



EUROPEAN MEDICINES AGENCY
SCIENCE MEDICINES HEALTH

10 November 2022
EMA/759336/2022
Human Medicines Division

List of nationally authorised medicinal products

Active substance(s): amoxicillin

Procedure No. PSUSA/00000187/202203



Product Name (in authorisation country)	MRP/DCP Authorisation number	National Authorisation Number	MAH of product in the member state	Member State where product is authorised
Ospamox 125 mg/5 ml – Pulver für orale Suspension	AT/H/0116/003	17.661	SANDOZ GMBH	AT
Amoxicillin 1A Pharma 500 mg – Filmtabletten	AT/H/0189/001	1-28793	1A PHARMA GMBH	AT
Amoxicillin 1A Pharma 750 mg – Filmtabletten	AT/H/0189/002	1-28794	1A PHARMA GMBH	AT
Amoxicillin 1A Pharma 1000 mg – Filmtabletten	AT/H/0189/003	1-28795	1A PHARMA GMBH	AT
Ospamox 750 mg - lösbare Tabletten	NL/H/0455/002	1-25541	SANDOZ GMBH	AT
Ospamox 500 mg - lösbare Tabletten	NL/H/0455/001	1-25540	SANDOZ GMBH	AT
Ospamox 500 mg/5 ml – Pulver für orale Suspension	AT/H/0116/005	1-24881	SANDOZ GMBH	AT
Amoxicillin Sandoz 750 mg – Filmtabletten	AT/H/0187/002	1-28901	SANDOZ GMBH	AT
Amoxicillin Sandoz 1000 mg – Filmtabletten	AT/H/0187/003	1-28902	SANDOZ GMBH	AT
Amoxicillin Sandoz 500 mg – Filmtabletten	AT/H/0187/001	1-28900	SANDOZ GMBH	AT
Ospamox 250 mg – lösbare Tabletten	not available	137432	SANDOZ GMBH	AT
Ospamox 500 mg – Filmtabletten	not available	17659	SANDOZ GMBH	AT
Ospamox 750 mg – Filmtabletten	not available	17657	SANDOZ GMBH	AT
Ospamox 1000 mg - Filmtabletten	not available	17653	SANDOZ GMBH	AT
Delamoxyle 1 g, poudre pour solution injectable/pour perfusion	FR/H/0780/002	BE109417	LABORATOIRES DELBERT	BE
Delamoxyle 1 g, Pulver zur Herstellung einer Injektions-/Infusionslösung	FR/H/0780/002	BE109417	LABORATOIRES DELBERT	BE
Delamoxyle 1 g, poeder voor oplossing voor injectie/infusie	FR/H/0780/002	BE109417	LABORATOIRES DELBERT	BE

Product Name (in authorisation country)	MRP/DCP Authorisation number	National Authorisation Number	MAH of product in the member state	Member State where product is authorised
Clamoxyl 250 mg/5 ml, poeder voor orale suspensie	FR/H/0600/003	BE080577	GLAXOSMITHKLINE PHARMACEUTICALS SA	BE
Clamoxyl 250 mg/5 ml, Pulver zur Herstellung einer Suspension zum Einnehmen	FR/H/0600/003	BE080577	GLAXOSMITHKLINE PHARMACEUTICALS SA	BE
Clamoxyl 125 mg/5 ml, poeder voor orale suspensie	FR/H/0600/002	BE080595	GLAXOSMITHKLINE PHARMACEUTICALS SA	BE
Clamoxyl 125 mg/5 ml, Pulver zur Herstellung einer Suspension zum Einnehmen	FR/H/0600/002	BE080595	GLAXOSMITHKLINE PHARMACEUTICALS SA	BE
Clamoxyl 1 g, disperseerbare tabletten	FR/H/0600/001	BE164306	GLAXOSMITHKLINE PHARMACEUTICALS SA	BE
Clamoxyl 1 g, Tabletten zur Herstellung einer Suspension zum Einnehmen	FR/H/0600/001	BE164306	GLAXOSMITHKLINE PHARMACEUTICALS SA	BE
Clamoxyl 250 mg/5 ml, poudre pour suspension buvable	FR/H/0600/003	BE080577	GLAXOSMITHKLINE PHARMACEUTICALS SA	BE
Clamoxyl 125 mg/5 ml, poudre pour suspension buvable	FR/H/0600/002	BE080595	GLAXOSMITHKLINE PHARMACEUTICALS SA	BE
Clamoxyl 1 g, comprimés dispersibles	FR/H/0600/001	BE164306	GLAXOSMITHKLINE PHARMACEUTICALS SA	BE
Clamoxyl 500 mg, harde capsules	FR/H/0649/001	BE191807	GLAXOSMITHKLINE PHARMACEUTICALS SA	BE
Clamoxyl 500 mg Hartkapseln	FR/H/0649/001	BE191807	GLAXOSMITHKLINE PHARMACEUTICALS SA	BE
Clamoxyl 500 mg, gélules	FR/H/0649/001	BE191807	GLAXOSMITHKLINE PHARMACEUTICALS SA	BE
Amoxil Forte 250 mg/5 ml κόνις για πόσιμο εναιώρημα	FR/H/600/03	3617	GLAXOSMITHKLINE (IRELAND) LIMITED	CY
Amoxil 500 mg καψάκια, σκληρά	FR/H/0649/001	19966	GLAXOSMITHKLINE (IRELAND) LIMITED	CY

Product Name (in authorisation country)	MRP/DCP Authorisation number	National Authorisation Number	MAH of product in the member state	Member State where product is authorised
Amoxicillin Micro Labs 500 mg Filmtabletten	DE/H/4385/002	94793.00.00	MICRO LABS GMBH	DE
Amoxicillin Micro Labs 500 mg Filmtabletten	DE/H/4385/002	94793.00.00	MICRO LABS GMBH	DE
Amoxicillin Micro Labs 500 mg Filmtabletten	DE/H/4385/002	94793.00.00	MICRO LABS GMBH	DE
Amoxicillin Micro Labs 500 mg Filmtabletten	DE/H/4385/002	94793.00.00	MICRO LABS GMBH	DE
Amoxicillin Micro Labs 500 mg Filmtabletten	DE/H/4385/002	94793.00.00	MICRO LABS GMBH	DE
Amoxicillin Micro Labs 500 mg Filmtabletten	DE/H/4385/002	94793.00.00	MICRO LABS GMBH	DE
Amoxicillin Micro Labs 500 mg Filmtabletten	DE/H/4385/002	94793.00.00	MICRO LABS GMBH	DE
Amoxicillin Micro Labs 500 mg Filmtabletten	DE/H/4385/002	94793.00.00	MICRO LABS GMBH	DE
Amoxicillin Micro Labs 500 mg Filmtabletten	DE/H/4385/002	94793.00.00	MICRO LABS GMBH	DE
Amoxicillin Micro Labs 500 mg Filmtabletten	DE/H/4385/002	94793.00.00	MICRO LABS GMBH	DE
Amoxicillin Micro Labs 500 mg Filmtabletten	DE/H/4385/002	94793.00.00	MICRO LABS GMBH	DE
Amoxicillin Micro Labs 500 mg Filmtabletten	DE/H/4385/002	94793.00.00	MICRO LABS GMBH	DE
Amoxicillin Micro Labs 500 mg Filmtabletten	DE/H/4385/002	94793.00.00	MICRO LABS GMBH	DE
Amoxicillin Micro Labs 500 mg Filmtabletten	DE/H/4385/002	94793.00.00	MICRO LABS GMBH	DE
Amoxicillin Micro Labs 500 mg Filmtabletten	DE/H/4385/002	94793.00.00	MICRO LABS GMBH	DE
Amoxicillin Micro Labs 500 mg Filmtabletten	DE/H/4385/002	94793.00.00	MICRO LABS GMBH	DE
Amoxicillin Micro Labs 500 mg Filmtabletten	DE/H/4385/002	94793.00.00	MICRO LABS GMBH	DE
Amoxicillin Micro Labs 500 mg Filmtabletten	DE/H/4385/002	94793.00.00	MICRO LABS GMBH	DE
Amoxicillin Micro Labs 500 mg Filmtabletten	DE/H/4385/002	94793.00.00	MICRO LABS GMBH	DE
Amoxicillin Micro Labs 500 mg Filmtabletten	DE/H/4385/002	94793.00.00	MICRO LABS GMBH	DE
Amoxicillin Micro Labs 500 mg Filmtabletten	DE/H/4385/002	94793.00.00	MICRO LABS GMBH	DE

Product Name (in authorisation country)	MRP/DCP Authorisation number	National Authorisation Number	MAH of product in the member state	Member State where product is authorised
Amoxicillin Micro Labs 750 mg Filmtabletten	DE/H/4385/003	94794.00.00	MICRO LABS GMBH	DE
Amoxicillin Micro Labs 750 mg Filmtabletten	DE/H/4385/003	94794.00.00	MICRO LABS GMBH	DE
Amoxicillin Micro Labs 750 mg Filmtabletten	DE/H/4385/003	94794.00.00	MICRO LABS GMBH	DE
Amoxicillin Micro Labs 750 mg Filmtabletten	DE/H/4385/003	94794.00.00	MICRO LABS GMBH	DE
Amoxicillin Micro Labs 750 mg Filmtabletten	DE/H/4385/003	94794.00.00	MICRO LABS GMBH	DE
Amoxicillin Micro Labs 750 mg Filmtabletten	DE/H/4385/003	94794.00.00	MICRO LABS GMBH	DE
Amoxicillin Micro Labs 750 mg Filmtabletten	DE/H/4385/003	94794.00.00	MICRO LABS GMBH	DE
Amoxicillin Micro Labs 750 mg Filmtabletten	DE/H/4385/003	94794.00.00	MICRO LABS GMBH	DE
Amoxicillin Micro Labs 750 mg Filmtabletten	DE/H/4385/003	94794.00.00	MICRO LABS GMBH	DE
Amoxicillin Micro Labs 750 mg Filmtabletten	DE/H/4385/003	94794.00.00	MICRO LABS GMBH	DE
Amoxicillin Micro Labs 750 mg Filmtabletten	DE/H/4385/003	94794.00.00	MICRO LABS GMBH	DE
Amoxicillin Micro Labs 750 mg Filmtabletten	DE/H/4385/003	94794.00.00	MICRO LABS GMBH	DE
Amoxicillin Micro Labs 750 mg Filmtabletten	DE/H/4385/003	94794.00.00	MICRO LABS GMBH	DE
Amoxicillin Micro Labs 750 mg Filmtabletten	DE/H/4385/003	94794.00.00	MICRO LABS GMBH	DE
Amoxicillin Micro Labs 750 mg Filmtabletten	DE/H/4385/003	94794.00.00	MICRO LABS GMBH	DE
Amoxicillin Micro Labs 750 mg Filmtabletten	DE/H/4385/003	94794.00.00	MICRO LABS GMBH	DE
Amoxicillin Micro Labs 750 mg Filmtabletten	DE/H/4385/003	94794.00.00	MICRO LABS GMBH	DE
Amoxicillin Micro Labs 750 mg Filmtabletten	DE/H/4385/003	94794.00.00	MICRO LABS GMBH	DE
Amoxicillin Micro Labs 750 mg Filmtabletten	DE/H/4385/003	94794.00.00	MICRO LABS GMBH	DE
Amoxicillin Micro Labs 750 mg Filmtabletten	DE/H/4385/003	94794.00.00	MICRO LABS GMBH	DE
Amoxicillin Micro Labs 750 mg Filmtabletten	DE/H/4385/003	94794.00.00	MICRO LABS GMBH	DE

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Imacillin, granulat til oral suspension	not available	11372	VIATRIS APS	DK
Imacillin, opløselige tabletter	not available	15288	VIATRIS APS	DK
Imacillin, opløselige tabletter	not available	15289	VIATRIS APS	DK
Imacillin, opløselige tabletter	not available	15290	VIATRIS APS	DK
Imacillin, opløselige tabletter	not available	15291	VIATRIS APS	DK
Imacillin, opløselige tabletter	not available	16713	VIATRIS APS	DK
Clamoxyl 1 g comprimidos dispersables	FR/H/0600/001	59.133	GLAXOSMITHKLINE S.A.	ES
Clamoxyl 500 mg cápsulas duras	FR/H/0649/001	50.239	GLAXOSMITHKLINE S.A.	ES
AMOXICILLINE ZENTIVA LAB 500 mg gélule	not available	NL43719	LABORATOIRE GLAXOSMITHKLINE	FR
AMOXICILLINE ZENTIVA LAB 1 g, comprimé dispersible	not available	NL43720	LABORATOIRE GLAXOSMITHKLINE	FR
XYLLOMAC 500 mg, poudre pour solution injectable/pour perfusion (IM-IV)	FR/H/0780/001	3400932252480	LABORATOIRES DELBERT	FR
XYLLOMAC 500 mg, poudre pour solution injectable/pour perfusion (IM-IV)	FR/H/0780/001	3400955648710	LABORATOIRES DELBERT	FR
XYLLOMAC 1 g, poudre pour solution injectable/pour perfusion (I.M.-I.V.)	FR/H/0780/002	34009 300 240 9 6	LABORATOIRES DELBERT	FR
XYLLOMAC 1 g, poudre pour solution injectable/pour perfusion (I.M.-I.V.)	FR/H/0780/002	34009 322 526 0 2	LABORATOIRES DELBERT	FR
XYLLOMAC 1 g, poudre	FR/H/0780/002	34009 552 561 2 0	LABORATOIRES DELBERT	FR

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pour solution injectable/pour perfusion (I.M.-I.V.)				
XYLLOMAC 1 g, poudre pour solution injectable/pour perfusion (I.M.-I.V.)	FR/H/0780/002	34009 555 775 3 9	LABORATOIRES DELBERT	FR
AMOXICILLINE ZENTIVA LAB 125 mg/5 ml, poudre pour suspension buvable	not available	NL43721	LABORATOIRE GLAXOSMITHKLINE	FR
AMOXICILLINE ZENTIVA LAB 250 mg/5 ml, poudre pour suspension buvable	not available	NL43722	LABORATOIRE GLAXOSMITHKLINE	FR
AMOXICILLINE ZENTIVA LAB 500 mg/5 ml, poudre pour suspension buvable	not available	NL43723	LABORATOIRE GLAXOSMITHKLINE	FR
XYLLOMAC 2 g, poudre pour solution injectable/pour perfusion (IV)	FR/H/0780/003	3400932252770	LABORATOIRES DELBERT	FR
XYLLOMAC 2 g, poudre pour solution injectable/pour perfusion (IV)	FR/H/0780/003	3400955249955	LABORATOIRES DELBERT	FR
CLAMOXYL 1 g, poudre et solvant pour solution injectable (IM)	FR/H/0600/006	VNL11851	LABORATOIRE GLAXOSMITHKLINE	FR
CLAMOXYL 125 mg/5 ml, poudre pour suspension buvable	FR/H/0600/002	VNL11074	LABORATOIRE GLAXOSMITHKLINE	FR
CLAMOXYL 250 mg/5 ml, poudre pour suspension buvable	FR/H/0600/003	VNL11075	LABORATOIRE GLAXOSMITHKLINE	FR
CLAMOXYL 1 g, comprimé dispersible	FR/H/0600/001	NL15119	LABORATOIRE GLAXOSMITHKLINE	FR
CLAMOXYL 500 mg/5 ml, poudre pour suspension buvable	FR/H/0600/004	VNL11666	LABORATOIRE GLAXOSMITHKLINE	FR

Product Name (in authorisation country)	MRP/DCP Authorisation number	National Authorisation Number	MAH of product in the member state	Member State where product is authorised
AMOXICILLINE BIOGARAN 1 g, comprimé dispersible	not available	NL19733	LABORATOIRE GLAXOSMITHKLINE	FR
AMOXICILLINE BIOGARAN 500 mg/5 ml, poudre pour suspension buvable	not available	NL19721	LABORATOIRE GLAXOSMITHKLINE	FR
AMOXICILLINE BIOGARAN 250 mg/5 ml, poudre pour suspension buvable	not available	NL19720	LABORATOIRE GLAXOSMITHKLINE	FR
AMOXICILLINE BIOGARAN 125 mg/5 ml, poudre pour suspension buvable	not available	NL19719	LABORATOIRE GLAXOSMITHKLINE	FR
CLAMOXYL 500 mg, gélule	FR/H/0649/001	VNL9721	LABORATOIRE GLAXOSMITHKLINE	FR
AMOXICILLINE SANDOZ 500 mg, gélule	not available	34009 367 960 1 0	SANDOZ	FR
AMOXICILLINE SANDOZ 250 mg/5 ml, poudre pour suspension buvable	not available	34009 365 824 3 9	SANDOZ	FR
AMOXICILLINE SANDOZ 125 mg/5 ml, poudre pour suspension buvable	not available	34009 368 829 6 6	SANDOZ	FR
Amoxil 1 g κόνις για ενέσιμο διάλυμα ή διάλυμα προς έγχυση	BE/H/0259/001	0933607	GLAXOSMITHKLINE SINGLE MEMBER A.E.B.E.	GR
Amoxil 500 mg/5 ml κόνις για πόσιμο εναιώρημα	FR/H/600/04	0933603	GLAXOSMITHKLINE SINGLE MEMBER A.E.B.E.	GR
Amoxil 1 g διασπειρόμενα δισκία	FR/H/0600/001	0933610	GLAXOSMITHKLINE SINGLE MEMBER A.E.B.E.	GR
Amoxil 250 mg/5 ml κόνις για πόσιμο εναιώρημα	FR/H/600/03	0933602	GLAXOSMITHKLINE SINGLE MEMBER A.E.B.E.	GR
Amoxil 500 mg καψάκια, σκληρά	FR/H/0649/001	0933605	GLAXOSMITHKLINE SINGLE MEMBER A.E.B.E.	GR
Amoxicillin Delbert 500 mg, powder for solution for injection or infusion	FR/H/0780/001	PA23217/001/001	LABORATOIRES DELBERT	IE
Amoxil Vials 500mg, powder for solution for injection or infusion	FR/H/0649/002	PA 1077/33/4	GLAXOSMITHKLINE (IRELAND) LIMITED	IE

Product Name (in authorisation country)	MRP/DCP Authorisation number	National Authorisation Number	MAH of product in the member state	Member State where product is authorised
Amoxicillin 250 mg hard capsules	not available	PA0126/282/001	CLONMEL HEALTHCARE LTD.	IE
Amoxicillin 125mg / 5ml powder for oral suspension	not available	PA0126/282/003	CLONMEL HEALTHCARE LTD.	IE
Amoxicillin 250mg / 5ml powder for oral suspension	not available	PA0126/282/004	CLONMEL HEALTHCARE LTD.	IE
Amoxicillin 500 mg hard capsules	not available	PA0126/282/002	CLONMEL HEALTHCARE LTD.	IE
Amoxicillin Sandoz 100 mg/ml mixtúruđuft, dreifa	AT/H/0116/005	IS/1/15/062/01	SANDOZ A/S	IS
ZIMOX 1 g compresse solubili e masticabili	not available	023086174	PFIZER ITALIA S.R.L.	IT
ZIMOX 500 mg capsule rigide	not available	023086061	PFIZER ITALIA S.R.L.	IT
ZIMOX 500 mg compresse solubili e masticabili	not available	023086162	PFIZER ITALIA S.R.L.	IT
ZIMOX 1 g compresse	not available	023086150	PFIZER ITALIA S.R.L.	IT
ZIMOX 250 mg/5 ml polvere per sospensione orale	not available	023086097	PFIZER ITALIA S.R.L.	IT
ZIMOX 100 mg/ml gocce orali, sospensione	not available	023086186	PFIZER ITALIA S.R.L.	IT
Velamox 250 mg/7 ml polvere per sospensione orale.	not available	023097037	NEOPHARMED GENTILI SPA	IT
Velamox 1 g compresse dispersibili.	not available	023097102	NEOPHARMED GENTILI SPA	IT
Velamox 500 mg capsule rigide.	not available	023097013	NEOPHARMED GENTILI SPA	IT
SINTOPEN " 1 G COMPRESSE"	not available	023053123	MAGIS FARMACEUTICI SRL	IT
Amoxina 1 g compresse dispersibili	not available	023966094	AESCLAPIUS FARMACEUTICI S.R.L.	IT
Amoxina 250mg/5ml polvere per sospensione orale	not available	023966082	AESCLAPIUS FARMACEUTICI S.R.L.	IT
SINTOPEN " 250 MG/5 ML POLVERE PER	not available	023053135	MAGIS FARMACEUTICI SRL	IT

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SOSPENSIONE ORALE"				
Amoxina 250mg/5ml polvere per sospensione orale	not available	023966106	AESCUAPIUS FARMACEUTICI S.R.L.	IT
Delamoxyle 1000 mg, poudre pour solution injectable/pour perfusion	FR/H/0780/002	2008100038	LABORATOIRES DELBERT	LU
Delamoxyle 1000 mg, Pulver zur Herstellung einer Injektions-/Infusionslösung	FR/H/0780/002	2008100038	LABORATOIRES DELBERT	LU
Clamoxyl 1 g, Tabletten zur Herstellung einer Suspension zum Einnehmen	FR/H/0600/001	0260/08 10 0030	GLAXOSMITHKLINE PHARMACEUTICALS SA	LU
Clamoxyl 125 mg/5 ml, Pulver zur Herstellung einer Suspension zum Einnehmen	FR/H/0600/002	2008 10 0032	GLAXOSMITHKLINE PHARMACEUTICALS SA	LU
Clamoxyl 250 mg/5 ml, Pulver zur Herstellung einer Suspension zum Einnehmen	FR/H/0600/003	2008 10 0033	GLAXOSMITHKLINE PHARMACEUTICALS SA	LU
Clamoxyl 250 mg/5 ml, poudre pour suspension buvable	FR/H/600/03/MR	2008 10 0033	GLAXOSMITHKLINE PHARMACEUTICALS SA	LU
Clamoxyl 1 g, comprimés dispersibles	FR/H/600/01/MR	0260/08 10 0030	GLAXOSMITHKLINE PHARMACEUTICALS SA	LU
Clamoxyl 500 mg Hartkapseln	FR/H/0649/001	260/08 10 0031	GLAXOSMITHKLINE PHARMACEUTICALS SA	LU
Clamoxyl 500 mg, gélules	FR/H/0649/001	0260/08 10 0031	GLAXOSMITHKLINE PHARMACEUTICALS SA	LU
Amoxil 500 mg cietas kapsulas	FR/H/0649/001	04-0196	GLAXOSMITHKLINE TRADING SERVICES LIMITED	LV
Ospamox 1000 mg apvalkotās tabletes	AT/H/0187/003	10-0081	SANDOZ PHARMACEUTICALS D.D.	LV
Amoxil 250 mg/5 ml powder for oral suspension	FR/H/0600/003	MA192/03201	GLAXOSMITHKLINE (IRELAND) LIMITED	MT

Product Name (in authorisation country)	MRP/DCP Authorisation number	National Authorisation Number	MAH of product in the member state	Member State where product is authorised
Amoxil 500 mg capsules, hard	FR/H/0649/001	MA192/03101	GLAXOSMITHKLINE (IRELAND) LIMITED	MT
Amoxicilline RIA 500 mg, capsules	not available	RVG 55123	RIA GENERICS LIMITED	NL
Amoxicilline dispers RIA 500 mg, dispergeerbare tabletten	not available	RVG 56827	RIA GENERICS LIMITED	NL
Amoxicilline dispers RIA 750 mg, dispergeerbare tabletten	not available	RVG 56828	RIA GENERICS LIMITED	NL
Ospamox 500 mg, 500 mg, tabletki powlekane	AT/H/0188/001	16801	SANDOZ GMBH	PL
OSPAMOX 750 MG, 750 MG, TABLETKI POWLEKANE	AT/H/0188/002	16802	SANDOZ GMBH	PL
OSPAMOX 1000 MG, 1000 MG, TABLETKI POWLEKANE	AT/H/0188/003	16687	SANDOZ GMBH	PL
Clamoxyl 250 mg/5 ml pó para suspensão oral	FR/H/600/03	8352203	BEECHAM PORTUGUESA, PRODUTOS FARMACÊUTICOS E QUÍMICOS, LDA	PT
Clamoxyl 500 mg/5 ml pó para suspensão oral	FR/H/600/04	8352211	BEECHAM PORTUGUESA, PRODUTOS FARMACÊUTICOS E QUÍMICOS, LDA	PT
Clamoxyl 1 g comprimidos dispersíveis	FR/H/0600/001	4667580	BEECHAM PORTUGUESA, PRODUTOS FARMACÊUTICOS E QUÍMICOS, LDA	PT
Clamoxyl 500 mg cápsulas	FR/H/0649/001	4667481	BEECHAM PORTUGUESA, PRODUTOS FARMACÊUTICOS E QUÍMICOS, LDA	PT
Clamoxyl 500 mg cápsulas	FR/H/0649/001	8352112	BEECHAM PORTUGUESA, PRODUTOS FARMACÊUTICOS E QUÍMICOS, LDA	PT
Ospamox 125 mg/5 ml Pó para suspensão oral	AT/H/0116/003	5022082	SANDOZ FARMACÊUTICA LDA.	PT
AMOXICILINA SANDOZ 1000 mg comprimate filmate	AT/H/0187/003	5475/2013/08	S.C. SANDOZ S.R.L.	RO
AMOXICILINA SANDOZ 500 mg comprimate filmate	AT/H/0187/001	5474/2013/01	S.C. SANDOZ S.R.L.	RO

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AMOXICILINA SANDOZ 500 mg comprimate filmate	AT/H/0187/001	5474/2013/02	S.C. SANDOZ S.R.L.	RO
AMOXICILINA SANDOZ 500 mg comprimate filmate	AT/H/0187/001	5474/2013/04	S.C. SANDOZ S.R.L.	RO
Amoxicilină Sandoz 500 mg comprimate filmate	AT/H/0187/001	5474/2013/03	S.C. SANDOZ S.R.L.	RO
AMOXICILINA SANDOZ 500 mg comprimate filmate	AT/H/0187/001	5474/2013/05	S.C. SANDOZ S.R.L.	RO
AMOXICILINA SANDOZ 500 mg comprimate filmate	AT/H/0187/001	5474/2013/06	S.C. SANDOZ S.R.L.	RO
AMOXICILINA SANDOZ 500 mg comprimate filmate	AT/H/0187/001	5474/2013/07	S.C. SANDOZ S.R.L.	RO
AMOXICILINA SANDOZ 500 mg comprimate filmate	AT/H/0187/001	5474/2013/09	S.C. SANDOZ S.R.L.	RO
AMOXICILINA SANDOZ 500 mg comprimate filmate	AT/H/0187/001	5474/2013/08	S.C. SANDOZ S.R.L.	RO
AMOXICILINA SANDOZ 500 mg comprimate filmate	AT/H/0187/001	5474/2013/10	S.C. SANDOZ S.R.L.	RO
AMOXICILINA SANDOZ 1000 mg comprimate filmate	AT/H/0187/003	5475/2013/01	S.C. SANDOZ S.R.L.	RO
Amoxicilină Sandoz 1000 mg comprimate filmate	AT/H/0187/003	5475/2013/02	S.C. SANDOZ S.R.L.	RO
Amoxicilină Sandoz 1000 mg comprimate filmate	AT/H/0187/003	5475/2013/04	S.C. SANDOZ S.R.L.	RO
AMOXICILINA SANDOZ 1000 mg comprimate filmate	AT/H/0187/003	5475/2013/03	S.C. SANDOZ S.R.L.	RO
Amoxicilină Sandoz 1000 mg comprimate filmate	AT/H/0187/003	5475/2013/05	S.C. SANDOZ S.R.L.	RO
Amoxicilină Sandoz 1000 mg comprimate filmate	AT/H/0187/003	5475/2013/06	S.C. SANDOZ S.R.L.	RO
AMOXICILINA SANDOZ 1000 mg comprimate filmate	AT/H/0187/003	5475/2013/07	S.C. SANDOZ S.R.L.	RO
Amoxicilină Sandoz 1000 mg comprimate filmate	AT/H/0187/003	5475/2013/10	S.C. SANDOZ S.R.L.	RO
AMOXICILINA SANDOZ	AT/H/0187/003	5475/2013/09	S.C. SANDOZ S.R.L.	RO

Product Name (in authorisation country)	MRP/DCP Authorisation number	National Authorisation Number	MAH of product in the member state	Member State where product is authorised
1000 mg comprimate filmate				
Amimox 375 mg filmdragerade tabletter	not available	11538	MEDA AB	SE
Amimox 750 mg filmdragerade tabletter	not available	11539	MEDA AB	SE
Amimox 100 mg/ml granulat till oral suspension	not available	11837	MEDA AB	SE
Amimox 500 mg filmdragerade tabletter	not available	11647	MEDA AB	SE
Amimox 50 mg/ml granulat till oral suspension	not available	11541	MEDA AB	SE
Imacillin 1 g tabletter	not available	11802	VIATRIS AB	SE
Amimox 125 mg granulat till oral suspension, dospåse	not available	11540	MEDA AB	SE
Duomox 1 000	not available	15/0383/95-S	ASTELLAS PHARMA S.R.O.	SK
Duomox 125	not available	15/0149/19-S	ASTELLAS PHARMA S.R.O.	SK
Duomox 250	not available	15/0150/19-S	ASTELLAS PHARMA S.R.O.	SK
Duomox 375	not available	15/0151/19-S	ASTELLAS PHARMA S.R.O.	SK
Duomox 750	not available	15/0153/19-S	ASTELLAS PHARMA S.R.O.	SK
Duomox 500	not available	15/0152/19-S	ASTELLAS PHARMA S.R.O.	SK
Amoxicillin 500mg Capsules	not available	PL 20416/0574	CRESCENT PHARMA LIMITED	XI
Amoxicillin 250mg Capsules	not available	PL 20416/0573	CRESCENT PHARMA LIMITED	XI
Amoxil Capsules 500 mg	FR/H/0649/001	PL 00038/0105	BEECHAM GROUP PLC	XI
Amoxicillin 125mg/5ml Suspension	AT/H/0116/003	PL 04416/0484	SANDOZ LTD	XI
Amoxicillin 500 mg hard capsules	AT/H/0116/002	PL 04416/0577	SANDOZ LTD	XI
Amoxicillin 250mg hard capsules	AT/H/0116/001	PL 04416/0576	SANDOZ LTD	XI
Amoxicillin 250mg/5ml Suspension	AT/H/0116/004	PL 04416/0485	SANDOZ LTD	XI
Amoxicillin 250mg Capsules	not available	PL 36722/0107	SPECIAL CONCEPT DEVELOPMENT (UK) LTD	XI

Product Name (in authorisation country)	MRP/DCP Authorisation number	National Authorisation Number	MAH of product in the member state	Member State where product is authorised
Amoxicillin 500mg Capsules	not available	PL 36722/0108	SPECIAL CONCEPT DEVELOPMENT (UK) LTD	XI
Amoxicillin 250mg Capsules	not available	PL 36722/0107	SPECIAL CONCEPT DEVELOPMENT (UK) LTD	XI
Amoxicillin 500mg Capsules	not available	PL 36722/0108	SPECIAL CONCEPT DEVELOPMENT (UK) LTD	XI
Amoxicillin 250mg Capsules	not available	PL 36722/0107	SPECIAL CONCEPT DEVELOPMENT (UK) LTD	XI
Amoxicillin 500mg Capsules	not available	PL 36722/0108	SPECIAL CONCEPT DEVELOPMENT (UK) LTD	XI
Amoxicillin 250 mg Capsules, hard	not available	PL 20117/0214	MORNINGSIDE HEALTHCARE LTD	XI
Amoxicillin 500 mg Capsules, hard	not available	PL 20117/0215	MORNINGSIDE HEALTHCARE LTD	XI
Amoxicillin 250 mg Capsules	not available	PL 40496/0041	BRILLPHARMA LIMITED	XI
Amoxicillin 500 mg Capsules	not available	PL 40496/0042	BRILLPHARMA LIMITED	XI