

Clinical trial results:

A phase 2b, multicenter, randomized, double-blind, placebo-controlled, parallel-group, dose-selection study of S-600918 in patients with refractory chronic cough

Summary

EudraCT number	2019-002283-27	
Trial protocol	GB CZ PL	
Global end of trial date	28 December 2020	
Results information		
Result version number	v1 (current)	
This version publication date	11 January 2022	
First version publication date	11 January 2022	

Trial information

Trial identification		
Sponsor protocol code	1812VA323	
Additional study identifiers		
ISRCTN number	-	
ClinicalTrials.gov id (NCT number)	NCT04110054	
WHO universal trial number (UTN)	-	

Notes:

Sponsors	
Sponsor organisation name	Shionogi B.V.
Sponsor organisation address	Kingsfordweg 151, Amsterdam, Netherlands, 1043 GR
Public contact	Regulatory Affairs, Shionogi B.V., +44 2030534200, shionogiclintrials-admin@shionogi.co.jp
Scientific contact	Regulatory Affairs, Shionogi B.V., +44 2030534200, shionogiclintrials-admin@shionogi.co.jp

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

Notes:

Results analysis stage	
Analysis stage	Final
Date of interim/final analysis	28 December 2020
Is this the analysis of the primary completion data?	Yes
Primary completion date	28 December 2020
Global end of trial reached?	Yes
Global end of trial date	28 December 2020
Was the trial ended prematurely?	No

Notes:

General information about the trial

Main objective of the trial:

To determine the optimal dose of S-600918 in patients with refractory chronic cough by evaluating the change from baseline in 24-hour cough frequency (coughs per hour) with S-600918 compared with placebo

Protection of trial subjects:

The study was conducted in accordance with the protocol approved by the IRB/IEC, all applicable regulatory requirements (including patient privacy requirements), current ICH GCP, and the ethical principles outlined in the Declaration of Helsinki.

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Background therapy: -		
Evidence for comparator: -		
Actual start date of recruitment	26 January 2020	
Long term follow-up planned	No	
Independent data monitoring committee (IDMC) involvement?	No	
Notes:		

Population of trial subjects

Population of trial subjects	
Subjects enrolled per country	
Country: Number of subjects enrolled	Poland: 49
Country: Number of subjects enrolled	United Kingdom: 38
Country: Number of subjects enrolled	Czechia: 29
Country: Number of subjects enrolled	United States: 150
Country: Number of subjects enrolled	Japan: 91
Country: Number of subjects enrolled	Ukraine: 49
Worldwide total number of subjects	406
FFA total number of subjects	78

Notes:

Subjects enrolled per age group	
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)	285
From 65 to 84 years	121
85 years and over	0

Subject disposition

Recruitment Recruitment details: -**Pre-assignment** Screening details: Subjects were screened prior to enrollment Period 1 Period 1 title Treatment (overall period) Yes Is this the baseline period? Allocation method Randomised - controlled Blinding used Double blind Roles blinded Subject, Investigator, Monitor, Data analyst **Arms** Are arms mutually exclusive? Yes Arm title Active 50mg Arm description: -Arm type Experimental S-600918 Investigational medicinal product name Investigational medicinal product code Other name Pharmaceutical forms Tablet Routes of administration Oral use Dosage and administration details: 50mg, 150mg or 300mg, once daily for 4 weeks **Arm title** Active 150mg Arm description: -Arm type Experimental Investigational medicinal product name S-600918 Investigational medicinal product code Other name Pharmaceutical forms **Tablet** Routes of administration Oral use Dosage and administration details: 50mg, 150mg or 300mg, once daily for 4 weeks Arm title Active 300mg Arm description: -Experimental Arm type Investigational medicinal product name S-600918 Investigational medicinal product code Other name Pharmaceutical forms **Tablet** Routes of administration Oral use Dosage and administration details: 50mg, 150mg or 300mg, once daily for 4 weeks **Arm title** Placebo Arm description: -

Placebo

EU-CTR publication date: 11 January 2022

Arm type

Investigational medicinal product name	placebo
Investigational medicinal product code	
Other name	
Pharmaceutical forms	Tablet
Routes of administration	Oral use

Dosage and administration details:

once daily for 4 weeks

Number of subjects in period 1	Active 50mg	Active 150mg	Active 300mg
Started	101	103	100
Completed	100	99	92
Not completed	1	4	8
Consent withdrawn by subject	-	3	2
Adverse event, non-fatal	1	1	4
Covid-19	-	-	2

Number of subjects in period 1	Placebo	
Started	102	
Completed	99	
Not completed	3	
Consent withdrawn by subject	-	
Adverse event, non-fatal	3	
Covid-19	-	

Baseline characteristics

Reporting groups		
Reporting group title	Active 50mg	
Reporting group description: -		
Reporting group title	Active 150mg	
Reporting group description: -		
Reporting group title	Active 300mg	
Reporting group description: -		
Reporting group title	Placebo	

Reporting group description: -

Reporting group values	Active 50mg	Active 150mg	Active 300mg
Number of subjects	101	103	100
Age categorical			
Units: Subjects			
Adults (18-64 years)	63	70	74
From 65-84 years	38	33	26
Gender categorical			
Units: Subjects			
Female	79	76	68
Male	22	27	32

Reporting group values	Placebo	Total	
Number of subjects	102	406	
Age categorical			
Units: Subjects			
Adults (18-64 years)	78	285	
From 65-84 years	24	121	
Gender categorical			
Units: Subjects			
Female	77	300	
Male	25	106	

Adverse events

Adverse events information		
Timeframe for reporting adverse events:		
4 weeks		
Assessment type	Systematic	
Dictionary used		
Dictionary name	MedDRA	
Dictionary version	23	
Reporting groups		
Reporting group title	Active - 50mg	
Reporting group description: -		
Reporting group title	Active - 150 mg	
Reporting group description: -		
Reporting group title	Active - 300mg	
Reporting group description: -		
Reporting group title	Placebo	·
Reporting group description: -		

Serious adverse events	Active - 50mg	Active - 150 mg	Active - 300mg
Total subjects affected by serious adverse events			
subjects affected / exposed	0 / 101 (0.00%)	0 / 103 (0.00%)	0 / 100 (0.00%)
number of deaths (all causes)	0	0	0
number of deaths resulting from adverse events			

Serious adverse events	Placebo	
Total subjects affected by serious adverse events		
subjects affected / exposed	0 / 102 (0.00%)	
number of deaths (all causes)	0	
number of deaths resulting from adverse events		

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

Non-serious adverse events	Active - 50mg	Active - 150 mg	Active - 300mg
Total subjects affected by non-serious adverse events			
subjects affected / exposed	5 / 101 (4.95%)	15 / 103 (14.56%)	34 / 100 (34.00%)
Nervous system disorders			

Dysgeusia subjects affected / exposed occurrences (all)	2 / 101 (1.98%)	12 / 103 (11.65%) 12	27 / 100 (27.00%) 27
Headache subjects affected / exposed occurrences (all)	3 / 101 (2.97%)	2 / 103 (1.94%)	1 / 100 (1.00%)
	3	2	1
Hypogeusia subjects affected / exposed occurrences (all)	0 / 101 (0.00%)	1 / 103 (0.97%)	6 / 100 (6.00%)
	0	1	6

Non-serious adverse events	Placebo	
Total subjects affected by non-serious adverse events		
subjects affected / exposed	10 / 102 (9.80%)	
Nervous system disorders		
Dysgeusia		
subjects affected / exposed	3 / 102 (2.94%)	
occurrences (all)	3	
Headache		
subjects affected / exposed	7 / 102 (6.86%)	
occurrences (all)	7	
Hypogeusia		
subjects affected / exposed	0 / 102 (0.00%)	
occurrences (all)	0	

More information

Substantial protocol amendments (globally)

Were there any global substantial amendments to the protocol? Yes

Date	Amendment
11 September 2020	Inclusion of COVID-19 measures. Amendment was not considered substantial in all countries

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Notes:

Interruptions (globally)

Were there any global interruptions to the trial? No

Limitations and caveats

None reported