Clinical trial results:

A Phase III, Multicenter, Randomized, Double-blind, Parallel-group Study to Evaluate the Safety of Once Daily Esomeprazole for the Treatment of Clinically Diagnosed Gastroesophageal Reflux Disease (GERD) in Pediatric and Adolescent Patients 12 to 17 Years, Inclusive

Summary

EudraCT number	2012-001136-61	
Trial protocol	Outside EU/EEA	
Global end of trial date 19 September 2008		
Results information		
Result version number	v1 (current)	
This version publication date 01 February 2017		
First version publication date	16 August 2015	

Trial information

Trial identification		
Sponsor protocol code	D9614C00098	
Additional study identifiers		
ISRCTN number	-	
ClinicalTrials.gov id (NCT number)	-	
WHO universal trial number (UTN)	-	
Notes:		

Sponsors	
Sponsor organisation name	AstraZeneca LP
Sponsor organisation address	1800 Concord Pike, Wilmington, Delaware, United States, 19850
Public contact	AZ Clinical Trial Transparency group, AstraZeneca R&D, ClinicalTrialTransparency@astrazeneca.com
Scientific contact	Marta Illueca, MD, FAAP, AstraZeneca LP, 1 302-885-5514,
Notes:	

Paediatric regulatory details

Is trial part of an agreed paediatric investigation plan (PIP)	Yes
EMA paediatric investigation plan number(s)	EMEA-000331-PIP01-08
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	19 September 2008	
Is this the analysis of the primary completion data?	Yes	
Primary completion date	19 September 2008	
Global end of trial reached?	Yes	
Global end of trial date	19 September 2008	
Was the trial ended prematurely?	No	
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Notes:

General information about the trial

Main objective of the trial:

The primary objective of this study was to evaluate the safety and tolerability of once daily treatment with esomeprazole in pediatric and adolescent patients 12 to 17 years of age, inclusive, with clinically diagnosed GERD.

Protection of trial subjects:

The study was performed in accordance with the ethical principles that have their origin in the Declaration of Helsinki and that are consistent with ICH/Good Clinical Practice and applicable regulatory requirements and the AstraZeneca policy on Bioethics. The final Clinical Study Protocol (CSP), including the final version of the Informed Consent Form and all advertising used to recruit patients, had to be approved by an Institutional Review Board (IRB) before enrollment of any patient into the study.All patients (and their parents/quardians) were provided with all the information necessary to make an informed decision about their participation in the study, including the nature and intended purpose of the study, possible benefits and possible risks. All information in the informed consent form was provided in language understandable to the patients and their parents/guardians. The patient's parent/guardian's signed and dated informed consent as well as the patient's signed and dated assent were obtained before conducting any procedure specifically for the study. If the patient or patient's parents/guardians could not read, the informed consent form (or assent) was signed and dated by an impartial witness. The person who discussed the informed consent information with the patient also signed and dated the form. Patients could be discontinued from study treatment and assessments at any time, voluntary by the patient who was free to discontinue at anay time, or at the discretion of the investigator(s)

Background therapy:

PPI use within 14 days prior to randomization (Visit 2), including over-the-counter (OTC) PRILOSEC® (AstraZeneca LP) was exclusion criteria, as was any prescription or OTC treatment use for symptoms of GERD, such as H2RA or prokinetics, within 3 days (72 hours) prior to randomization (Visit 2). Antacids were allowed, except for those containing bismuth.

Evidence for comparator:		
No comparator group		
Actual start date of recruitment	20 February 2004	
Long term follow-up planned	No	
Independent data monitoring committee (IDMC) involvement?	No	
Notes:		

Notes:

Population of trial subjects

Subjects enrolled per country		
Country: Number of subjects enrolled	France: 5	
Country: Number of subjects enrolled	Italy: 1	
Country: Number of subjects enrolled	Canada: 21	
Country: Number of subjects enrolled	United States: 122	

Worldwide total number of subjects	149
EEA total number of subjects	6

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)	149	
Adults (18-64 years)	0	
From 65 to 84 years	0	
85 years and over	0	

Recruitment

Recruitment details:

First patient enrolled: 20 February 2004 Last patient completed: 04 May 2005

Pre-assignment

Screening details:

Out of 157 patients screened for this study, 8 patients failed to be eligible and never received study medication.

Pre-assignment period milestones		
Number of subjects started	149	
Number of subjects completed	149	

Period 1		
Period 1 title	Treatment (overall period)	
Is this the baseline period?	Yes	
Allocation method	Randomised - controlled	
Blinding used	Double blind	
Roles blinded	Investigator, Monitor, Carer, Data analyst, Assessor, Subject	
Blinding implementation details:		
Double-blind		
Arms		
Are arms mutually exclusive?	Yes	
Arm title	Esomeprazole 20 mg	
Arm description:	•	
Esomeprazole magnesium 20 mg		
Arm type	Experimental	
Investigational medicinal product name	Esomeprazole magnesium	
Investigational medicinal product code		
Other name	NEXIUM	
Pharmaceutical forms	Capsule	
Routes of administration	Oral use	
Dosage and administration details:		
20 mg od orally		
Arm title	Esomeprazole 40 mg	
Arm description:		
Esomeprazole magnesium 40 mg		
Arm type	Experimental	
Investigational medicinal product name	Esomeprazole magnesium	
Investigational medicinal product code		
Other name		
Pharmaceutical forms	Capsule	
Routes of administration	Oral use	
Dosage and administration details:		

40 mg od orally

Number of subjects in period 1	Esomeprazole 20 mg	Esomeprazole 40 mg
Started	76	73
Completed	60	67
Not completed	16	6
Consent withdrawn by subject	4	1
Adverse event, non-fatal	5	1
Lost to follow-up	1	3
Lack of efficacy	2	-
Not eligible, developed discont. crit, 2 not spec.	4	1

Reporting groups		
Reporting group title	Esomeprazole 20 mg	
Reporting group description:		
Esomeprazole magnesium 20 mg		
Reporting group title	Esomeprazole 40 mg	
Reporting group description:		
Esomeprazole magnesium 40 mg		

Reporting group values	Esomeprazole 20 mg	Esomeprazole 40 mg	Total
Number of subjects	76	73	149
Age Categorical			
Age years		·	
Units: Subjects			
In utero	0	0	0
Preterm newborn infants (gestational age < 37 wks)	0	0	0
Newborns (0-27 days)	0	0	0
Infants and toddlers (28 days-23 months)	0	0	0
Children (2-11 years)	0	0	0
Adolescents (12-17 years)	76	73	149
Adults (18-64 years)	0	0	0
From 65-84 years	0	0	0
85 years and over	0	0	0
Age Continuous			
Age (years)			
Units: years			
median	14	14	
full range (min-max)	12 to 17	12 to 17	-
Gender Categorical			
Units: Subjects			
Female	47	42	89
Male	29	31	60

Subject analysis sets			
Subject analysis set title	ITT		
Subject analysis set type	Intention-to-treat		
Subject analysis set description:			
ПТ			
			-
Reporting group values	ITT		
Number of subjects	149		
Age Categorical			

Age Categorical		
Age years		
Units: Subjects		
In utero	0	

Reporting group title	Esomeprazole 20 mg
Reporting group description:	
Esomeprazole magnesium 20 mg	
Reporting group title	Esomeprazole 40 mg
Reporting group description:	
Esomeprazole magnesium 40 mg	
Subject analysis set title	ІТТ
Subject analysis set type	Intention-to-treat
Subject analysis set description:	
ITT	

Primary: Number of patients whose Physician Global Assessment score improved from baseline at their final visit (ITT population)

End point title	Number of patients whose Physician Global Assessment score
	improved from baseline at their final visit (ITT population) ^[1]

End point description:

Number of patients whose Physician's Globle Assessment score improved from baseline

End point type	Primary
End point timeframe:	

From baseline to final Visit

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.

 $\label{eq:constraint} \ensuremath{\mathsf{Justification}}: \ensuremath{\mathsf{Only}}\xspace \ensuremath{\mathsf{Discriptive}}\xspace \ensuremath{\mathsf{statistics}}\xspace \ensuremath{\mathsf{areprox}}\xspace \ensuremath{\mathsf{statistics}}\xspace \ensuremath{\mathsf{areprox}}\xspace \ensuremath{\mathsf{areprox}}\xspace$

End point values	Esomeprazole 20 mg	Esomeprazole 40 mg	ITT	
Subject group type	Reporting group	Reporting group	Subject analysis set	
Number of subjects analysed	76	73	149	
Units: Number of subjects	55	55	110	

Statistical analyses

No statistical analyses for this end point

Secondary: Heartburn Symptom - average change from baseline at final week				
End point title Heartburn Symptom - average change from baseline at fina week				
End point description:				
symptoms were rated on a 4-point scale as follows: 0=none; 1=mild; 2=moderate; 3=severe.				
End point type Secondary				
End point timeframe:				
Baseline to final week				

End point values	Esomeprazole 20 mg	Esomeprazole 40 mg	ITT	
Subject group type	Reporting group	Reporting group	Subject analysis set	
Number of subjects analysed	73 ^[2]	72 ^[3]	145 ^[4]	
Units: Change in average score				
arithmetic mean (standard deviation)	-0.72 (± 0.97)	-0.71 (± 0.92)	-0.72 (± 0.95)	

Notes:

[2] - All patients who had heartburn symptoms at baseline

[3] - All patients who had heartburn symptoms at baseline

[4] - All patients who had heartburn symptoms at baseline

Statistical analyses

No statistical analyses for this end point

Adverse events information		
Timeframe for reporting adverse events:		
During treatment (8weeks)		
Assessment type	Systematic	
Dictionary used		
Dictionary name MedDRA		
Dictionary version	8.0	
Reporting groups		
Reporting group title	Esomeprazole 40 mg	
Reporting group description:		
Esomeprazole magnesium 40 mg		
Reporting group title	Esomeprazole 20 mg	
Reporting group description:		
Esomeprazole magnesium 20 mg		

Serious adverse events	Esomeprazole 40 mg	Esomeprazole 20 mg	
Total subjects affected by serious adverse events			
subjects affected / exposed	0 / 73 (0.00%)	0 / 75 (0.00%)	
number of deaths (all causes)	0	0	
number of deaths resulting from adverse events	0	0	

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

Non-serious adverse events	Esomeprazole 40 mg	Esomeprazole 20 mg	
Total subjects affected by non-serious adverse events			
subjects affected / exposed	57 / 73 (78.08%)	56 / 75 (74.67%)	
Nervous system disorders			
Headache			
subjects affected / exposed	15 / 73 (20.55%)	12 / 75 (16.00%)	
occurrences (all)	57	56	
General disorders and administration site conditions pyrexia			
subjects affected / exposed	2 / 73 (2.74%)	5 / 75 (6.67%)	
occurrences (all)	57	56	
Gastrointestinal disorders			

Abdominal pain	1		I
subjects affected / exposed	9 / 73 (12.33%)	9 / 75 (12.00%)	
occurrences (all)	57	56	
Diarrhoea			
subjects affected / exposed	8 / 73 (10.96%)	6 / 75 (8.00%)	
occurrences (all)	57	56	
Vomiting			
subjects affected / exposed	7 / 73 (9.59%)	5 / 75 (6.67%)	
occurrences (all)	57	56	
Nausea			
subjects affected / exposed	4 / 73 (5.48%)	6 / 75 (8.00%)	
occurrences (all)	57	56	
(,	1	50	
constipation			
subjects affected / exposed	4 / 73 (5.48%)	2 / 75 (2.67%)	
occurrences (all)	57	56	
Respiratory, thoracic and mediastinal			
disorders			
Cough subjects affected / exposed			
	5 / 73 (6.85%)	6 / 75 (8.00%)	
occurrences (all)	57	56	
pharyngolaryngeal pain			
subjects affected / exposed	9 / 73 (12.33%)	1 / 75 (1.33%)	
occurrences (all)	57	56	
Nasal congestion			
subjects affected / exposed	1 / 73 (1.37%)	4 / 75 (5.33%)	
occurrences (all)			
	57	56	
Infections and infestations			
Nasopharyngitis			
subjects affected / exposed	9 / 73 (12.33%)	9 / 75 (12.00%)	
occurrences (all)	57	56	
Upper respiratory tract infection			
subjects affected / exposed	4 / 73 (5.48%)	9 / 75 (12.00%)	
occurrences (all)	57	56	
sinusitis subjects affected / exposed			
	4 / 73 (5.48%)	3 / 75 (4.00%)	
occurrences (all)	57	56	

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