



EUROPEAN MEDICINES AGENCY
SCIENCE MEDICINES HEALTH

31 August 2023
EMA/547473/2023
Human Medicines Evaluation Division

List of nationally authorised medicinal products

Active substance(s): furosemide

Procedure no.: PSUSA/00001491/202301



Product full name (in authorisation country as per Art. 57 data)	MRP/DCP or CP Authorisation number (if not available then insert not available)	National Authorisation Number of product in the member state	MAH of product in the member state	Member State where product is authorised
Furesis 20 mg tabletit	not available	8681	ORION CORPORATION	FI
Furesis 20 mg tabletter	not available	8681	ORION CORPORATION	FI
Furesis 40 mg tabletit	not available	2675	ORION CORPORATION	FI
Furesis 40 mg tabletter	not available	2675	ORION CORPORATION	FI
Furesis 500 mg tabletit	not available	7115	ORION CORPORATION	FI
Furesis 500 mg tabletter	not available	7115	ORION CORPORATION	FI
Furorese 120 mg long Hartkapseln, retardiert	not available	27324.02.00	HEXAL AG	DE
Furorese 125 mg Tabletten	not available	6913686.01.00	HEXAL AG	DE
FUORESE 125 mg tablety	not available	50/048/99-C	HEXAL AG	CZ
FUORESE 250 mg tablety	not available	50/049/99-C	HEXAL AG	CZ
Furorese 40 mg tablety	not available	50/047/99-C	HEXAL AG	CZ
FUORESE 500 mg tablety	not available	50/050/99-C	HEXAL AG	CZ
Furorese® 20 mg/2 ml injekt Injektionslösung	not available	7259.00.02	HEXAL AG	DE

Product full name (in authorisation country as per Art. 57 data)	MRP/DCP or CP Authorisation number (if not available then insert not available)	National Authorisation Number of product in the member state	MAH of product in the member state	Member State where product is authorised
Furorese® 250 mg Tabletten	not available	37633.00.00	HEXAL AG	DE
Furorese® 250 mg/25 ml Infusionslösung	not available	7259.00.00	HEXAL AG	DE
Furorese® 40 mg/4 ml injekt, Injektionslösung	not available	39036.00.00	HEXAL AG	DE
Furorese® 80 mg Tabletten	not available	37865.01.00	HEXAL AG	DE
Furosemid 125 - 1 A Pharma®	not available	40792.00.00	1 A PHARMA GMBH	DE
Furosemid 250 - 1 A Pharma®	not available	37630.00.00	1 A PHARMA GMBH	DE
Furosemide 10 mg/ml oral solution	IE/H/1026/003	PA22697/011/003	SYRI PHARMA LIMITED	IE
Furosemide 20 mg tablets	not available	PL 28278/0097	IPCA LABORATORIES UK LIMITED	XI
Furosemide 20 mg Tablets	not available	PL 40496/0043	BRILLPHARMA LIMITED	XI
Furosemide 20mg in 2ml Solution for Injection	not available	PA0073/059/004	MERCURY PHARMACEUTICALS (IRELAND) LTD.	IE
Furosemide 20mg in 2ml Solution for Injection	not available	MA079/00601	MERCURY PHARMACEUTICALS (IRELAND) LTD.	MT
Furosemide 20mg Tablets	not available	PL 11311/0710	TILLOMED LABORATORIES LTD	XI
Furosemide 20mg Tablets	not available	PL 11311/0710	TILLOMED LABORATORIES LTD	XI

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Furosemide 20mg Tablets	not available	PL 11311/0710	TILLOMED LABORATORIES LTD	XI
Furosemide 20mg Tablets	not available	PL 11311/0710	TILLOMED LABORATORIES LTD	XI
Furosemide 20mg Tablets	not available	PL 11311/0710	TILLOMED LABORATORIES LTD	XI
Furosemide 20mg Tablets	not available	PL 11311/0710	TILLOMED LABORATORIES LTD	XI
Furosemide 20mg Tablets	not available	PL 11311/0710	TILLOMED LABORATORIES LTD	XI
Furosemide 20mg Tablets	not available	PL 11311/0710	TILLOMED LABORATORIES LTD	XI
Furosemide 250mg in 25ml Solution for Injection/Infusion	not available	PA0073/059/006	MERCURY PHARMACEUTICALS (IRELAND) LTD.	IE
Furosemide 40 mg Tablets	not available	PL 40496/0044	BRILLPHARMA LIMITED	XI
Furosemide 40mg Tablets	not available	PL 11311/0711	TILLOMED LABORATORIES LTD	XI
Furosemide 40mg Tablets	not available	PL 11311/0711	TILLOMED LABORATORIES LTD	XI
Furosemide 40mg Tablets	not available	PL 11311/0711	TILLOMED LABORATORIES LTD	XI
Furosemide 40mg Tablets	not available	PL 11311/0711	TILLOMED LABORATORIES LTD	XI
Furosemide 40mg Tablets	not available	PL 11311/0711	TILLOMED LABORATORIES LTD	XI

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Furosemide 40mg Tablets	not available	PL 11311/0711	TILLOMED LABORATORIES LTD	XI
Furosemide 40mg Tablets	not available	PL 11311/0711	TILLOMED LABORATORIES LTD	XI
Furosemide 40mg Tablets	not available	PL 11311/0711	TILLOMED LABORATORIES LTD	XI
Furosemide 40mg Tablets	not available	PL 11311/0711	TILLOMED LABORATORIES LTD	XI
Furosemide 40mg Tablets	not available	PL 11311/0711	TILLOMED LABORATORIES LTD	XI
Furosemide 40mg Tablets	not available	PL 11311/0711	TILLOMED LABORATORIES LTD	XI
Furosemide 4mg/ml Oral Solution	IE/H/1026/001	PA22697/011/001	SYRI PHARMA LIMITED	IE
Furosemide 4mg/ml Oral Solution	UK/H/5871/001	PL 39307/0047	SYRI LIMITED T/A THAME LABORATORIES	XI
Furosemide 50mg in 5ml Solution for Injection	not available	PA 73/59/5	MERCURY PHARMACEUTICALS (IRELAND) LTD.	IE
Furosemide 50mg in 5ml Solution for Injection	not available	MA079/00603	MERCURY PHARMACEUTICALS (IRELAND) LTD.	MT
Furosemide 8mg/ml Oral Solution	IE/H/1026/002	PA22697/011/002	SYRI PHARMA LIMITED	IE
Furosemide 8mg/ml Oral Solution	UK/H/5871/002	PL 39307/0048	SYRI LIMITED T/A THAME LABORATORIES	XI
Furosemide Teva 20 mg, tabletten	not available	RVG 10185	TEVA NEDERLAND B.V.	NL

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FUROSEMIDE ZENTIVA 500 mg, comprimé sécable	not available	34009 550 008 3 9	ZENTIVA FRANCE	FR
FUROSEMIDE ZENTIVA 500 mg, comprimé sécable	not available	34009 300 022 1 6	ZENTIVA FRANCE	FR
FUROSEMIDE ZENTIVA 500 mg, comprimé sécable	not available	34009 300 022 0 9	ZENTIVA FRANCE	FR
Impugan 10 mg/ml dropar til inntöku, lausn	SE/H/1341/003	IS/1/13/039/03	TEVA B.V	IS
Impugan 10 mg/ml orala droppar, lösning	SE/H/1341/003	10777	ACTAVIS GROUP PTC EHF.	SE
Impugan 20 mg tabletter	SE/H/1341/002	14915	ACTAVIS GROUP PTC EHF.	SE
Impugan 20 mg töflur	SE/H/1341/002	IS/1/13/039/01	TEVA B.V	IS
LASILIX 10 mg/ml, solution buvable	not available	34009 330 009 1 2	SANOFI-AVENTIS FRANCE	FR
LASILIX 20 mg/2 ml, solution injectable en ampoule	not available	586 880-3	SANOFI-AVENTIS FRANCE	FR
LASILIX 20 mg/2 ml, solution injectable en ampoule	not available	305 733-1	SANOFI-AVENTIS FRANCE	FR
LASILIX 20 mg/2 ml, solution injectable en ampoule	not available	581 250-1	SANOFI-AVENTIS FRANCE	FR
LASILIX 20 mg/2 ml, solution injectable en ampoule	not available	553 192-0	SANOFI-AVENTIS FRANCE	FR
LASILIX 20 mg/2 ml, solution injectable en ampoule	not available	559 215-2	SANOFI-AVENTIS FRANCE	FR

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Lasilix 40 mg scored tablets	not available	MA1359/02201	SANOFI S.R.L.	MT
LASILIX 40 mg, comprimé sécable	not available	34009 372 377 9 6	SANOFI-AVENTIS FRANCE	FR
LASILIX 40 mg, comprimé sécable	not available	34009 305 734 8 8	SANOFI-AVENTIS FRANCE	FR
LASILIX 40 mg, comprimé sécable	not available	34009 352 813 8 8	SANOFI-AVENTIS FRANCE	FR
LASILIX 40 mg, comprimé sécable	not available	34009 553 193 7 5	SANOFI-AVENTIS FRANCE	FR
LASILIX FAIBLE 20 mg, comprimé	not available	34009 321 033 0 0	SANOFI-AVENTIS FRANCE	FR
LASILIX FAIBLE 20 mg, comprimé	not available	34009 556 653 9 7	SANOFI-AVENTIS FRANCE	FR
LASILIX FAIBLE 20 mg, comprimé	not available	34009 321 555 6 5	SANOFI-AVENTIS FRANCE	FR
LASILIX FAIBLE 20 mg, comprimé	not available	34009 321 032 4 9	SANOFI-AVENTIS FRANCE	FR
LASILIX FAIBLE 20 mg, comprimé	not available	34009 372 376 2 8	SANOFI-AVENTIS FRANCE	FR
LASILIX RETARD 60 mg, gélule	not available	34009 324 922 0 6	SANOFI-AVENTIS FRANCE	FR
LASILIX RETARD 60 mg, gélule	not available	34009 556 652 2 9	SANOFI-AVENTIS FRANCE	FR
LASILIX RETARD 60 mg, gélule	not available	34009 324 923 7 4	SANOFI-AVENTIS FRANCE	FR

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LASILIX RETARD 60 mg, gélule	not available	34009 372 378 5 7	SANOFI-AVENTIS FRANCE	FR
LASILIX SPECIAL 250 mg/25 ml, solution injectable en ampoule	not available	34009 278 193 5 3	SANOFI-AVENTIS FRANCE	FR
LASILIX SPECIAL 250 mg/25 ml, solution injectable en ampoule	not available	34009 321 390 8 8	SANOFI-AVENTIS FRANCE	FR
LASILIX SPECIAL 500 mg, comprimé sécable	not available	34009 321 391 4 9	SANOFI-AVENTIS FRANCE	FR
LASILIX SPECIAL 500 mg, comprimé sécable	not available	34009 575 374 4 9	SANOFI-AVENTIS FRANCE	FR
LASILIX SPECIAL 500 mg, comprimé sécable	not available	34009 338 453 8 4	SANOFI-AVENTIS FRANCE	FR
Lasix	not available	19672	SANOFI WINTHROP INDUSTRIE	CY
Lasix	not available	41423/07/27-05-2008	SANOFI-AVENTIS MONOPROSOPI A.E.B.E	GR
Lasix	not available	41423/07/27-05-2008	SANOFI-AVENTIS MONOPROSOPI A.E.B.E	GR
Lasix	not available	41423/07/27-05-2008	SANOFI-AVENTIS MONOPROSOPI A.E.B.E	GR
Lasix	not available	41426/07/27-05-2008	SANOFI-AVENTIS MONOPROSOPI A.E.B.E	GR
Lasix	not available	41423/07/27-05-2008	SANOFI-AVENTIS MONOPROSOPI A.E.B.E	GR
Lasix 20 mg/ 2 ml oplossing voor injectie	not available	BE067401	SANOFI BELGIUM	BE

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Lasix 20 mg/ 2 ml oplossing voor injectie	not available	BE067401	SANOFI BELGIUM	BE
Lasix 20 mg/ 2 ml oplossing voor injectie	not available	BE067401	SANOFI BELGIUM	BE
Lasix 20 mg/ 2 ml solution injectable	not available	BE067401	SANOFI BELGIUM	BE
Lasix 20 mg/ 2 ml solution injectable	not available	BE067401	SANOFI BELGIUM	BE
Lasix 20 mg/ 2 ml solution injectable	not available	BE067401	SANOFI BELGIUM	BE
Lasix 20 mg/ 2 ml solution injectable	not available	0689075	SANOFI BELGIUM	LU
Lasix 20 mg/ 2 ml solution injectable	not available	0063935	SANOFI BELGIUM	LU
LASIX 10 mg/ml soluzione orale	not available	023993052	SANOFI S.R.L.	IT
Lasix 20 mg/2 ml Injektionslösung	not available	BE067401	SANOFI BELGIUM	BE
Lasix 20 mg/2 ml Injektionslösung	not available	BE067401	SANOFI BELGIUM	BE
Lasix 20 mg/2 ml Injektionslösung	not available	BE067401	SANOFI BELGIUM	BE
Lasix 20 mg/2 ml solução injetável	not available	8113704	SANOFI - PRODUTOS FARMACEUTICOS LDA	PT
Lasix 20mg/2ml Solution for Injection or Infusion	not available	PA 540/52/2	SANOFI-AVENTIS IRELAND LTD. T/A SANOFI	IE

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LASIX 25 mg compresse	not available	023993013	SANOFI S.R.L.	IT
Lasix 250 mg Konzentrat zur Herstellung einer Infusionslösung	not available	15.861	SANOFI-AVENTIS GMBH OSTERREICH	AT
Lasix 250 mg Konzentrat zur Herstellung einer Infusionslösung	not available	15.861	SANOFI-AVENTIS GMBH OSTERREICH	AT
Lasix 250 mg/ 25 ml concentraat voor oplossing voor infusie	not available	BE091506	SANOFI BELGIUM	BE
Lasix 250 mg/ 25 ml concentraat voor oplossing voor infusie	not available	BE091506	SANOFI BELGIUM	BE
Lasix 250 mg/25 ml Konzentrat zur Herstellung einer Infusionslösung	not available	BE091506	SANOFI BELGIUM	BE
Lasix 250 mg/25 ml Konzentrat zur Herstellung einer Infusionslösung	not available	BE091506	SANOFI BELGIUM	BE
Lasix 250 mg/25 ml solution à diluer pour perfusion.	not available	BE091506	SANOFI BELGIUM	BE
Lasix 250 mg/25 ml solution à diluer pour perfusion.	not available	BE091506	SANOFI BELGIUM	BE
Lasix 250 mg/25 ml solution à diluer pour perfusion.	not available	0743507	SANOFI BELGIUM	LU
Lasix 250 mg/25 ml solution à diluer pour perfusion.	not available	0063952	SANOFI BELGIUM	LU
LASIX 250 mg/25 ml soluzione per infusione	not available	023993049	SANOFI S.R.L.	IT
Lasix 30 mg Prolongatum capsules met verlengde afgifte, hard	not available	BE115367	SANOFI BELGIUM	BE

Product full name (in authorisation country as per Art. 57 data)	MRP/DCP or CP Authorisation number (if not available then insert not available)	National Authorisation Number of product in the member state	MAH of product in the member state	Member State where product is authorised
Lasix 30 mg Prolongatum gélules à libération prolongée	not available	BE115367	SANOFI BELGIUM	BE
Lasix 30 mg Prolongatum gélules à libération prolongée	not available	0064039	SANOFI BELGIUM	LU
Lasix 30 mg Prolongatum Hartkapseln, retardiert	not available	BE115367	SANOFI BELGIUM	BE
Lasix 40 mg comprimés	not available	BE067663	SANOFI BELGIUM	BE
Lasix 40 mg comprimés	not available	BE067663	SANOFI BELGIUM	BE
Lasix 40 mg comprimés	not available	BE067663	SANOFI BELGIUM	BE
Lasix 40 mg comprimés	not available	BE067663	SANOFI BELGIUM	BE
Lasix 40 mg comprimés	not available	0063899	SANOFI BELGIUM	LU
Lasix 40 mg comprimés	not available	0063904	SANOFI BELGIUM	LU
Lasix 40 mg comprimés	not available	0656544	SANOFI BELGIUM	LU
Lasix 40 mg comprimidos	not available	4597381	SANOFI - PRODUTOS FARMACEUTICOS LDA	PT
Lasix 40 mg comprimidos	not available	8113845	SANOFI - PRODUTOS FARMACEUTICOS LDA	PT
Lasix 40 mg comprimidos	not available	4597282	SANOFI - PRODUTOS FARMACEUTICOS LDA	PT

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Lasix 40 mg comprimidos	not available	8113837	SANOFI - PRODUTOS FARMACEUTICOS LDA	PT
Lasix 40 mg Injektionslösung	not available	35018.00.00	SANOFI-AVENTIS DEUTSCHLAND GMBH	DE
Lasix 40 mg Injektionslösung	not available	35018.00.00	SANOFI-AVENTIS DEUTSCHLAND GMBH	DE
Lasix 40 mg Injektionslösung	not available	35018.00.00	SANOFI-AVENTIS DEUTSCHLAND GMBH	DE
Lasix 40 mg tablete	not available	HR-H-797987648-01	SANOFI-AVENTIS GROUPE	HR
Lasix 40 mg tablete	not available	HR-H-797987648-03	SANOFI-AVENTIS GROUPE	HR
Lasix 40 mg tablete	not available	HR-H-797987648	SANOFI-AVENTIS GROUPE	HR
Lasix 40 mg tablete	not available	H/92/00877/001	SANOFI-AVENTIS GROUPE	SI
Lasix 40 mg Tabletten	not available	12.585	SANOFI-AVENTIS GMBH OSTERREICH	AT
Lasix 40 mg Tabletten	not available	12.585	SANOFI-AVENTIS GMBH OSTERREICH	AT
Lasix 40 mg tabletten	not available	BE067663	SANOFI BELGIUM	BE
Lasix 40 mg Tabletten	not available	BE067663	SANOFI BELGIUM	BE
Lasix 40 mg tabletten	not available	BE067663	SANOFI BELGIUM	BE

Product full name (in authorisation country as per Art. 57 data)	MRP/DCP or CP Authorisation number (if not available then insert not available)	National Authorisation Number of product in the member state	MAH of product in the member state	Member State where product is authorised
Lasix 40 mg tabletten	not available	BE067663	SANOFI BELGIUM	BE
Lasix 40 mg tabletten	not available	BE067663	SANOFI BELGIUM	BE
Lasix 40 mg Tabletten	not available	BE067663	SANOFI BELGIUM	BE
Lasix 40 mg Tabletten	not available	BE067663	SANOFI BELGIUM	BE
Lasix 40 mg Tabletten	not available	BE067663	SANOFI BELGIUM	BE
LASIX 500 mg compresse	not available	023993037	SANOFI S.R.L.	IT
Lasix 500 mg comprimés	not available	0063921	SANOFI BELGIUM	LU
Lasix 500 mg comprimés.	not available	BE091497	SANOFI BELGIUM	BE
Lasix 500 mg Tabletten	not available	15.860	SANOFI-AVENTIS GMBH OSTERREICH	AT
Lasix 500 mg tabletten	not available	BE091497	SANOFI BELGIUM	BE
Lasix 500 mg Tabletten	not available	BE091497	SANOFI BELGIUM	BE
Lasix 500, tabletten 500 mg	not available	RVG 15018	GENZYME EUROPE B.V.	NL
Lasix 500, tabletten 500 mg	not available	RVG 15018	GENZYME EUROPE B.V.	NL

Product full name (in authorisation country as per Art. 57 data)	MRP/DCP or CP Authorisation number (if not available then insert not available)	National Authorisation Number of product in the member state	MAH of product in the member state	Member State where product is authorised
Lasix 80 mg Tabletten	not available	17.821	SANOFI-AVENTIS GMBH OSTERREICH	AT
Lasix 80 mg Tabletten	not available	17.821	SANOFI-AVENTIS GMBH OSTERREICH	AT
Lasix 80 mg Tabletten	not available	17.821	SANOFI-AVENTIS GMBH OSTERREICH	AT
LASIX FIALE 20 mg/2 ml soluzione iniettabile	not available	020465011	SANOFI-AVENTIS DEUTSCHLAND GMBH	IT
Lasix liquidum 10 mg/ml Lösung zum Einnehmen	not available	12043.00.00	SANOFI-AVENTIS DEUTSCHLAND GMBH	DE
Lasix liquidum 10 mg/ml Lösung zum Einnehmen	not available	12043.00.00	SANOFI-AVENTIS DEUTSCHLAND GMBH	DE
LASIX RETARD	not available	6896	SANOFI-AVENTIS NORGE AS	NO
LASIX RETARD	not available	6896	SANOFI-AVENTIS NORGE AS	NO
Lasix Retard 30 mg depotkapsler	not available	6895	SANOFI-AVENTIS NORGE AS	NO
Lasix Retard 30 mg hård depotkapsel	not available	9803	SANOFI AB	SE
Lasix Retard 30 mg hård depotkapsel	not available	9803	SANOFI AB	SE
Lasix Retard 30 mg hård depotkapsel	not available	9803	SANOFI AB	SE
Lasix Retard 30 mg hård depotkapsel	not available	9803	SANOFI AB	SE

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Lasix Retard 30 mg hörð forðahylki.	not available	920254	SANOFI-AVENTIS NORGE AS	IS
Lasix retard 30 mg Kapseln	not available	17.113	SANOFI-AVENTIS GMBH OSTERREICH	AT
Lasix retard 30 mg Kapseln	not available	17.113	SANOFI-AVENTIS GMBH OSTERREICH	AT
Lasix Retard 60 mg cápsulas de libertação prolongada	not available	5829890	SANOFI - PRODUTOS FARMACEUTICOS LDA	PT
Lasix Retard 60 mg cápsulas de libertação prolongada	not available	9485102	SANOFI - PRODUTOS FARMACEUTICOS LDA	PT
Lasix Retard 60 mg hård depotkapsel	not available	9804	SANOFI AB	SE
Lasix Retard 60 mg hård depotkapsel	not available	9804	SANOFI AB	SE
Lasix Retard 60 mg hård depotkapsel	not available	9804	SANOFI AB	SE
Lasix Retard 60 mg hörð forðahylki	not available	920255	SANOFI-AVENTIS NORGE AS	IS
Lasix Retard 60 mg hörð forðahylki	not available	920255	SANOFI-AVENTIS NORGE AS	IS
Lasix retard 60 mg Kapseln	not available	1-22210	SANOFI-AVENTIS GMBH OSTERREICH	AT
Lasix retard 60 mg Kapseln	not available	1-22210	SANOFI-AVENTIS GMBH OSTERREICH	AT
Lasix retard 60 mg Kapseln	not available	1-22210	SANOFI-AVENTIS GMBH OSTERREICH	AT

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Lasix Retard 60 mg pailginto atpalaidavimo kietosios kapsulės	not available	LT/1/99/1059/002	SANOFI-AVENTIS GROUPE	LT
Lasix Retard 60 mg pailginto atpalaidavimo kietosios kapsulės	not available	LT/1/99/1059/001	SANOFI-AVENTIS GROUPE	LT
Lasix Retard 60, capsules met gereguleerde afgifte 60 mg	not available	RVG 10129	GENZYME EUROPE B.V.	NL
Lasix Retard, 60 mg toimeainet prolongeeritult vabastavad kõvakapslid	not available	327200	SANOFI-AVENTIS GROUPE	EE
Lasix Retard, depotkapsler	not available	09940	SANOFI A/S	DK
Lasix®	not available	19673	SANOFI WINTHROP INDUSTRIE	CY
Lasix® 20 mg/2 ml Ampullen	not available	12.584	SANOFI-AVENTIS GMBH OSTERREICH	AT
Lasix® 20 mg/2 ml Ampullen	not available	12.584	SANOFI-AVENTIS GMBH OSTERREICH	AT
Lasix® 20 mg/2 ml Ampullen	not available	12.584	SANOFI-AVENTIS GMBH OSTERREICH	AT
Lasix® 20 mg/2 ml Ampullen	not available	12.584	SANOFI-AVENTIS GMBH OSTERREICH	AT
Lasix® 40 mg/4 ml Ampullen	not available	17.822	SANOFI-AVENTIS GMBH OSTERREICH	AT
Seguril 20 mg/2 ml solución inyectable	not available	39968	SANOFI-AVENTIS, S.A.	ES
Seguril 250 mg/25 ml solución inyectable	not available	56508	SANOFI-AVENTIS, S.A.	ES

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Seguril 40 mg comprimidos	not available	39969	SANOFI-AVENTIS, S.A.	ES
Seguril 40 mg comprimidos	not available	39969	SANOFI-AVENTIS, S.A.	ES