

Clinical trial results:

A phase III, randomised, comparative, open-label study of intravenous iron isomaltoside 1000 (Monofer®) administered as maintenance therapy by single or repeated bolus injections in comparison with intravenous iron sucrose in subjects with stage 5 chronic kidney disease on dialysis therapy (CKD-5D)

Summary

EudraCT number	2010-023471-26	
Trial protocol	GB SE DK PL	
Global end of trial date	28 October 2013	
Results information		
Result version number	v2 (current)	
This version publication date	07 April 2016	
First version publication date	15 July 2015	
Version creation reason	Correction of full data set Incorrect data was discovered during the review process.	

Trial information

Trial identification		
Sponsor protocol code	P-Monofer-CKD-03	
Additional study identifiers		
ISRCTN number	-	
ClinicalTrials.gov id (NCT number)	NCT01222884	
WHO universal trial number (UTN)	-	
Notes:	•	

Sponsors

Sponsor organisation name	Pharmacosmos A/S
Sponsor organisation address	Roervangsvej 30, Holbaek, Denmark, Dk-4300
Public contact	Clinical trial disclosure desk, Pharmacosmos A/S, 045 59485935, trial@pharmacosmos.com
Scientific contact	Clinical trial disclosure desk, Pharmacosmos A/S, 045 59485935, trial@pharmacosmos.com

Notes:

Paediatric regulatory details		
	Is trial part of an agreed paediatric investigation plan (PIP)	No
	Does article 45 of REGULATION (EC) No	No

1901/2006 apply to this trial?

Does article 46 of REGULATION (EC) No No 1901/2006 apply to this trial?

Notes:

Results analysis stage	
Analysis stage	Final
Date of interim/final analysis	28 October 2013
Is this the analysis of the primary completion data?	Yes
Primary completion date	28 October 2013
Global end of trial reached?	Yes
Global end of trial date	28 October 2013
Was the trial ended prematurely?	No

Notes:

General information about the trial

Main objective of the trial:

The primary efficacy objective of the study was to demonstrate that IV iron isomaltoside 1000 is non-inferior to IV iron sucrose determined as ability to maintain haemoglobin (Hb) between 9.5 and 12.5 g/dL in subjects with CKD-5D who were on maintenance iron therapy.

Protection of trial subjects:

The protocol and amendments were approved by local ethics committees/Institutional Review Boards and competent authorities. The trial was conducted in accordance with good clinical practice and the Declaration of Helsinki. Informed consent was obtained in writing prior to any trial-related activities.

Background therapy: -	
Evidence for comparator: -	
Actual start date of recruitment	14 June 2011
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	Poland: 13
Country: Number of subjects enrolled	Sweden: 10
Country: Number of subjects enrolled	United Kingdom: 187
Country: Number of subjects enrolled	Denmark: 11
Country: Number of subjects enrolled	India: 72
Country: Number of subjects enrolled	Romania: 26
Country: Number of subjects enrolled	Russian Federation: 9
Country: Number of subjects enrolled	Switzerland: 19
Country: Number of subjects enrolled	United States: 4
Worldwide total number of subjects	351

247

Notes:

Subjects enrolled per age group	
In utero	0
Preterm newborn - gestational age < 37 wk	0
Newborns (0-27 days)	0

EEA total number of subjects

Infants and toddlers (28 days-23 months)	0
Children (2-11 years)	0
Adolescents (12-17 years)	0
Adults (18-64 years)	195
From 65 to 84 years	146
85 years and over	10

Subject disposition

Recruitment

Recruitment details:

Subjects were screened in the period 14 June 2011 to 10 September 2013. The trial took place at 48 sites (hospitals or private dialysis clinics); 16 centres in India, 14 centres in the UK, 4 in Russia, 4 in Poland, 3 in Sweden, 3 in Switzerland, 2 in Romania, 1 in Denmark, and 1 in the USA.

Pre-assignment

Screening details:

Subjects \geq 18 years of age with a diagnosis of CKD and on haemodialysis therapy for at least 90 days, Hb between 9.5 and 12.5 g/dL (inclusive both values) both at screening visit 1a and screening visit 1b, serum-ferritin < 800 ng/mL, TSAT < 35%, and receiving ESA treatment with stable dose for the previous 4 weeks prior to screening were enrolled.

Arm description:

All subjects received a cumulative dose of 500 mg iron isomaltoside 1000. Subjects in subgroup A1 were administered iron isomaltoside 1000 as a single undiluted IV bolus injection of 500 mg over approximately 2 min at baseline, subjects in subgroup A2 were administered undiluted iron isomaltoside 1000 in split doses of 100 mg at baseline and 200 mg each at week 2 and 4 as IV bolus injections over approximately 2 min.

Arm type	Experimental
Investigational medicinal product name	Iron isomaltoside 1000
Investigational medicinal product code	ATC code: B03AC
Other name	Monofer, Monover, Monofar, Monoferro
Pharmaceutical forms	Solution for injection/infusion
Routes of administration	Intravenous use

Dosage and administration details:

All subjects received a cumulative dose of 500 mg iron isomaltoside 1000. Subjects in subgroup A1 were administered iron isomaltoside 1000 as a single undiluted IV bolus injection of 500 mg over approximately 2 min at baseline, subjects in subgroup A2 were administered undiluted iron isomaltoside 1000 in split doses of 100 mg at baseline and 200 mg each at week 2 and 4 as IV bolus injections over approximately 2 min.

Iron isomaltoside 1000 is available as a dark brown, non-transparent aqueous solution for injection/infusion containing 100 mg iron/mL with pH between 5.0 and 7.0.

Arm title Group B, iron sucrose	

Arm description:

All subjects received a cumulative dose of 500 mg iron sucrose.

Subjects in group B were administered undiluted iron sucrose in split doses of 100 mg at baseline and 200 mg each at week 2 and 4.

Arm type	Active comparator
Investigational medicinal product name	Iron sucrose
Investigational medicinal product code	ATC code: B03AB02,B03AC02
Other name	Venofer
Pharmaceutical forms	Concentrate for solution for infusion, Solution for injection
Routes of administration	Intravenous use

Dosage and administration details:

Subjects in group B were administered undiluted iron sucrose in split doses of 100 mg at baseline and 200 mg each at week 2 and 4. The doses of iron sucrose were administered as per local summary product of characteristics or package insert and/or local hospital guidelines, as applicable. All dosages were administered during dialysis, at least 30 min after the start and at least 1 h before the end of dialysis.

Number of subjects in period 1	Group A, iron isomaltoside 1000	Group B, iron sucrose
Started	234	117
Completed	210	113
Not completed	24	4
Adverse event, serious fatal	3	-
Consent withdrawn by subject	6	2
Physician decision	1	-
Subject decided to go on holiday for 8 we	1	-
Kidney transplantation	1	-
Patient had been given IV iron by dialysis staff	1	-
Adverse event, non-fatal	8	1
Did not fulfil the inclusion/exclusion criteria	3	-
Transplant	-	1

Baseline characteristics

Reporting groups

Reporting group title	Group A, iron isomaltoside 1000

Reporting group description:

All subjects received a cumulative dose of 500 mg iron isomaltoside 1000. Subjects in subgroup A1 were administered iron isomaltoside 1000 as a single undiluted IV bolus injection of 500 mg over approximately 2 min at baseline, subjects in subgroup A2 were administered undiluted iron isomaltoside 1000 in split doses of 100 mg at baseline and 200 mg each at week 2 and 4 as IV bolus injections over approximately 2 min.

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Reporting group title	Group B, iron sucrose

Reporting group description:

All subjects received a cumulative dose of 500 mg iron sucrose.

Subjects in group B were administered undiluted iron sucrose in split doses of 100 mg at baseline and 200 mg each at week 2 and 4.

Reporting group values	Group A, iron isomaltoside 1000	Group B, iron sucrose	Total
Number of subjects	234	117	351
Age categorical			
Units: Subjects			
In utero			0
Preterm newborn infants (gestational age < 37 wks)			0
Newborns (0-27 days)			0
Infants and toddlers (28 days-23 months)			0

Subject analysis set description:

The full analysis set (FAS) population (N=341) included all subjects who were randomized into the trial, received at least one dose of the trial drug, and had at least one post-baseline Hb assessment.

Subject analysis set title	Per protocol (PP) analysis set
Subject analysis set type	Per protocol

Subject analysis set description:

The per protocol (PP) population (N=306) included all subjects in the FAS who did not have any major protocol deviation of clinical or statistical relevance.

Reporting group values	Safety analysis set	Full analysis set (FAS)	Per protocol (PP) analysis set
Number of subjects	344	341	306
Age categorical			
Units: Subjects			
In utero			
Preterm newborn infants (gestational age < 37 wks)			
Newborns (0-27 days)			
Infants and toddlers (28 days-23 months)			
Children (2-11 years)			
Adolescents (12-17 years)			
Adults (18-64 years)			
From 65-84 years			
85 years and over			
Age continuous			
Age is calculated by subtracting the scre	ening visit date with t	he birth date.	
Units: years			
arithmetic mean	60	60.1	59.6
standard deviation	± 16	± 15.9	± 15.9
Gender categorical			
Units: Subjects			
Female	117	117	100
Male	227	224	206

End points

End points reporting groups

Reporting group title	Group A, iron isomaltoside 1000

Reporting group description:

All subjects received a cumulative dose of 500 mg iron isomaltoside 1000. Subjects in subgroup A1 were administered iron isomaltoside 1000 as a single undiluted IV bolus injection of 500 mg over approximately 2 min at baseline, subjects in subgroup A2 were administered undiluted iron isomaltoside 1000 in split doses of 100 mg at baseline and 200 mg each at week 2 and 4 as IV bolus injections over approximately 2 min.

Reporting group title Group B, iron sucrose

Reporting group description:

All subjects received a cumulative dose of 500 mg iron sucrose.

Subjects in group B were administered undiluted iron sucrose in split doses of 100 mg at baseline and 200 mg each at week 2 and 4.

Subject analysis set title	Safety analysis set
Subject analysis set type	Safety analysis

Subject analysis set description:

The safety population (N=344) included all subjects who were randomized and received at least one dose of the trial drug.

Subject analysis set title	Full analysis set (FAS)
Subject analysis set type	Full analysis

Subject analysis set description:

The full analysis set (FAS) population (N=341) included all subjects who were randomized into the trial, received at least one dose of the trial drug, and had at least one post-baseline Hb assessment.

Subject analysis set title	Per protocol (PP) analysis set
Subject analysis set type	Per protocol

Subject analysis set description:

The per protocol (PP) population (N=306) included all subjects in the FAS who did not have any major protocol deviation of clinical or statistical relevance.

Primary: Proportion of subjects who were able to maintain Hb between 9.5 and 12.5 g/dL (both values included) at week 6, FAS

End point title	Proportion of subjects who were able to maintain Hb between
	9.5 and 12.5 g/dL (both values included) at week 6, FAS

End point description:

Proportion of subjects who were able to maintain Hb between 9.5 and 12.5 g/dL (both values included) at week 6.

The analysis was performed on the FAS.

End point type	Primary

End point timeframe:

Proportion of subjects who were able to maintain Hb between 9.5 and 12.5 g/dL (both values included) at week 6.

End point values	Group A, iron isomaltoside 1000	Group B, iron sucrose	
Subject group type	Reporting group	Reporting group	
Number of subjects analysed	226	115	
Units: Proportion of subjects			
Responder	187	95	

Non-responder	39	20	
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Statistical analysis title	Non-inferiority tested by risk difference
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Statistical analysis description:

A generalised linear model using the identity link function was used to compare the proportion of subjects with Hb concentration between 9.5 and 12.5 g/dL (both values included) at week 6 using the last observation carried forward approach.

The number of subjects may differ from the analysis population if data is missing.

Comparison groups	Group A, iron isomaltoside 1000 v Group B, iron sucrose
Number of subjects included in analysis	341
Analysis specification	Pre-specified
Analysis type	non-inferiority ^[1]
P-value	= 0.0106 [2]
Method	Risk differences
Parameter estimate	Risk difference (RD)
Point estimate	1
Confidence interval	
level	95 %
sides	2-sided
lower limit	-7.4
upper limit	9.4

Notes:

[1] - Treatment and stratum (serum-ferritin (<100 versus \ge 100 ng/mL) were used as factors and baseline value as a covariate.

With a 2:1 randomisation, a two-sided significance level of 0.05, and a non-inferiority margin of 10 % points, there was approximately 80 % power to demonstrate non-inferiority with 214 subjects in group A and 107 subjects in group B.

[2] - As the trial was designed to demonstrate non-inferiority, the analyses of FAS and PP population would lead to similar conclusions and therefore the analyses for both analysis sets needed to be powered properly.

Primary: Proportion of subjects who were able to maintain Hb between 9.5 and 12.5 g/dL (both values included) at week 6, PP

End point title	Proportion of subjects who were able to maintain Hb between
	9.5 and 12.5 g/dL (both values included) at week 6, PP

End point description:

Proportion of subjects who were able to maintain Hb between 9.5 and 12.5 g/dL (both values included) at week 6.

The analysis was performed on the PP analysis set.

End point type Primary		Primary
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End point timeframe:

Proportion of subjects who were able to maintain Hb between 9.5 and 12.5 g/dL (both values included) at week 6.

End point values	Group A, iron isomaltoside 1000	Group B, iron sucrose	
Subject group type	Reporting group	Reporting group	
Number of subjects analysed	199	107	
Units: Proportion of subjects			
Responder	167	88	
Non-responder	32	19	

Statistical analysis description:

A generalised linear model using the identity link function was used to compare the proportion of subjects with Hb concentration between 9.5 and 12.5 g/dL (both values included) at week 6 using the last observation carried forward approach.

The number of subjects may differ from the analysis population if data is missing.

The named of subjects may affer from	the analysis population in data is imposing.
Comparison groups	Group B, iron sucrose v Group A, iron isomaltoside 1000
Number of subjects included in analysis	306
Analysis specification	Pre-specified
Analysis type	non-inferiority ^[3]
P-value	= 0.0057 [4]
Method	Risk difference
Parameter estimate	Risk difference (RD)
Point estimate	2.2
Confidence interval	
level	95 %
sides	2-sided
lower limit	-6.4
upper limit	10.9

Notes:

- [3] Treatment and stratum (serum-ferritin (<100 versus ≥100 ng/mL) were used as factors and baseline value as a covariate. With a 2:1 randomisation, a two-sided significance level of 0.05, and a non-inferiority margin of 10 % points, there was approximately 80 % power to demonstrate non-inferiority with 214 subjects in group A and 107 subjects in group B.
- [4] As the trial was designed to demonstrate non-inferiority, the analyses of FAS and PP population would lead to similar conclusions and therefore the analyses for both analysis sets needed to be powered properly.

End point title	Change in Hb concentration from baseline to week 2
End point description:	
Change in Hb concentration	from baseline to week 2.
Analysis performed on the	FAS.
Analysis performed on the End point type	FAS. Secondary

End point values	Group A, iron isomaltoside 1000	Group B, iron sucrose	
Subject group type	Reporting group	Reporting group	
Number of subjects analysed	219	115	
Units: g/dL			
arithmetic mean (standard deviation)	0.04 (± 0.71)	-0.05 (± 0.69)	

Statistical alialyses			
Statistical analysis title	Test for superiority, MMRM		
Statistical analysis description:			
	s includes treatment, visit, treatment*visit interactions and /mL)) ,country as factors and baseline values as covariates.		
Comparison groups	Group A, iron isomaltoside 1000 v Group B, iron sucrose		
Number of subjects included in analysis	334		
Analysis specification	Pre-specified		
Analysis type	superiority		
P-value	= 0.1239		
Method	MMRM		
Parameter estimate	Mean difference (final values)		
Point estimate	0.1138		
Confidence interval			
level	95 %		
sides	2-sided		
lower limit	-0.031		
upper limit	0.259		
Variability estimate	Standard error of the mean		
Dispersion value	0.0737		

Secondary: Change in Hb concentration from baseline to week 4		
End point title Change in Hb concentration from baseline to week 4		
End point description:		
Change in Hb concentration	from baseline to week 4.	
Analysis performed on the	FAS.	
End point type Secondary		
End point timeframe:		
Change in Hb concentration	from baseline to week 4	

End point values	Group A, iron isomaltoside 1000	Group B, iron sucrose	
Subject group type	Reporting group	Reporting group	
Number of subjects analysed	213	114	
Units: g/dL			
arithmetic mean (standard deviation)	0.01 (± 0.91)	-0.03 (± 0.68)	

Statistical allalyses				
Statistical analysis title	Test for superiority, MMRM			
Statistical analysis description:				
	s includes treatment, visit, treatment*visit interactions and /mL)), country as factors and baseline values as covariates.			
Comparison groups	Group A, iron isomaltoside 1000 v Group B, iron sucrose			
Number of subjects included in analysis	327			
Analysis specification	Pre-specified			
Analysis type	superiority			
P-value	= 0.5233			
Method	MMRM			
Parameter estimate	Mean difference (final values)			
Point estimate	0.0546			
Confidence interval				
level	95 %			
sides	2-sided			
lower limit	-0.113			
upper limit	0.223			
Variability estimate	Standard error of the mean			
Dispersion value	0.0854			

Secondary: Change in Hb concentration from baseline to week 6.		
End point title Change in Hb concentration from baseline to week 6.		
End point description:		
Change in Hb concentration from basel	ine to week 6.	
Analysis performed on the FAS.		
End point type Secondary		
End point timeframe:		
Change in Hb concentration from baseli	ine to week 6	

End point values	Group A, iron isomaltoside 1000	Group B, iron sucrose	
Subject group type	Reporting group	Reporting group	
Number of subjects analysed	216	113	
Units: g/dL			
arithmetic mean (standard deviation)	-0.07 (± 1.11)	-0.06 (± 0.99)	

Statistical analyses			
Statistical analysis title	Test for superiority, MMRM		
Statistical analysis description:			
	includes treatment, visit, treatment*visit interactions and mL)) ,country as factors and baseline values as covariates.		
Comparison groups	Group A, iron isomaltoside 1000 v Group B, iron sucrose		
Number of subjects included in analysis	329		
Analysis specification	Pre-specified		
Analysis type	superiority		
P-value	= 0.8557		
Method	MMRM		
Parameter estimate	Mean difference (final values)		
Point estimate	-0.0069		
Confidence interval			
level	95 %		
sides	2-sided		
lower limit	-0.204		
upper limit	0.246		
Variability estimate	Standard error of the mean		
Dispersion value	0.1143		

Secondary: Change in s-iron concentration from baseline to week 1		
End point title Change in s-iron concentration from baseline to week 1		
End point description:	•	
Change in s-iron concentration	n from baseline to week 1.	
Analysis performed on the FAS	5.	
End point type Secondary		
Life point type	,	
End point timeframe:	,	

End point values	Group A, iron isomaltoside 1000	Group B, iron sucrose	
Subject group type	Reporting group	Reporting group	
Number of subjects analysed	221	112	
Units: micromol/L			
arithmetic mean (standard deviation)	1.45 (± 4.35)	0.75 (± 3.3)	

Statistical analyses	statistical analyses			
Statistical analysis title	Test for superiority, MMRM			
Statistical analysis description:	•			
	s includes treatment, visit, treatment*visit interactions and /mL)) ,country as factors and baseline values as covariates.			
Comparison groups	Group A, iron isomaltoside 1000 v Group B, iron sucrose			
Number of subjects included in analysis	333			
Analysis specification	Pre-specified			
Analysis type	superiority			
P-value	= 0.1429			
Method	MMRM			
Parameter estimate	Mean difference (final values)			
Point estimate	0.5793			
Confidence interval	·			
level	95 %			
sides	2-sided			
lower limit	-0.197			
upper limit	1.355			
Variability estimate	Standard error of the mean			
Dispersion value	0.3942			

Secondary: Change in s-iron concentration from baseline to week 2		
End point title	Change in s-iron concentration from baseline to week 2	
End point description:		

Change in s-iron concentration from baseline to week 2.

Analysis performed on the FAS.

End point type Secondary

End point timeframe:

Change in s-iron concentration from baseline to week 2.

End point values	Group A, iron isomaltoside 1000	Group B, iron sucrose	
Subject group type	Reporting group	Reporting group	
Number of subjects analysed	220	115	
Units: micromol/L			
arithmetic mean (standard deviation)	1.07 (± 4.12)	0.64 (± 5.75)	

Statistical analysis title Test for superiority, MMRM				
Statistical analysis description:				
	includes treatment, visit, treatment*visit interactions and (mL)), country as factors and baseline values as covariates.			
Comparison groups	Group A, iron isomaltoside 1000 v Group B, iron sucrose			
Number of subjects included in analysis	335			
Analysis specification	Pre-specified			
Analysis type	superiority			
P-value	= 0.5277			
Method	MMRM			
Parameter estimate	Mean difference (final values)			
Point estimate	0.3761			
Confidence interval				
level	95 %			
sides	2-sided			
lower limit	-0.797			
upper limit	1.549			
Variability estimate	Standard error of the mean			
Dispersion value	0.5943			

End point title Change in s-iron concentration from baseline to week				
End point description:	•			
Change in a iron concentration	on from baseline to week 4			
Change in S-non concentration	on from baseline to week 4.			
Analysis performed on the FA				
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Analysis performed on the FA	AS.			

End point values	Group A, iron isomaltoside 1000	Group B, iron sucrose	
Subject group type	Reporting group	Reporting group	
Number of subjects analysed	212	114	
Units: micromol/L			
arithmetic mean (standard deviation)	0.81 (± 4.14)	1.08 (± 4.21)	

Statistical alialyses			
Statistical analysis title	Test for superiority, MMRM		
Statistical analysis description:			
	s includes treatment, visit, treatment*visit interactions and /mL)) ,country as factors and baseline values as covariates.		
Comparison groups	Group A, iron isomaltoside 1000 v Group B, iron sucrose		
Number of subjects included in analysis	326		
Analysis specification	Pre-specified		
Analysis type	superiority		
P-value	= 0.438		
Method	MMRM		
Parameter estimate	Mean difference (final values)		
Point estimate	-0.3576		
Confidence interval			
level	95 %		
sides	2-sided		
lower limit	-1.265		
upper limit	0.549		
Variability estimate	Standard error of the mean		
Dispersion value	0.4603		

Secondary: Change in s-iron concentration from baseline to week 6					
End point title Change in s-iron concentration from baseline to week					
End point description:					
Change in s-iron concentration from baseline to week 6.					
Analysis performed on the FAS)•				
Analysis performed on the FAS End point type	Secondary				

End point values	Group A, iron isomaltoside 1000	Group B, iron sucrose	
Subject group type	Reporting group	Reporting group	
Number of subjects analysed	216	113	
Units: micromol/L			
arithmetic mean (standard deviation)	0.82 (± 5.21)	0.76 (± 4.18)	

Statistical analysis title	Test for superiority, MMRM				
Statistical analysis description:					
	includes treatment, visit, treatment*visit interactions and (mL)), country as factors and baseline values as covariates.				
Comparison groups	Group A, iron isomaltoside 1000 v Group B, iron sucrose				
Number of subjects included in analysis	329				
Analysis specification	Pre-specified				
Analysis type	superiority				
P-value	= 0.9894				
Method	MMRM				
Parameter estimate	Mean difference (final values)				
Point estimate	0.0066				
Confidence interval					
level	95 %				
sides	2-sided				
lower limit	-0.965				
upper limit	0.978				
Variability estimate	Standard error of the mean				
Dispersion value	0.4932				

Secondary: Change in transferrin saturation (TSAT) concentration from baseline to week ${\bf 1}$

End point title	Change in transferrin saturation (TSAT) concentration from
	baseline to week 1

End point description:

Change in transferrin saturation (TSAT) concentration from baseline to week 1.

Analysis performed on the FAS.

End point type Secondary

End point timeframe:

Change in transferrin saturation (TSAT) concentration from baseline to week 1.

End point values	Group A, iron isovnRltds0de0 1000	J 0 241 0.5 240	4 761.5 S 1 w 0	J000RG[]
Subject group type	Reporting group			
Number of subjects analysed	221			
Units: percentage				
arithmetic mean (standard deviation)	2.8 (± 18.47)			

Statistical analysis title

End point values	Group A, iron isomaltoside 1000	Group B, iron sucrose	
Subject group type	Reporting group	Reporting group	
Number of subjects analysed	220	115	
Units: percentage			
arithmetic mean (standard deviation)	2.45 (± 20.75)	1.44 (± 9.62)	

Statistical analysis title Test for superiority, MMRM				
Statistical analysis description:				
	s includes treatment, visit, treatment*visit interactions and 'mL)) ,country as factors and baseline values as covariates.			
Comparison groups	Group A, iron isomaltoside 1000 v Group B, iron sucrose			
Number of subjects included in analysis	335			
Analysis specification	Pre-specified			
Analysis type	superiority			
P-value	= 0.355			
Method	MMRM			
Parameter estimate	Mean difference (final values)			
Point estimate	1.1992			
Confidence interval				
level	95 %			
sides	2-sided			
lower limit	-1.35			
upper limit	3.748			
Variability estimate	Standard error of the mean			
Dispersion value	1.294			

Secondary: Change in transferrin saturation (TSAT) concentration from baseline to week 4

End point title	Change in transferrin saturation (TSAT) concentration from
	baseline to week 4

End point description:

Change in transferrin saturation (TSAT) concentration from baseline to week 4.

Analysis performed on the FAS.

End point type Secondary

End point timeframe:

Change in transferrin saturation (TSAT) concentration from baseline to week 4.

End point values	Group A, iron isomaltoside 1000	Group B, iron sucrose	
Subject group type	Reporting group	Reporting group	
Number of subjects analysed	212	114	
Units: percentage			
arithmetic mean (standard deviation)	1.8 (± 19.26)	2.85 (± 8.98)	

Statistical analysis title Test for superiority, MMRM				
Statistical analysis description:				
	s includes treatment, visit, treatment*visit interactions and 'mL)) ,country as factors and baseline values as covariates.			
Comparison groups	Group A, iron isomaltoside 1000 v Group B, iron sucrose			
Number of subjects included in analysis	326			
Analysis specification	Pre-specified			
Analysis type	superiority			
P-value	= 0.3487			
Method	MMRM			
Parameter estimate	Mean difference (final values)			
Point estimate	-0.9972			
Confidence interval				
level	95 %			
sides	2-sided			
lower limit	-3.09			
upper limit	1.095			
Variability estimate	Standard error of the mean			
Dispersion value	1.0617			

Secondary: Change in transferrin saturation (TSAT) concentration from baseline to week 6

End point title	Change in transferrin saturation (TSAT) concentration from
	baseline to week 6

End point description:

Change in transferrin saturation (TSAT) concentration from baseline to week 6.

Analysis performed on the FAS.

End point type Secondary

End point timeframe:

Change in transferrin saturation (TSAT) concentration from baseline to week 6.

End point values	Group A, iron isomaltoside 1000	Group B, iron sucrose	
Subject group type	Reporting group	Reporting group	
Number of subjects analysed	216	113	
Units: percentage			
arithmetic mean (standard deviation)	2.29 (± 19.43)	2.42 (± 8.62)	

Statistical analysis title	Test for superiority, MMRM
Statistical analysis description:	
	includes treatment, visit, treatment*visit interactions and (mL)), country as factors and baseline values as covariates.
Comparison groups	Group A, iron isomaltoside 1000 v Group B, iron sucrose
Number of subjects included in analysis	329
Analysis specification	Pre-specified
Analysis type	superiority
P-value	= 0.9845
Method	MMRM
Parameter estimate	Mean difference (final values)
Point estimate	-0.0207
Confidence interval	
level	95 %
sides	2-sided
lower limit	-2.118
upper limit	2.077
Variability estimate	Standard error of the mean
Dispersion value	1.0654

Secondary: Change in s-ferritin concentration from baseline to week 1			
End point title	Change in s-ferritin concentration from baseline to week 1		
End point description:			
Change in s-ferritin concent	ration from baseline to week 1.		
Analysis performed on the F	FAS.		
Analysis performed on the F End point type	FAS. Secondary		

End point values	Group A, iron isomaltoside 1000	Group B, iron sucrose	
Subject group type	Reporting group	Reporting group	
Number of subjects analysed	221	113	
Units: microg/L			
arithmetic mean (standard deviation)	156.75 (± 148.91)	48.43 (± 75.36)	

Statistical analysis title	Test for superiority, MMRM
Statistical analysis description:	
The mixed model for repeated measures	s includes treatment, visit, treatment*visit interactions and (mL)), country as factors and baseline values as covariates.
Comparison groups	Group A, iron isomaltoside 1000 v Group B, iron sucrose
Number of subjects included in analysis	334
Analysis specification	Pre-specified
Analysis type	superiority
P-value	< 0.0001
Method	MMRM
Parameter estimate	Mean difference (final values)
Point estimate	107.8382
Confidence interval	
level	95 %
sides	2-sided
lower limit	87.987
upper limit	127.689
Variability estimate	Standard error of the mean
Dispersion value	10.0818

Secondary: Change in s-ferritin concentration from baseline to week 2				
End point title	Change in s-ferritin concentration from baseline to week 2			
End point description:				
Change in s-ferritin concentration from baseline to week 2.				
Analysis performed on the FA	AS.			
	AS. Secondary			
Analysis performed on the FA End point type End point timeframe:				

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End point values	Group A, iron isomaltoside 1000	Group B, iron sucrose	
Subject group type	Reporting group	Reporting group	
Number of subjects analysed	220	115	
Units: microg/L			
arithmetic mean (standard deviation)	142.58 (± 187.82)	20.85 (± 94.84)	

Statistical analysis title	rsis title Test for superiority, MMRM				
Statistical analysis description:					
	s includes treatment, visit, treatment*visit interactions and /mL)) ,country as factors and baseline values as covariates.				
Comparison groups	Group A, iron isomaltoside 1000 v Group B, iron sucrose				
Number of subjects included in analysis	335				
Analysis specification	Pre-specified				
Analysis type	superiority				
P-value	< 0.0001				
Method	MMRM				
Parameter estimate	Mean difference (final values)				
Point estimate	123.36				
Confidence interval					
level	95 %				
sides	2-sided				
lower limit	96.449				
upper limit	150.271				
Variability estimate	Standard error of the mean				
Dispersion value	13.6719				

Secondary: Change in s-ferritin concentration from baseline to week 4				
End point title	Change in s-ferritin concentration from baseline to week 4			
End point description:				
Change in s-ferritin concentration from baseline to week 4.				
Analysis performed on the FAS.				
, ,				
End point type	Secondary			
, ,	Secondary			

End point values	Group A, iron isomaltoside 1000	Group B, iron sucrose	
Subject group type	Reporting group	Reporting group	
Number of subjects analysed	212	114	
Units: microg/L			
arithmetic mean (standard deviation)	128.04 (± 157.75)	86.33 (± 126.79)	

Statistical analysis title Test for superiority, MMRM			
Statistical analysis description:			
	s includes treatment, visit, treatment*visit interactions and 'mL)), country as factors and baseline values as covariates		
Comparison groups	Group A, iron isomaltoside 1000 v Group B, iron sucrose		
Number of subjects included in analysis	326		
Analysis specification	Pre-specified		
Analysis type	superiority		
P-value	= 0.002		
Method	MMRM		
Parameter estimate	Mean difference (final values)		
Point estimate	49.3393		
Confidence interval			
level	95 %		
sides	2-sided		
lower limit	18.174		
upper limit	80.505		
Variability estimate	Standard error of the mean		
Dispersion value	15.8282		

Secondary: Change in s-ferritin concentration from baseline to week 6				
End point title	Change in s-ferritin concentration from baseline to week 6			
End point description:				
Change in s-ferritin concentration from baseline to week 6.				
Analysis performed on the FAS.				
Analysis periorities on the LAS.				
· ·	Secondary			
End point type End point timeframe:	Secondary			

End point values	Group A, iron isomaltoside 1000	Group B, iron sucrose	
Subject group type	Reporting group	Reporting group	
Number of subjects analysed	216	114	
Units: microg/L			
arithmetic mean (standard deviation)	136.2 (± 154.59)	156.3 (± 183.63)	

Statistical analysis title Test for superiority, MMRM				
Statistical analysis description:				
	s includes treatment, visit, treatment*visit interactions and (mL)), country as factors and baseline values as covariates.			
Comparison groups	Group A, iron isomaltoside 1000 v Group B, iron sucrose			
Number of subjects included in analysis	330			
Analysis specification	Pre-specified			
Analysis type	superiority			
P-value	= 0.4489			
Method	MMRM			
Parameter estimate	Mean difference (final values)			
Point estimate	-15.0585			
Confidence interval				
level	95 %			
sides	2-sided			
lower limit	-54.196			
upper limit	24.079			
Variability estimate	Standard error of the mean			
Dispersion value	19.8434			

Secondary: Change in reticulocyte count from baseline to week 1			
End point title	Change in reticulocyte count from baseline to week 1		
End point description:			
Change in reticulocyte count from baseline to week 1.			
Analysis performed on the FAS.			
End point type	Secondary		
	Secondary		

End point values	Group A, iron isomaltoside 1000	Group B, iron sucrose	
Subject group type	Reporting group	Reporting group	
Number of subjects analysed	212	108	
Units: percentage			
arithmetic mean (standard deviation)	0.12 (± 0.42)	-0.02 (± 0.38)	

Statistical analysis title	Test for superiority, MMRM
Statistical analysis description:	
	includes treatment, visit, treatment*visit interactions and (mL)), country as factors and baseline values as covariates.
Comparison groups	Group A, iron isomaltoside 1000 v Group B, iron sucrose
Number of subjects included in analysis	320
Analysis specification	Pre-specified
Analysis type	superiority
P-value	= 0.0006
Method	MMRM
Parameter estimate	Mean difference (final values)
Point estimate	0.154
Confidence interval	
level	95 %
sides	2-sided
lower limit	0.066
upper limit	0.242
Variability estimate	Standard error of the mean
Dispersion value	0.0445

Secondary: Change in reticulocyte count from baseline to week 2		
End point title	Change in reticulocyte count from baseline to week 2	
End point description:		
Change in reticulocyte cour	t from baseline to week 2.	
Analysis performed on the	FAS.	
End point type	Secondary	
End point timeframe:	•	
Change in reticulocyte cour	t from baseline to week 2	

End point values	Group A, iron isomaltoside 1000	Group B, iron sucrose	
Subject group type	Reporting group	Reporting group	
Number of subjects analysed	211	111	
Units: percentage			
arithmetic mean (standard deviation)	0.05 (± 0.45)	0.02 (± 0.4)	

Statistical analysis title	Test for superiority, MMRM
Statistical analysis description:	
	includes treatment, visit, treatment*visit interactions and (mL)), country as factors and baseline values as covariates.
Comparison groups	Group A, iron isomaltoside 1000 v Group B, iron sucrose
Number of subjects included in analysis	322
Analysis specification	Pre-specified
Analysis type	superiority
P-value	= 0.3448
Method	MMRM
Parameter estimate	Mean difference (final values)
Point estimate	0.0439
Confidence interval	
level	95 %
sides	2-sided
lower limit	-0.047
upper limit	0.135
Variability estimate	Standard error of the mean
Dispersion value	0.0464

Secondary: Change in reticulocyte count from baseline to week 4				
End point title	Change in reticulocyte count from baseline to week 4			
End point description:				
Change in reticulocyte count from baseline to week 4.				
Analysis performed on the FA	S.			
Analysis performed on the FA End point type	S. Secondary			

End point values	Group A, iron isomaltoside 1000	Group B, iron sucrose	
Subject group type	Reporting group	Reporting group	
Number of subjects analysed	204	110	
Units: percentage			
arithmetic mean (standard deviation)	0.05 (± 0.47)	0.03 (± 0.36)	

Statistical analysis title	Test for superiority, MMRM			
Statistical analysis description:				
	includes treatment, visit, treatment*visit interactions and (mL)), country as factors and baseline values as covariates.			
Comparison groups	Group A, iron isomaltoside 1000 v Group B, iron sucrose			
Number of subjects included in analysis	314			
Analysis specification	Pre-specified			
Analysis type	superiority			
P-value	= 0.5171			
Method	MMRM			
Parameter estimate	Mean difference (final values)			
Point estimate	0.0302			
Confidence interval				
level	95 %			
sides	2-sided			
lower limit	-0.061			
upper limit	0.122			
Variability estimate	Standard error of the mean			
Dispersion value	0.0465			

Secondary: Change in reticulocyte count from baseline to week 6			
End point title Change in reticulocyte count from baseline to week 6			
End point description:	<u> </u>		
Change in reticulocyte count	t from baseline to week 6.		
Analysis performed on the F	AS.		
End point type	Secondary		
End point timeframe:	•		
Change in reticulocyte count	t from haseline to week 6		

End point values	Group A, iron isomaltoside 1000	Group B, iron sucrose	
Subject group type	Reporting group	Reporting group	
Number of subjects analysed	207	109	
Units: percentage			
arithmetic mean (standard deviation)	0.06 (± 0.53)	0 (± 0.4)	

otation and your				
Statistical analysis title	Test for superiority, MMRM			
Statistical analysis description:				
	includes treatment, visit, treatment*visit interactions and (mL)), country as factors and baseline values as covariates.			
Comparison groups	Group A, iron isomaltoside 1000 v Group B, iron sucrose			
Number of subjects included in analysis	316			
Analysis specification	Pre-specified			
Analysis type	superiority			
P-value	= 0.1564			
Method	MMRM			
Parameter estimate	Mean difference (final values)			
Point estimate	0.0727			
Confidence interval				
level	95 %			
sides	2-sided			
lower limit	-0.028			
upper limit	0.173			
Variability estimate	Standard error of the mean			
Dispersion value	0.0512			

Secondary: Number of subjects in each randomisation group who discontinued
study because of lack of response or intolerance of investigational drugs

•	Number of subjects in each randomisation group who
	discontinued study because of lack of response or intolerance
	of investigational drugs

End point description:

Number of subjects in each randomisation group who discontinued study because of lack of response or intolerance of investigational drugs.

The analysis was performed on the FAS.

End point type	Secondary
End point timeframe:	

The endpoint covers the complete trial period.

End point values	Group A, iron isomaltoside 1000	Group B, iron sucrose	
Subject group type	Reporting group	Reporting group	
Number of subjects analysed	226	115	
Units: Number of subjects			
Discontinued due to intolerance/lack of response	1	0	
Discontinued due to other reasons	15	2	

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Sta	LIS	.ıcaı	alla	Iyってっ

Statistical analysis title

Statistical analysis description:	
• • • • • • • • • • • • • • • • • • •	s includes treatment, visit, treatment*visit interactions and /mL)), country as factors and baseline values as covariates.
Comparison groups	Group A, iron isomaltoside 1000 v Group B, iron sucrose
Number of subjects included in analysis	319
Analysis specification	Pre-specified
Analysis type	superiority
P-value	= 0.4653
Method	MMRM
Parameter estimate	Mean difference (final values)
Point estimate	1.4173
Confidence interval	
level	95 %
sides	2-sided
lower limit	-2.402
upper limit	5.237
Variability estimate	Standard error of the mean
Dispersion value	1.9379

Secondary: Change in total quality of life (QoL) score (LASA: Energy level) from baseline to week 6			
End point title	Change in total quality of life (QoL) score (LASA: Energy level) from baseline to week 6		
End point description:			
Change in total quality of life (QoL)	score (LASA: Energy level) from baseline to week 6.		
The analysis was performed on the I	FAS.		
End point type	Secondary		
End point timeframe:			
Change in total quality of life (QoL)	score (LASA: Energy level) from baseline to week 6.		

End point values	Group A, iron isomaltoside 1000	Group B, iron sucrose	
Subject group type	Reporting group	Reporting group	
Number of subjects analysed	204	113	
Units: QoL score			
arithmetic mean (standard deviation)	3.9 (± 18.91)	2.3 (± 17.54)	

Statistical analysis title	Superiority tested by MMRM	
Statistical analysis description:		
The mixed model for repeated measures includes treatment, visit, treatment*visit interactions and stratum (s-ferritin ($<100 \text{ vs.} >=100 \text{ ng/mL}$), country as factors and baseline values as covariates.		
Comparison groups	Group A, iron isomaltoside 1000 v Group B, iron sucrose	

Number of subjects included in analysis	317
Analysis specification	Pre-specified
Analysis type	superiority
P-value	= 0.9539
Method	MMRM
Parameter estimate	Mean difference (final values)
Point estimate	0.1111
Confidence interval	
level	95 %
sides	2-sided
lower limit	-3.667
upper limit	3.889
Variability estimate	Standard error of the mean
Dispersion value	1.9182

Secondary: Change in total quality of life (QoL) score (LASA: Ability to do daily activities) from baseline to week 4		
End point title	Change in total quality of life (QoL) score (LASA: Ability to do daily activities) from baseline to week 4	
End point description:		
Change in total quality of life (QoL) score (LASA: Ability to do daily activities) from baseline to week 4.		
The analysis was performed	on the FAS.	
End point type	Secondary	
End point timeframe:	•	
Change in total quality of life	e (OoL) score (LASA: Ability to do daily activities) from baseline to week 4.	

End point values	Group A, iron isomaltoside 1000	Group B, iron sucrose	
Subject group type	Reporting group	Reporting group	
Number of subjects analysed	208	111	
Units: QoL score			
arithmetic mean (standard deviation)	2.2 (± 21.03)	-0.2 (± 14.25)	

Statistical analysis title Test for superiority, MMRM		
Statistical analysis description:		
The mixed model for repeated measures includes treatment, visit, treatment*visit interactions and stratum (s-ferritin ($<100 \text{ vs.} >=100 \text{ ng/mL}$)), country as factors and baseline values as covariates.		
Comparison groups	Group A, iron isomaltoside 1000 v Group B, iron sucrose	

Number of subjects included in analysis	319
Analysis specification	Pre-specified
Analysis type	superiority
P-value	= 0.4086
Method	MMRM
Parameter estimate	Mean difference (final values)
Point estimate	1.5719
Confidence interval	
level	95 %
sides	2-sided
lower limit	-2.168
upper limit	5.312
Variability estimate	Standard error of the mean
Dispersion value	1.8991

Secondary: Change in total quality of life (QoL) score (LASA: Ability to do daily activities) from baseline to week 6		
End point title	Change in total quality of life (QoL) score (LASA: Ability to do daily activities) from baseline to week 6	
End point description:		
Change in total quality of li	e (QoL) score (LASA: Ability to do daily activities) from baseline to week 6.	
The analysis was performed	on the FAS.	
End point type	Secondary	
End point timeframe:	•	
Change in total quality of life	e (QoL) score (LASA: Ability to do daily activities) from baseline to week 6.	

End point values	Group A, iron isomaltoside 1000	Group B, iron sucrose	
Subject group type	Reporting group	Reporting group	
Number of subjects analysed	204	113	
Units: QoL score			
arithmetic mean (standard deviation)	3.3 (± 19.6)	2.8 (± 17.91)	

,		
Statistical analysis title Test for superiority, MMRM		
Statistical analysis description:		
The mixed model for repeated measures includes treatment, visit, treatment*visit interactions and stratum (s-ferritin ($<100 \text{ vs.} >=100 \text{ ng/mL}$), country as factors and baseline values as covariates.		
Comparison groups	group A, iron isomaltoside 1000 v Group B, iron sucrose	

317
Pre-specified
superiority
= 0.6734
MMRM
Mean difference (final values)
-0.8519
95 %
2-sided
-4.829
3.125
Standard error of the mean
2.0186

Secondary: Change in total qualifrom baseline to week 4	ty of life (QoL) score (LASA: Overall quality of life)
•	Change in total quality of life (QoL) score (LASA: Overall quality of life) from baseline to week 4

End point description:

Change in total quality of life (QoL) score (LASA: Overall quality of life) from baseline to week 4.

The analysis was performed on the FAS.

End point type	Secondary
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End point timeframe:

Change in total quality of life (QoL) score (LASA: Overall quality of life) from baseline to week 4.

End point values	Group A, iron isomaltoside 1000	Group B, iron sucrose	
Subject group type	Reporting group	Reporting group	
Number of subjects analysed	208	111	
Units: QoL score			
arithmetic mean (standard deviation)	1.7 (± 17.88)	-0.3 (± 15.63)	

Statistical analysis title Test for superiority, MMRM		
Statistical analysis description:		
The mixed model for repeated measures includes treatment, visit, treatment*visit interactions and stratum (s-ferritin ($<100 \text{ vs.} >=100 \text{ ng/mL}$)), country as factors and baseline values as covariates.		
Comparison groups	Group A, iron isomaltoside 1000 v Group B, iron sucrose	

Number of subjects included in analysis	319
Analysis specification	Pre-specified
Analysis type	superiority
P-value	= 0.5711
Method	MMRM
Parameter estimate	Mean difference (final values)
Point estimate	1.0565
Confidence interval	
level	95 %
sides	2-sided
lower limit	-2.614
upper limit	4.727
Variability estimate	Standard error of the mean
Dispersion value	1.8621

Secondary: Change in total quality of life (QoL) score (LASA: Overall quality of life) from baseline to week 6		
End point title	Change in total quality of life (QoL) score (LASA: Overall quality of life) from baseline to week 6	
End point description:		
Change in total quality of life (QoL) score (LASA: Overall quality of life) from baseline to week 6.		
The analysis was performed on the FAS.		
End point type	Secondary	

End point timeframe: Change in total quality of life (QoL) score (LASA: Overall quality of life) from baseline to week 6.

End point values	Group A, iron isomaltoside 1000	Group B, iron sucrose	
Subject group type	Reporting group	Reporting group	
Number of subjects analysed	204	113	
Units: QoL score			
arithmetic mean (standard deviation)	2 (± 18.56)	0.1 (± 15.65)	

Statistical analysis title Test for superiority, MMRM		
Statistical analysis description:		
The mixed model for repeated measures includes treatment, visit, treatment*visit interactions and stratum (s-ferritin ($<100 \text{ vs.} >=100 \text{ ng/mL}$), country as factors and baseline values as covariates.		
Comparison groups Group A, iron isomaltoside 1000 v Group B, iron sucrose		

Number of subjects included in analysis	317
Analysis specification	Pre-specified
Analysis type	superiority
P-value	= 0.7964
Method	MMRM
Parameter estimate	Mean difference (final values)
Point estimate	0.4718
Confidence interval	
level	95 %
sides	2-sided
lower limit	-3.125
upper limit	4.069
Variability estimate	Standard error of the mean
Dispersion value	1.8264

Secondary: Change in restless legs syndrome (RLS) symptoms (Cambridge Hopkins-RLS questionnaire (CH-RLSq) score) from baseline to week 6 in subjects with RLS symptoms at baseline

End point title	Change in restless legs syndrome (RLS) symptoms (Cambridge
	Hopkins-RLS questionnaire (CH-RLSq) score) from baseline to
	week 6 in subjects with RLS symptoms at baseline

End point description:

Change in restless legs syndrome (RLS) symptoms (Cambridge Hopkins-RLS questionnaire (CH-RLSq) score) from baseline to week 6 in subjects with RLS symptoms at baseline.

The analysis was performed on the FAS.

End point type	Secondary

End point timeframe:

Change in restless legs syndrome (RLS) symptoms (Cambridge Hopkins-RLS questionnaire (CH-RLSq) score) from baseline to week 6 in subjects with RLS symptoms at baseline.

End point values	Group A, iron isomaltoside 1000	Group B, iron sucrose	
Subject group type	Reporting group	Reporting group	
Number of subjects analysed	68	40	
Units: RLS score			
arithmetic mean (standard deviation)	-0.7 (± 7.6)	-1.3 (± 5.37)	

Statistical analysis title Superiority tested by ANCOVA		
Statistical analysis description:		
The ANCOVA mixed model includes treatment and stratum as factors and baseline value as covariates.		
Comparison groups	Group A, iron isomaltoside 1000 v Group B, iron sucrose	

Number of subjects included in analysis	108
Analysis specification	Pre-specified
Analysis type	superiority
P-value	= 0.7267
Method	ANCOVA
Parameter estimate	Mean difference (final values)
Point estimate	0.4033
Confidence interval	
level	95 %
sides	2-sided
lower limit	-1.88
upper limit	2.686
Variability estimate	Standard error of the mean
Dispersion value	1.1507

Adverse events

Adverse events information

Timeframe for reporting adverse events:

From the time a subject had signed the ICF and until he/she had completed the study, all AEs/SAEs were collected in the CRF. The SAEs occurring after study termination were re-ported if considered related to the study treatment.

Adverse event reporting additional description:

The principle investigator (PI) was responsible for ensuring that all AEs observed by PI or reported by the subject were properly collected and recorded in the subject's medical record as well as on the AE form.

Assessment type	Systematic
Dictionary used	
Dictionary name	MedDRA
Dictionary version	16.1
Reporting groups	
Reporting group title	Group B, iron sucrose
Reporting group description:	
-	ive dose of 500 mg iron sucrose. Subjects in group B were administered loses of 100 mg at baseline and 200 mg each at week 2 and 4.

Reporting group description:

Reporting group title

All subjects received a cumulative dose of 500 mg iron isomaltoside 1000. Subjects in subgroup A1 were administered iron isomaltoside 1000 as a single undiluted IV bolus injection of 500 mg over approximately 2 min at baseline, subjects in subgroup A2 were administered undiluted iron isomaltoside 1000 in split doses of 100 mg at baseline and 200 mg each at week 2 and 4 as IV bolus injections over approximately 2 min.

Group A, iron isomaltoside 1000

Serious adverse events	Group B, iron sucrose	Group A, iron isomaltoside 1000	
Total subjects affected by serious adverse events			
subjects affected / exposed	6 / 114 (5.26%)	22 / 230 (9.57%)	
number of deaths (all causes)	0	3	
number of deaths resulting from adverse events			
Vascular disorders			
Aortic stenosis			
subjects affected / exposed	0 / 114 (0.00%)	1 / 230 (0.43%)	
occurrences causally related to treatment / all	0 / 0	0 / 1	
deaths causally related to treatment / all	0 / 0	0 / 0	
General disorders and administration site conditions			
Puncture site haemorrhage			
subjects affected / exposed	0 / 114 (0.00%)	1 / 230 (0.43%)	
occurrences causally related to treatment / all	0 / 0	0 / 1	
deaths causally related to treatment / all	0 / 0	0 / 0	

Sudden death			
subjects affected / exposed	0 / 114 (0.00%)	1 / 230 (0.43%)	
occurrences causally related to treatment / all	0 / 0	0 / 1	
deaths causally related to treatment / all	0 / 0	0 / 1	
Immune system disorders			
Hypersensitivity			
subjects affected / exposed	0 / 114 (0.00%)	1 / 230 (0.43%)	
occurrences causally related to treatment / all	0 / 0	1 / 1	
deaths causally related to treatment / all	0 / 0	0 / 0	
Reproductive system and breast disorders			
Prostatitis			
subjects affected / exposed	1 / 114 (0.88%)	0 / 230 (0.00%)	
occurrences causally related to treatment / all	0 / 1	0 / 0	
deaths causally related to treatment / all	0/0	0 / 0	
Respiratory, thoracic and mediastinal disorders			
Dyspnoea			
subjects affected / exposed	1 / 114 (0.88%)	0 / 230 (0.00%)	
occurrences causally related to treatment / all	1 / 1	0 / 0	
deaths causally related to treatment / all	0/0	0 / 0	
Injury, poisoning and procedural complications			
Anaesthetic complication			
subjects affected / exposed	1 / 114 (0.88%)	0 / 230 (0.00%)	
occurrences causally related to treatment / all	0 / 1	0 / 0	
deaths causally related to treatment / all	0/0	0 / 0	
Arteriovenous fistula site haemorrhage			<u> </u>
subjects affected / exposed	0 / 114 (0.00%)	2 / 230 (0.87%)	
occurrences causally related to treatment / all	0 / 0	0 / 2	
deaths causally related to treatment / all	0/0	0 / 0	
Fall			
subjects affected / exposed	0 / 114 (0.00%)	3 / 230 (1.30%)	
occurrences causally related to treatment / all	0 / 0	0/3	
deaths causally related to treatment / all	0/0	0 / 0	
Femoral neck fracture	l i		
	•		

subjects affected / exposed	0 / 114 (0.00%)	1 / 230 (0.43%)	
occurrences causally related to treatment / all	0 / 0	0 / 1	
deaths causally related to treatment / all	0 / 0	0 / 0	
Road traffic accident			
subjects affected / exposed	1 / 114 (0.88%)	0 / 230 (0.00%)	
occurrences causally related to treatment / all	0 / 1	0 / 0	
deaths causally related to treatment / all	0 / 0	0 / 0	
Vascular graft occlusion			
subjects affected / exposed	0 / 114 (0.00%)	1 / 230 (0.43%)	
occurrences causally related to treatment / all	0 / 0	0 / 1	
deaths causally related to treatment / all	0 / 0	0 / 1	
Cardiac disorders			
Acute coronary syndrome			
subjects affected / exposed	0 / 114 (0.00%)	1 / 230 (0.43%)	
occurrences causally related to treatment / all	0 / 0	0 / 1	
deaths causally related to treatment / all	0 / 0	0 / 0	
Acute myocardial infarction			
subjects affected / exposed	0 / 114 (0.00%)	1 / 230 (0.43%)	
occurrences causally related to treatment / all	0 / 0	0 / 1	
deaths causally related to treatment / all	0 / 0	0 / 0	
Nervous system disorders			
Brain stem infarction			
subjects affected / exposed	0 / 114 (0.00%)	1 / 230 (0.43%)	
occurrences causally related to treatment / all	0 / 0	0 / 1	
deaths causally related to treatment / all	0 / 0	0 / 1	
Gastrointestinal disorders			
Duodenal ulcer haemorrhage			
subjects affected / exposed	0 / 114 (0.00%)	1 / 230 (0.43%)	
occurrences causally related to treatment / all	0 / 0	0 / 1	
deaths causally related to treatment / all	0 / 0	0 / 0	
Gingival bleeding			
subjects affected / exposed	0 / 114 (0.00%)	1 / 230 (0.43%)	
occurrences causally related to treatment / all	0 / 0	0 / 1	
deaths causally related to treatment / all	0 / 0	0 / 0	

Subjects affected / exposed occurrences causally related to treatment / all deaths causally related to treatm	Lower gastrointestinal haemorrhage	1		
Occurrences causally related to treatment / all O / 0		0 / 114 (0.00%)	1 / 230 (0.43%)	
Hepatobiliary disorders				
Biliary colic subjects affected / exposed occurrences causally related to treatment / all deaths causally related to treatment / all		0 / 0	0 / 0	
subjects affected / exposed occurrences causally related to treatment / all deaths causally related to treatment / all Renal and urinary disorders Nephrolithiasis subjects affected / exposed occurrences causally related to treatment / all deaths causally related to treatment / all	Hepatobiliary disorders			
occurrences causally related to treatment / all deaths causally related to treatment / all o/0 0/0 0/0 0/0 0/0 0/0 0/0 0/0 0/0 0/0	' '			
treatment / all deaths causally related to treatment / all occurrences causally related to treatment / all deaths causally related to treatment / all occurrences causally related to occurren	subjects affected / exposed	0 / 114 (0.00%)	1 / 230 (0.43%)	
treatment / all		0 / 0	0 / 1	
Nephrolithiasis subjects affected / exposed occurrences causally related to treatment / all deaths causally related to treatment / all occurrences causally related to treatment / all deaths causally related to treatment / all		0 / 0	0 / 0	
subjects affected / exposed occurrences causally related to treatment / all deaths causally related to treatment / all musculoskeletal and connective tissue disorders Musculoskeletal pain subjects affected / exposed occurrences causally related to treatment / all deaths causally related to treatment / all	Renal and urinary disorders			
occurrences causally related to treatment / all deaths causally related to treatment / all of treatment / al	·			
treatment / all deaths causally related to treatment / all o/0 0/0 0/0 Musculoskeletal and connective tissue disorders Musculoskeletal pain subjects affected / exposed occurrences causally related to treatment / all deaths causally related to treatment / all 0/0 0/0 0/0 Infections and infestations Arteriovenous fistula site infection subjects affected / exposed occurrences causally related to treatment / all 0/0 0/1 0/1 0/1 0/1 0/1 0/1 0/1 0/1 0/1	subjects affected / exposed	0 / 114 (0.00%)	1 / 230 (0.43%)	
Musculoskeletal and connective tissue disorders Musculoskeletal pain subjects affected / exposed occurrences causally related to treatment / all deaths causally related to		0 / 0	0 / 1	
disorders Musculoskeletal pain subjects affected / exposed occurrences causally related to treatment / all deaths causally related to treatment subjects affected / exposed occurrences causally related to treatment / all Infections and infestations Arteriovenous fistula site infection subjects affected / exposed occurrences causally related to treatment / all deaths causally related to treatment / all Device related infection subjects affected / exposed occurrences causally related to treatment / all Device related infection subjects affected / exposed occurrences causally related to treatment / all deaths causally related to treatment / all Infected fistula subjects affected / exposed occurrences causally related to treatment / all deaths causally related to treatment / all		0 / 0	0 / 0	
subjects affected / exposed occurrences causally related to treatment / all deaths causally related to treatment / all deaths causally related to treatment / all Infections and infestations Arteriovenous fistula site infection subjects affected / exposed occurrences causally related to treatment / all deaths causally related to treatment / all Device related infection subjects affected / exposed O / 114 (0.00%) O / 0 I / 230 (0.43%) O / 0 O / 1 Device related infection subjects affected / exposed O / 114 (0.00%) O / 0 I / 230 (0.43%) O / 0 O / 0 I / 230 (0.43%) O / 0 O / 1 Infected fistula subjects affected / exposed O / 0 O / 0 Infected fistula subjects affected / exposed O / 114 (0.00%) O / 0 Infected fistula subjects affected / exposed O / 114 (0.00%) O / 0 Infected fistula subjects affected / exposed O / 114 (0.00%) O / 0 Infected fistula subjects affected / exposed O / 114 (0.00%) O / 0 Infected fistula subjects affected / exposed O / 114 (0.00%) O / 0 Infected fistula subjects affected / exposed O / 0 / 0 O / 0 Infected fistula subjects affected / exposed O / 0 / 0 O / 0 Infected fistula subjects affected / exposed O / 0 / 0 O / 0 Infected fistula subjects affected / exposed O / 0 / 0 O / 0 Infected fistula subjects affected / exposed O / 0 / 0 O / 0 Infected fistula subjects affected / exposed O / 0 / 0 O / 0				
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Infections and infestations Arteriovenous fistula site infection subjects affected / exposed occurrences causally related to treatment / all deaths causally related to treatment / all deaths causally related to treatment / all Infected fistula subjects affected / exposed occurrences causally related to treatment / all Infected fistula subjects affected / exposed occurrences causally related to treatment / all deaths causally related to treatment / all Infected fistula subjects affected / exposed occurrences causally related to treatment / all deaths causally related to treatment / all O / 0 O / 0 O / 0 O / 1 Infected fistula subjects affected / exposed occurrences causally related to treatment / all deaths causally related to treatment / all O / 0 O / 0 O / 0		0 / 0	0 / 1	
Arteriovenous fistula site infection subjects affected / exposed occurrences causally related to treatment / all deaths causally related to treatment / all occurrences causally related to occurrences causal		0 / 0	0 / 0	
subjects affected / exposed	Infections and infestations			
occurrences causally related to treatment / all deaths causally related to treatment / all Device related infection subjects affected / exposed occurrences causally related to treatment / all deaths causally related to treatment / all Infected fistula subjects affected / exposed occurrences causally related to treatment / all Infected fistula subjects affected / exposed occurrences causally related to treatment / all deaths causally related to treatment / all	Arteriovenous fistula site infection			
treatment / all deaths causally related to treatment / all Device related infection subjects affected / exposed occurrences causally related to treatment / all deaths causally related to treatment / all Infected fistula subjects affected / exposed occurrences causally related to treatment / all occurrences causally related to treatment / all occurrences causally related to treatment / all deaths causally related to treatment / all occurrences causally related to treatment / all	subjects affected / exposed	0 / 114 (0.00%)	1 / 230 (0.43%)	
treatment / all 0 / 0 0 / 0 Device related infection subjects affected / exposed 0 / 114 (0.00%) 1 / 230 (0.43%) occurrences causally related to treatment / all deaths causally related to treatment / all		0 / 0	0 / 1	
subjects affected / exposed occurrences causally related to treatment / all deaths causally related to treatment / all Infected fistula subjects affected / exposed occurrences causally related to treatment / all occurrences causally related to treatment / all deaths causally related to treatment / all deaths causally related to treatment / all occurrences causally related to treatment / all		0 / 0	0 / 0	
occurrences causally related to treatment / all deaths causally related to treatment / all	Device related infection			
treatment / all deaths causally related to treatment / all Infected fistula subjects affected / exposed occurrences causally related to treatment / all deaths causally related to treatment / all deaths causally related to treatment / all o / 0 0 / 0 1 / 230 (0.43%) 0 / 0 0 / 0 0 / 0	subjects affected / exposed	0 / 114 (0.00%)	1 / 230 (0.43%)	
treatment / all 0 / 0 0 / 0 Infected fistula subjects affected / exposed 0 / 114 (0.00%) 1 / 230 (0.43%) occurrences causally related to treatment / all deaths causally related to treatment / all 0 / 0 0 / 0		0 / 0	0 / 1	
subjects affected / exposed occurrences causally related to treatment / all deaths causally related to treatment / all 0 / 114 (0.00%) 0 / 0 0 / 1 0 / 0 0 / 0		0 / 0	0 / 0	
subjects affected / exposed occurrences causally related to treatment / all deaths causally related to treatment / all 0 / 114 (0.00%) 0 / 0 0 / 1 0 / 0 0 / 0	Infected fistula]		ļ
occurrences causally related to treatment / all		0 / 114 (0.00%)	1 / 230 (0.43%)	
deaths causally related to treatment / all 0 / 0 0 / 0				
	deaths causally related to	0 / 0	0 / 0	
	Lower respiratory tract infection			

subjects affected / exposed	1 / 114 (0.88%)	1 / 230 (0.43%)	
occurrences causally related to treatment / all	0 / 1	0 / 1	
deaths causally related to treatment / all	0 / 0	0 / 0	
Respiratory tract infection			
subjects affected / exposed	0 / 114 (0.00%)	1 / 230 (0.43%)	
occurrences causally related to treatment / all	0 / 0	0 / 1	
deaths causally related to treatment / all	0 / 0	0 / 0	
Staphylococcal bacteraemia			
subjects affected / exposed	1 / 114 (0.88%)	0 / 230 (0.00%)	
occurrences causally related to treatment / all	1 / 1	0 / 0	
deaths causally related to treatment / all	0 / 0	0 / 0	
Metabolism and nutrition disorders			
Fluid overload			
subjects affected / exposed	0 / 114 (0.00%)	1 / 230 (0.43%)	
occurrences causally related to treatment / all	0 / 0	0 / 1	
deaths causally related to treatment / all	0 / 0	0 / 0	

Frequency threshold for reporting non-serious adverse events: 2 %

Non-serious adverse events	Group B, iron sucrose	Group A, iron isomaltoside 1000	
Total subjects affected by non-serious adverse events			
subjects affected / exposed	15 / 114 (13.16%)	38 / 230 (16.52%)	
Investigations			
C-reactive protein increased			

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Diarrhoea			
subjects affected / exposed	2 / 114 (1.75%)	5 / 230 (2.17%)	
occurrences (all)	2	5	
Musculoskeletal and connective tissue disorders			
Pain in extremity			
subjects affected / exposed	0 / 114 (0.00%)	5 / 230 (2.17%)	
occurrences (all)	0	6	
Infections and infestations			
Lower respiratory tract infection			
subjects affected / exposed	3 / 114 (2.63%)	3 / 230 (1.30%)	
occurrences (all)	3	4	
Nasopharyngitis			
subjects affected / exposed	1 / 114 (0.88%)	6 / 230 (2.61%)	
occurrences (all)	1	6	
Metabolism and nutrition disorders			
Hyperphosphataemia			
subjects affected / exposed	4 / 114 (3.51%)	5 / 230 (2.17%)	
occurrences (all)	4	5	

More information

Substantial protocol amendments (globally)

Were there any global substantial amendments to the protocol? Yes

Date	Amendment
11 January 2012	 Primary endpoint was changed from "change in Hb concentrations from baseline to week 6" to "proportion of subjects able to maintain Hb between 10 and 12.5 g/dl (both values included) at week 6" In the secondary endpoint "change in Hb concentration from baseline to week 2 and 4", additional time point "week 6" was added Study design was revised in terms of number of treatment groups and extension of enrolment period to 18 months, and study centre at Norway was removed from the list of participating countries Text regarding iron isomaltoside 1000 was updated Inclusion criteria pertaining to Hb, ESA treatment, and subjects not on IV iron and exclusion criterion 3 were modified to bring more clarity to text Study flowchart was revised to clarify that height should be measured only at screening The option of performing blood pregnancy test instead of UPT was added Iron sucrose infusion time was changed to "according to SmPC" Iron sucrose test dose administration was changed to "according to SmPC or local guidelines" Statistical section was revised as per changes in the study endpoints (primary endpoint and first secondary endpoint) The possibility of re-screening the screen-failure subjects once 2 weeks after the screening visit was added Appendix 2 related to CH-RLSq was updated
10 July 2012	 Total study duration was increased to approximately 19 months, and study centres were rephrased as Europe, USA, and India In inclusion criterion # 5, the target Hb range between "10 and 12.5 g/dL" was revised to "9.5 and 12.5 g/dL" Additional text was included in sections: dosage and administration, prohibited medication, screen failure, and rescreening Re-screening was allowed "up to 3 times" The frequency of planned review by safety review committee was decreased from one meeting every 2 months to one meeting every 4 months for a feasible study conduct

Notes:

Interruptions (globally)

Were there any global interruptions to the trial? No

Limitations and caveats

None reported