 1: People with type 2 diabetes mellitus and heart failure	People without heart failure
Strata 2: People with atherosclerotic cardiovascular diseases	
Strata 3: People with type 2 diabetes mellitus and chronic kidney disease	Not stated/unclear
Strata 4: People with type 2 diabetes mellitus and high cardiovascular risk	People at higher risk of developing cardiovascular disease Based on BMI, hypertension and presence of type 2 diabetes.
Subgroup 1: People with frailty	Not stated/unclear
Subgroup 2: Onset of type 2 diabetes mellitus	Not stated/unclear
Subgroup 3: People with non-alcoholic fatty liver disease	Not stated/unclear
Subgroup 4: People with obesity	People with obesity

Not stated/unclear
Not stated/unclear
5) All treatment naïve
No additional information.
Metformin N=26 Metformin 500mg twice daily for 4 weeks, then increased to 500mg three times a day for 4-12 weeks. If fasting plasma glucose >6.1 mmol/L at week 12, metformin administrations was increased to 1000mg twice daily. Concomitant therapy: On confirmed hypoglycaemic event (documented blood glucose <3.3 mmol/L) and 2 unconfirmed events allowed the dose of either drug to be decreased by 50% (additional episodes allowed further decrease or discontinuation).
59
26 weeks.
No additional information.
ITT
No additional information.

118.2.1. Exenatide (N = 33)

Exenatide 5 micrograms injected subcutaneously twice daily before morning and evening meals for 4 weeks, then increased to 10 micrograms twice daily thereafter. Given for a total of 26 weeks. Concomitant therapy: On confirmed hypoglycaemic event (documented blood glucose <3.3 mmol/L) and 2 unconfirmed events allowed the dose of either drug to be decreased by 50% (additional episodes allowed further decrease or discontinuation).

118.2.2. Metformin (N = 26)

Metformin 500mg twice daily for 4 weeks, then increased to 500mg three times a day for 4-12 weeks. If fasting plasma glucose >6.1 mmol/L at week 12, metformin administrations was increased to 1000mg twice daily. Concomitant therapy: On confirmed hypoglycaemic event (documented blood glucose <3.3 mmol/L) and 2 unconfirmed events allowed the dose of either drug to be decreased by 50% (additional episodes allowed further decrease or discontinuation).

118.3. Characteristics

118.3.1. Arm-level characteristics

7111110101101101101101101101101101101101	Tiolo.ii Aim level onalaoteristics			
Characteristic	Exenatide (N = 33)	Metformin (N = 26)		
% Male	n = 17; % = 51	n = 12; % = 46		
Sample size				
Mean age (SD) (years)	58.5 (10.6)	56.8 (7.6)		
Mean (SD)				
Ethnicity	n = NR ; % = NR	n = NR ; % = NR		
Sample size				
Comorbidities	n = NR ; % = NR	n = NR ; % = NR		
Sample size				
Presence of frailty	n = NR ; % = NR	n = NR ; % = NR		
Sample size				
Time since type 2 diabetes diagnosis	NR (NR)	NR (NR)		
Mean (SD)				
HbA1c (%)	8.27 (1.58)	8.11 (1.92)		
Mean (SD)				
Blood pressure (mmHg)	NA (NA)	NA (NA)		
Mean (SD)				
Systolic blood pressure	138 (12)	130 (10)		
Mean (SD)				
Diastolic blood pressure	87 (11)	85 (10)		
Mean (SD)				
Heart rate	NR (NR)	NR (NR)		

	P	BB (6) - 1 (2) - 25
Characteristic Magn (SD)	Exenatide (N = 33)	Metformin (N = 26)
Mean (SD)	- ND - 0/ - ND	
Smoking status	n = NR ; % = NR	n = NR; % = NR
Sample size		
Alcohol consumption	n = NR ; % = NR	n = NR ; % = NR
Sample size		
Presence of severe mental illness	n = NR ; % = NR	n = NR ; % = NR
Sample size		
People with significant cognitive impairment	n = NR ; % = NR	n = NR ; % = NR
Sample size	n = ND : 0/ ND	
People with a learning disability	n = NR ; % = NR	n = NR ; % = NR
Sample size	00 0 (40 0)	
Weight (kg)	82.2 (12.8)	83.7 (10.7)
Mean (SD)		
BMI (kg/m2)	30.6 (2.8)	29.3 (2.6)
Mean (SD)		
Number of people with obesity	n = NR ; % = NR	n = NR ; % = NR
Sample size	ND (ND)	
Cholesterol and lipid levels	NR (NR)	NR (NR)
Mean (SD)		
Albumin creatinine ratio	NR (NR)	NR (NR)
Mean (SD)		
eGFR (mL/min/1.73m2)	NR (NR)	NR (NR)
Mean (SD)		
Other antidiabetic medication used	n = NR ; % = NR	n = NR ; % = NR
Sample size		11 - 141C, 70 - 141C
Blood pressure-lowering medication used	n = NR ; % = NR	n = NR ; % = NR
Sample size		,
Statins/lipid-lowering medication used	n = NR ; % = NR	n = NR ; % = NR
Sample size		

Characteristic	Exenatide (N = 33)	Metformin (N = 26)
Other treatment being received	n = NR ; % = NR	n = NR ; % = NR
Sample size		

119. Zhang, 2020

Bibliographic Reference

Zhang, L. Y.; Qu, X. N.; Sun, Z. Y.; Zhang, Y.; Effect of liraglutide therapy on serum fetuin A in patients with type 2 diabetes and non-alcoholic fatty liver disease; Clinics & Research in Hepatology & Gastroenterology; 2020; vol. 44 (no. 5); 674-680

	tady dotains	
Secondary publication of another included study- see primary study for details	NA	
Other publications associated with this study included in review	None	
Trial name / registration number	Not reported	
Study type	Randomised controlled trial (RCT)	
Study location	Shandong, China	
Study setting	Community/Hospital	
Study dates	Not reported	
Sources of funding	Funded by Yantai Affiliated Hospital of Binzhou Medical University	
Inclusion criteria	 Aged ≥18 to ≤70 years Treatment naive to hypoglycaemic drugs≥3 months before trial inclusion HbA1c level ≥7.0 to ≤14% BMI ≥20 to ≤35 kg/m2 Weight fluctuation<10% ≤3 months before trial inclusion Non-alcoholic fatty liver disease (as defined by Chinese Association for the Study of Liver Disease [2010]: i. no history of alcohol-drinking habits or alcohol intake <140g/week for men or <70g/week for women, in past 12 months; ii. liver imaging results meet diagnostic criteria of diffuse fatty liver and could not be explained by other reasons, and/or participants with metabolic syndrome-related components showing persistent elevation of ALT or AST, and gamma-GT of unknown cause >6 months; iii. specific diseases that could lead to steatosis have been excluded (e.g. viral 	

	hepatitis, drug-induced liver disease, total parenteral nutrition, Wilson's disease, and autoimmune liver disease.)
Exclusion criteria	 Diagnosis of type 1 diabetes or secondary diabetes Presence of hypertension, kidney, cardiovascular, hyperthyroidism, hypothyroidism, acute stress state or any active inflammatory diseases Receiving insulin
Recruitment / selection of participants	Participants recruited from Yantai Affiliated Hospital of Binzhou Medical University, China. All participants had previous metformin therapy (500 mg thrice daily) three times per day but maintained HbA1c>7%. Randomisation 1:1 to groups using computer-generated sequence. All participants given information about diet and exercise.
Intervention(s)	Liraglutide 1.2 mg once daily
	Participants started on subcutaneous injections of liraglutide 0.6 mg once daily for 1 week, increased to 1.2 mg once daily for 23 weeks.
Strata 1: People with type 2 diabetes mellitus and heart failure	Not stated/unclear
Strata 2: People with atherosclerotic cardiovascular diseases	Not stated/unclear
Strata 3: People with type 2 diabetes mellitus and chronic kidney disease	Not stated/unclear
Strata 4: People with type 2 diabetes mellitus and high cardiovascular risk	Not stated/unclear
Subgroup 1: People with frailty	Not stated/unclear
Subgroup 2: Onset of type	Not stated/unclear

2 diabetes mellitus	
Subgroup 3: People with non-alcoholic fatty liver disease	People with non-alcoholic fatty liver disease
Subgroup 4: People with obesity	Not stated/unclear
Subgroup 5: eGFR category at baseline	Not stated/unclear
Subgroup 6: Albuminuria category at baseline	Not stated/unclear
Sensitivity analysis category: Enrichment trial status	6) No response criteria All participants had previously taken metformin but maintained HbA1c % level >7% on it. Washout period of ≥3-mo inclusion criteria.
Population subgroups	
Comparator	 Pioglitazone 30 mg once daily Oral pioglitazone tablets 15 mg once daily for 1 week, increased to 30 mg once daily for remaining 23 weeks.
Number of participants	N=60
Duration of follow-up	24 weeks
Method of analysis	Not stated/unclear

119.2.1. Liraglutide 1.2 mg once daily (N = 30)

Subcutaneous injection of liraglutide 0.6 mg once daily for 1 week and then 1.2 mg once daily for remaining 23 weeks.

119.2.2. **Pioglitazone 30 mg once daily (N = 30)**

Oral pioglitazone tablets 15 mg once daily for 1 week and then 30 mg once daily for remaining 23 weeks.

119.3. Characteristics

119.3.1. Arm-level characteristics

119.5.1. Allii-level C	, ilai acteristics	
Characteristic	Liraglutide 1.2 mg once daily (N = 30)	Pioglitazone 30 mg once daily (N = 30)
% Male	n = 13; % = 43.3	n = 15; % = 50
Sample size		
Mean age (SD) (years)	50.2 (11.5)	51.5 (12.1)
Mean (SD)		
Ethnicity	NR	NR
Nominal		
Comorbidities	NR	NR
Nominal		
Presence of frailty	NR	NR
Nominal		
Time since type 2 diabetes diagnosis	NR	NR
Nominal		
HbA1c (%)	8.1 (2)	8.1 (1.7)
Mean (SD)		
Blood pressure (mmHg)	NA (NA)	NA (NA)
Mean (SD)		
Systolic blood pressure	133.3 (17.1)	135.9 (12.6)
Mean (SD)		
Diastolic blood pressure	81.5 (13.4)	81.2 (12.1)
Mean (SD)		
Heart rate Nominal	NR	NR
	ND	
Smoking status	NR	NR
Nominal		
Alcohol consumption	NR	NR
Nominal		

Liraglutide 1.2 mg once daily (N = 30)	Pioglitazone 30 mg once daily (N = 30)
NR	NR
NR	NR
NR	NR
70.0 (0.0)	
19.3 (8.8)	78 (9.2)
27.6 (5.2)	27.1 (3.8)
NR	NR
NA (NA)	NA (NA)
5.2 (1.1)	5.1 (1.5)
1.1 (0.2)	1.1 (0.4)
3.3 (1)	
, ,	3.3 (1.3)
0.0 (0.5)	
0.9 (0.5)	0.9 (0.6)
NR	NR
NR (NR)	NR (NR)
	1417 (1417)
	daily (N = 30) NR NR NR 79.3 (8.8) 27.6 (5.2) NR NA (NA) 5.2 (1.1) 1.1 (0.2) 3.3 (1) 0.9 (0.5) NR

Characteristic	Liraglutide 1.2 mg once daily (N = 30)	Pioglitazone 30 mg once daily (N = 30)
Other antidiabetic medication used	NR	NR
Nominal		
Blood pressure-lowering medication used	NR	NR
Nominal		
Statins/lipid-lowering medication used	NR	NR
Nominal		
Other treatment being received	NR	NR
Nominal		

120. Zhou, 2022

Bibliographic Reference

Zhou, H.; Ding, J.; Mohammad, O. H.; Wu, L.; Yang, S.; Effects of Metformin Combined with Dapagliflozin on Homocysteine, Cystatin C and Beta-2 Microglobulin Levels in Patients with Diabetes Mellitus; Indian journal of pharmaceutical sciences; 2022; vol. 84; 153-157

	tudy details		
Secondary publication of another included study- see primary study for details	No additional information.		
Other publications associated with this study included in review	No additional information.		
Trial name / registration number	No additional information.		
Study type	Randomised controlled trial (RCT)		
Study location	China.		
Study setting	Outpatient follow-up.		
Study dates	No additional information.		
Sources of funding	No additional information.		
Inclusion criteria	Meeting the diagnostic criteria of T2DM; course of disease at least 12 months; people and their families know about the situations and give their informed consent.		
Exclusion criteria	Combined with acute and chronic infectious diseases of various tissues and organs; combined with malignant tumours; combined with immune system diseases; have allergic reactions to the drugs adopted in our study.		
Recruitment / selection of participants	No additional information.		
Intervention(s)	Dapagliflozin + Metformin N=75 Dapagliflozin 10mg once a day and metformin 250mg twice daily for 24 weeks. Concomitant therapy: No additional information.		

Strata 1: People with type 2 diabetes mellitus and heart failure	Not stated/unclear
Strata 2: People with atherosclerotic cardiovascular diseases	Not stated/unclear
Strata 3: People with type 2 diabetes mellitus and chronic kidney disease	Not stated/unclear
Strata 4: People with type 2 diabetes mellitus and high cardiovascular risk	Not stated/unclear
Subgroup 1: People with frailty	Not stated/unclear
Subgroup 2: Onset of type 2 diabetes mellitus	Not stated/unclear
Subgroup 3: People with non-alcoholic fatty liver disease	Not stated/unclear
Subgroup 4: People with obesity	Not stated/unclear
Subgroup 5: eGFR category at baseline	Not stated/unclear
Subgroup 6: Albuminuria category at baseline	Not stated/unclear
Sensitivity analysis	8) Not reported

category: Enrichment trial status	
Population subgroups	No additional information.
Comparator	Metformin N=75 Metformin 250mg twice daily for 24 weeks. Concomitant therapy: No additional information.
Number of participants	150
Duration of follow-up	24 weeks.
Indirectness	No additional information.
Method of analysis	Not stated/unclear
Additional comments	No additional information.

120.2.1. Dapagliflozin + Metformin (N = 75)

Dapagliflozin 10mg once a day and metformin 250mg twice daily for 24 weeks. Concomitant therapy: No additional information.

120.2.2. Metformin (N = 75)

Metformin 250mg twice daily for 24 weeks. Concomitant therapy: No additional information.

120.3. Characteristics

120.3.1. Arm-level characteristics

Characteristic	Dapagliflozin + Metformin (N = 75)	Metformin (N = 75)
% Male Sample size	n = 39; % = 52	n = 40 ; % = 53
Mean age (SD) (years) Mean (SD)	47.69 (5.32)	47.72 (5.26)

Characteristic	Dapagliflozin + Metformin (N = 75)	Metformin (N = 75)
Ethnicity	n = NR ; % = NR	n = NR ; % = NR
Sample size		
Comorbidities	n = NR ; % = NR	n = NR ; % = NR
Sample size		
Presence of frailty Sample size	n = NR ; % = NR	n = NR ; % = NR
Time since type 2 diabetes diagnosis	NR (NR)	
Mean (SD)	TVIX (TVIX)	NR (NR)
HbA1c	NR (NR)	
Mean (SD)		NR (NR)
Blood pressure	NR (NR)	NR (NR)
Mean (SD)		
Heart rate	NR (NR)	NR (NR)
Mean (SD)		
Smoking status	n = NR ; % = NR	n = NR ; % = NR
Sample size		
Alcohol consumption	n = NR ; % = NR	n = NR ; % = NR
Sample size		
Presence of severe mental illness Sample size	n = NR ; % = NR	n = NR ; % = NR
People with significant cognitive	n = NR ; % = NR	
impairment	11 - 1417, 70 - 1417	n = NR ; % = NR
Sample size		
People with a learning disability	n = NR ; % = NR	n = NR ; % = NR
Sample size		
Weight	NR (NR)	NR (NR)
Mean (SD)		
BMI (kg/m2)	27.69 (1.45)	27.61 (1.49)
Mean (SD)		

Characteristic	Dapagliflozin + Metformin (N = 75)	Metformin (N = 75)
Number of people with obesity	n = NR ; % = NR	n = NR ; % = NR
Sample size		
Cholesterol and lipid levels	NA (NA)	NA (NA)
Mean (SD)		
Total cholesterol	6.17 (0.33)	6.15 (0.34)
Mean (SD)		0.13 (0.54)
HDL cholesterol	1.15 (0.85)	4.40 (0.00)
Mean (SD)		1.16 (0.83)
LDL cholesterol	1.16 (0.35)	
Moon (SD)		1.14 (0.36)
Mean (SD) Triglycerides	3.03 (0.28)	
	0.00 (0.20)	3.05 (0.21)
Mean (SD)		
Albumin creatinine ratio	NR (NR)	NR (NR)
Mean (SD)		
eGFR (mL/min/1.73m2)	NR (NR)	NR (NR)
Mean (SD)		
Other antidiabetic medication used	n = NR ; % = NR	n = NR ; % = NR
Sample size		11 - 1417, 70 - 1417
Blood pressure-lowering medication	n = NR ; % = NR	NID 0/ NID
used		n = NR ; % = NR
Sample size		
Statins/lipid-lowering medication used	n = NR ; % = NR	n = NR ; % = NR
Sample size		
Other treatment being received	n = NR ; % = NR	n = NR ; % = NR
Sample size		

121. Zografou, 2015

Bibliographic Reference

Zografou, Ioanna; Sampanis, Christos; Gkaliagkousi, Eugenia; Iliadis, Fotios; Papageorgiou, Athanasios; Doukelis, Panagiotis; Vogiatzis, Konstantinos; Douma, Stella; Effect of vildagliptin on hsCRP and arterial stiffness in patients with type 2 diabetes mellitus.; Hormones (Athens, Greece); 2015; vol. 14 (no. 1); 118-25

Secondary publication of another included study- see primary study for details	No additional information.
Other publications associated with this study included in review	No additional information.
Trial name / registration number	No additional information.
Study type	Randomised controlled trial (RCT)
Study location	Greece.
Study setting	Outpatient follow-up.
Study dates	No additional information.
Sources of funding	None declared.
Inclusion criteria	People age at least 18 to no more than 70 years old; inadequate glycaemic control (HbA1c 7-9%) despite diet and exercise for 3 months.
Exclusion criteria	People with any macrovascular or microvascular diabetic complications; history of heart disease; uncontrolled hypertension (systolic blood pressure more than 180 mmHg and/or diastolic blood pressure more than 110 mmHg); excessive dyslipidaemic (total cholesterol more than 110 mmHg, triglycerides more than 400 mg/dL); elevated liver enzymes three times above the upper normal range and any other comorbidity that could interfere with the study; pregnant women and those who were breastfeeding or planning for pregnancy; women of childbearing age were advised to use contraception if they were sexually active.
Recruitment / selection of participants	No additional information.

Intervention(s)	Vildagliptin + Metformin N=32
	Vildagliptin 50mg twice daily plus metformin 850mg twice daily.
Cointervention	Concomitant therapy: Antihypertensive and lipid-lowering drugs were unchanged during the study period. People were instructed to maintain dietary habits and daily activities.
Strata 1: People with type 2 diabetes mellitus and heart failure	People without heart failure
Strata 2: People with atherosclerotic cardiovascular diseases	People without other cardiovascular diseases
Strata 3: People with type 2 diabetes mellitus and chronic kidney disease	People without chronic kidney disease
Strata 4: People with type 2 diabetes mellitus and high cardiovascular risk	Not stated/unclear
Subgroup 1: People with frailty	Not stated/unclear
Subgroup 2: Onset of type 2 diabetes mellitus	Not stated/unclear
Subgroup 3: People with non-alcoholic fatty liver disease	Not stated/unclear
Subgroup 4: People with obesity	Not stated/unclear
Subgroup 5: eGFR category at baseline	Not stated/unclear

Subgroup 6: Albuminuria category at baseline	Not stated/unclear
Sensitivity analysis category: Enrichment trial status	5) All treatment naïve
Population subgroups	No additional information.
Comparator	Metformin N=32 Metformin 850mg twice daily.
Number of participants	64
Duration of follow-up	6 months.
Indirectness	No additional information.
Method of analysis	Not stated/unclear
Additional comments	No additional information.

121.2.1. Vildagliptin + Metformin (N = 32)

Vildagliptin 50mg twice daily plus metformin 850mg twice daily. Concomitant therapy: Antihypertensive and lipid-lowering drugs were unchanged during the study period. People were instructed to maintain dietary habits and daily activities.

121.2.2. Metformin (N = 32)

Metformin 850mg twice daily. Concomitant therapy: Antihypertensive and lipid-lowering drugs were unchanged during the study period. People were instructed to maintain dietary habits and daily activities.

121.3. Characteristics

121.3.1. Arm-level characteristics

121.3.1. Allii-level cilalac		
Characteristic	Vildagliptin + Metformin (N = 32)	Metformin (N = 32)
% Male	n = 18; % = 56	n = 20 ; % = 63
Sample size		
Mean age (SD) (years)	52 (11.2)	56 (10.5)
Mean (SD)		
Ethnicity	n = NR ; % = NR	n = NR ; % = NR
Sample size		
Comorbidities	n = NR ; % = NR	n = NR ; % = NR
Sample size		
Presence of frailty	n = NR ; % = NR	n = NR ; % = NR
Sample size		
Time since type 2 diabetes diagnosis	NR (NR)	NR (NR)
Mean (SD)		
Smoking status Sample size	n = NR ; % = 12.5	n = NR ; % = 25
·	n = ND : 0/ = ND	
Alcohol consumption Sample size	n = NR ; % = NR	n = NR ; % = NR
Presence of severe mental illness	n = NR ; % = NR	
Sample size	11 - INK , 70 - INK	n = NR ; % = NR
People with significant cognitive	n = NR ; % = NR	
impairment	II - IVIX , 70 - IVIX	n = NR ; % = NR
Sample size		
People with a learning disability	n = NR ; % = NR	n = NR ; % = NR
Sample size		
Number of people with obesity	n = NR ; % = NR	n = NR ; % = NR
Sample size		
Other antidiabetic medication used	n = NR ; % = NR	n = NR ; % = NR
Sample size		

Characteristic	Vildagliptin + Metformin (N = 32)	Metformin (N = 32)
Blood pressure-lowering medication used	n = NR; % = 40.6	n = NR ; % = 43.8
Sample size		
Statins/lipid-lowering medication used	n = NR; % = 40.6	n = NR ; % = 53.1
Sample size		
Other treatment being received	n = NR ; % = NR	n = NR ; % = NR
Sample size		