

Clinical trial results:

A Randomized, Double-blind, Two Treatment, Two Period, Chronic Dosing (4 weeks), Cross-Over, Multi-Center Pilot Study to Evaluate the Effects of Budesonide/Glycopyrronium/Formoterol Fumarate and Glycopyrronium/Formoterol Fumarate on Specific Image-based Airway Volumes and Resistance in Subjects With Moderate to Severe Chronic Obstructive Pulmonary Disease Summary

EudraCT number	2018-001704-10		
Trial protocol	NL		
Global end of trial date	11 November 2019		
Results information			
Result version number	v2 (current)		
This version publication date	10 February 2021		
First version publication date	27 November 2020		
Version creation reason			

Trial information

Trial identification			
Sponsor protocol code	D5980C00019		
Additional study identifiers			
ISRCTN number	-		
ClinicalTrials.gov id (NCT number)	-		
WHO universal trial number (UTN)	-		

Notes:

Sponsors			
Sponsor organisation name	AstraZeneca AB		
Sponsor organisation address	Pepparedsleden 1, Molndal, Sweden,		
Public contact	Magnus Aurivillius, AstraZeneca AB, magnus.aurivillius@astrazeneca.com		
Scientific contact	Magnus Aurivillius, AstraZeneca AB, magnus.aurivillius@astrazeneca.com		

Notes:

Paediatric regulatory details		
Is trial part of an agreed paediatric investigation plan (PIP)	No	
Does article 45 of REGULATION (EC) No 1901/2006 apply to this trial?	No	
Does article 46 of REGULATION (EC) No 1901/2006 apply to this trial?	No	
Natar		

Notes:

Results analysis stage		
Analysis stage	Final	
Date of interim/final analysis	16 December 2019	
Is this the analysis of the primary completion data?	Yes	
Primary completion date	11 November 2019	
Global end of trial reached?	Yes	
Global end of trial date	11 November 2019	
Was the trial ended prematurely?	No	

Notes:

General information about the trial

Main objective of the trial:

To assess the effects of BGF MDI and GFF MDI on specific image-based airway volumes and resistance in subjects with moderate to severe COPD following chronic twice daily (BID) dosing after approximately 4 weeks of treatment.

Protection of trial subjects:

Ventolin HFA was provided throughout the study for subjects to take as needed for relief of symptoms.

Background therapy: -

Evidence for comparator: -

Actual start date of recruitment	26 February 2019	
Long term follow-up planned	No	
Independent data monitoring committee (IDMC) involvement?	No	

Notes:

Population of trial subjects

Subjects enrolled per country

Country: Number of subjects enrolled	Belgium: 7
Country: Number of subjects enrolled	Netherlands: 16
Worldwide total number of subjects	23
EEA total number of subjects	23

Notes:

In utero	0
Preterm newborn - gestational age < 37 wk	0
Newborns (0-27 days)	0
Infants and toddlers (28 days-23 months)	0
Children (2-11 years)	0
Adolescents (12-17 years)	0
Adults (18-64 years)	9
From 65 to 84 years	14
85 years and over	0

Subject disposition

Recruitment

Recruitment details:

This study randomized 23 subjects at 4 study centers (1 center in Belgium and 3 centers in the Netherlands) from 26 February 2019 to 11 November 2019.

Pre-assignment

Screening details:

Subjects were randomized into 1 of 2 treatment sequences. Sequence 1 received BGF MDI in Period 1 followed by GFF MDI in Period 2. Sequence 2 received GFF MDI in Period 1 followed by BGF MDI in Period 2

Period 1			
Period 1 title	Overall Study (overall period)		
Is this the baseline period?	Yes		
Allocation method	Randomised - controlled		
Blinding used	Double blind		
Roles blinded	Subject, Investigator, Monitor, Data analyst, Carer, Assessor		
Blinding implementation details:			
All Subjects Randomized			
Arms			
Arm title	Overall Study		
Arm description: -			
Arm type	Experimental		
Investigational medicinal product name	Budesonide/Glycopyrronium/Formoterol Fumarate Metered Dose Inhalation		
Investigational medicinal product code			
Other name			
Pharmaceutical forms	Pressurised inhalation, suspension		
Routes of administration	Inhalation use		
Dosage and administration details:			
BGF MDI 320/14.4/9.6 μg			
Investigational medicinal product name	Glycopyrronium/Formoterol Fumarate Metered Dose Inhalation		
Investigational medicinal product code			
Other name			
Pharmaceutical forms	Pressurised inhalation, suspension		
Routes of administration	Inhalation use		
Docago and administration dotails:			

Dosage and administration details:

GFF MDI 14.4/9.6 μg

Number of subjects in period 1	Overall Study	
Started	23	
Completed	21	
Not completed	2	
Protocol deviation	1	
Adverse event, non-fatal	1	

Baseline characteristics

Reporting groups Reporting group title Overall Study Reporting group description: All Subjects Randomized

Reporting group values	Overall Study	Total	
Number of subjects	23	23	
Age Categorical			
Units: Subjects			
In utero	0	0	
Preterm newborn infants (gestational age <37 wks)	0	0	
Newborns (0-27 days)	0	0	
Children (2-11 years)	0	0	
Adolescents (12-17 years)	0	0	
Adults (≥40-<65 years)	9	9	
Adults (≥65-<80 years)	14	14	
Age Continuous			
Units: years			
arithmetic mean	64.9		
standard deviation	± 7.6	-	
Gender Categorical			
Units: Subjects			
Female	5	5	
Male	18	18	
Race			
Units: Subjects			
American Indian or Alaska Native	0	0	
Asian	0	0	
Native Hawaiian or Other Pacific Islander	0	0	
Black or African American	0	0	
White	23	23	
More than one race	0	0	
Unknown or Not Reported	0	0	

Subject analysis sets		
Subject analysis set title	BGF MDI	
Subject analysis set type	Intention-to-treat	
Subject analysis set description:		
All Subjects Randomized		
Subject analysis set title	GFF MDI	_
Subject analysis set type	Intention-to-treat	
Subject analysis set description:		
All Subjects Randomized		

Reporting group values	BGF MDI	GFF MDI	
Number of subjects	22	23	
Age Categorical			
Units: Subjects			
In utero	0	0	
Preterm newborn infants (gestational age <37 wks)	0	0	
Newborns (0-27 days)	0	0	
Children (2-11 years)	0	0	
Adolescents (12-17 years)	0	0	
Adults (≥40-<65 years)	9	9	
Adults (≥65-<80 years)	13	14	
Age Continuous			
Units: years			
arithmetic mean	64.8	64.9	
standard deviation	± 7.8	± 7.6	
Gender Categorical			
Units: Subjects			
Female	5	5	
Male	17	18	
Race			
Units: Subjects			
American Indian or Alaska Native	0	0	
Asian	0	0	
Native Hawaiian or Other Pacific Islander	0	0	
Black or African American	0	0	
White	22	23	
More than one race	0	0	
Unknown or Not Reported	0	0	

End points

End points reporting groups		
Reporting group title	Overall Study	
Reporting group description: -		
Subject analysis set title	BGF MDI	
Subject analysis set type	Intention-to-treat	
Subject analysis set description:		
All Subjects Randomized		
Subject analysis set title	GFF MDI	
Subject analysis set type	Intention-to-treat	
Subject analysis set description:		
All Subjects Randomized		

Primary: Specific Image-based Airway Volume (siVaw)			
End point title Specific Image-based Airway Volume (siVaw)[1]			
End point description:			
Specific Image-based Airway Volume (silvolume. Reported as ratio to baseline.	Vaw) measured in mL/L. Average across lobes, adjusted for lobe		
End point type Primary			
End point timeframe:			
Baseline, Day 29			

Notes:

[1] - No statistical analyses have been specified for this primary end point. It is expected there is at least one statistical analysis for each primary end point.

Justification: According to the protocol, the primary analysis in this study was not a comparison between treatments, it was a comparison to baseline within each treatment. Therefore, no statistical analysis is included in this form as to only name a single treatment arm for each analysis generated validation errors. The estimates and confidence intervals for each comparison to baseline are however provided.

End point values	BGF MDI	GFF MDI	
Subject group type	Subject analysis set	Subject analysis set	
Number of subjects analysed	20	21	
Units: Ratio to baseline			
geometric mean (confidence interval 95%)	1.72 (1.38 to 2.13)	1.53 (1.28 to 1.83)	

Statistical analyses

No statistical analyses for this end point

Primary: Specific Image-based Airway Resistance (siRaw)			
End point title Specific Image-based Airway Resistance (siRaw)[2]			
End point description:			
Specific Image-based Airway lobe volume. Reported as rati	Resistance (siRaw) measured in kPa·s. Average across lobes, adjusted for to baseline.		
End point type Primary			
End point timeframe:	·		
Baseline, Day 29			

Notes:

[2] - No statistical analyses have been specified for this primary end point. It is expected there is at least one statistical analysis for each primary end point.

Justification: According to the protocol, the primary analysis in this study was not a comparison between treatments, it was a comparison to baseline within each treatment. Therefore, no statistical analysis is included in this form as to only name a single treatment arm for each analysis generated validation errors. The estimates and confidence intervals for each comparison to baseline are however provided.

End point values	BGF MDI	GFF MDI	
Subject group type	Subject analysis set	Subject analysis set	
Number of subjects analysed	20	21	
Units: Ratio to baseline			
geometric mean (confidence interval 95%)	0.50 (0.39 to 0.63)	0.52 (0.40 to 0.67)	

Statistical analyses

No statistical analyses for this end point

Secondary: Image-based Airway Volume (iVaw)		
End point title Image-based Airway Volume (iVaw)		
End point description:		
Image-based Airway Volume (iVaw) meavolume. Reported as ratio to baseline.	asured in mL. Average across lobes, without adjustment for lobe	
End point type Secondary		
End point timeframe:		
Baseline, Day 29		

End point values	BGF MDI	GFF MDI	
Subject group type	Subject analysis set	Subject analysis set	
Number of subjects analysed	20	21	
Units: Ratio to baseline			
geometric mean (confidence interval 95%)	1.70 (1.37 to 2.11)	1.51 (1.26 to 1.80)	

Statistical analyses

No statistical analyses for this end point

Secondary: Image-based Airway Resistance (iRaw)			
End point title Image-based Airway Resistance (iRaw)			
End point description:			
Image-based Airway Resistance (iRaw) measured in kPa·s/L. Average across lobes, without adjustment			

for lobe volume. Reported as ratio to baseline.

End point type

Secondary

End point type Secondary

EU-CTR publication date: 10 February 2021

End point timeframe:

End point values	BGF MDI	GFF MDI	
Subject group type	Subject analysis set	Subject analysis set	
Number of subjects analysed	20	21	
Units: Ratio to baseline			
geometric mean (confidence interval 95%)	0.50 (0.40 to 0.63)	0.52 (0.40 to 0.68)	

Statistical analyses

No statistical analyses for this end point

Secondary: Forced Expiratory Volume in 1 Second (Post-dose FEV1)			
End point title Forced Expiratory Volume in 1 Second (Post-dose FEV1)			
End point description:			
Change from baseline in Forced Expiratory Volume in One Second (Post-dose FEV1)			
End point type Secondary			
End point timeframe:			
Baseline, Day 29			

End point values	BGF MDI	GFF MDI	
Subject group type	Subject analysis set	Subject analysis set	
Number of subjects analysed	21	21	
Units: Liters			
arithmetic mean (confidence interval 95%)	0.346 (0.182 to 0.509)	0.273 (0.140 to 0.405)	

Statistical analyses

No statistical analyses for this end point

Secondary: Image-based Functional Residual Capacity (FRC)				
End point title Image-based Functional Residual Capacity (FRC)				
End point description:				
Change from baseline in Functional Residual Capacity (FRC).				
End point type Secondary				
End point timeframe:				
Baseline, Day 29				

End point values	BGF MDI	GFF MDI	
Subject group type	Subject analysis set	Subject analysis set	
Number of subjects analysed	22	21	
Units: Liters			
arithmetic mean (confidence interval 95%)	-0.28 (-0.77 to 0.21)	-0.50 (-0.81 to -0.18)	

EU-CTR publication date: 10 February 2021

Statistical analyses

No statistical analyses for this end point

Adverse events

Adverse events information

Timeframe for reporting adverse events:

Adverse events were collected from Visit 2 throughout the treatment period and including the follow-up period telephone call.

Adverse event reporting additional description:

The Safety Population was defined as all subjects who were randomized to treatment and received at least one dose of study treatment and for whom any post-dose data were available. Serious adverse events were collected from Visit 2 throughout the treatment period and including the follow-up period telephone call.

Assessment type	Systematic	
Dictionary used		
Dictionary name	MedDRA	
Dictionary version	22.1	
Reporting groups		
Reporting group title	BGF MDI	
Reporting group description:		
Budesonide/Glycopyrronium/Formoterol Fumarate Metered Dose Inhalation		
Reporting group title	GFF MDI	
Reporting group description:		
Glycopyrronium/Formoterol Fumarate Metered Dose Inhalation		

Serious adverse events	BGF MDI	GFF MDI	
Total subjects affected by serious adverse events			
subjects affected / exposed	1 / 22 (4.55%)	0 / 23 (0.00%)	
number of deaths (all causes)	0	0	
number of deaths resulting from adverse events	0	0	
Neoplasms benign, malignant and unspecified (incl cysts and polyps)			
Colon cancer			
subjects affected / exposed	1 / 22 (4.55%)	0 / 23 (0.00%)	
occurrences causally related to treatment / all	0 / 1	0 / 0	
deaths causally related to treatment / all	0 / 0	0 / 0	

Frequency threshold for reporting non-serious adverse events: 5 %

Non-serious adverse events	BGF MDI	GFF MDI	
Total subjects affected by non-serious adverse events			
subjects affected / exposed	0 / 22 (0.00%)	2 / 23 (8.70%)	
Respiratory, thoracic and mediastinal disorders			

Dyspnoea subjects affected / exposed	0 / 22 (0.00%)	2 / 23 (8.70%)	
occurrences (all)	0	2	

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More information

Substantial protocol amendments (globally)

Were there any global substantial amendments to the protocol? No

Interruptions (globally)

Were there any global interruptions to the trial? No

Limitations and caveats

None reported