



EUROPEAN MEDICINES AGENCY
SCIENCE MEDICINES HEALTH

11 January 2024
EMA/504337/2023
Human Medicines Division

List of nationally authorised medicinal products

Active substance(s): azithromycin (systemic use formulations)

Procedure No. PSUSA/00010491/202304



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
Zithromax® 500 mg - Filmtabletten	not available	1-21939	PFIZER CORPORATION AUSTRIA GES.M.B.H.	AT
Zithromax® 200 mg/5 ml - Trockensaft	not available	1-20313	PFIZER CORPORATION AUSTRIA GESELLSCHAFT M.B.H.	AT
Zithromax® i.v. 500 mg - Pulver zur Herstellung einer Infusionslösung	not available	1-24636	PFIZER CORPORATION AUSTRIA GES.M.B.H.	AT
Azithromycin Sandoz 500 mg – Filmtabletten	NL/H/4670/002	1-26352	SANDOZ GMBH	AT
ZITROMAX 500 mg Filmomhulde tabletten	not available	BE193243	PFIZER S.A.	BE
ZITROMAX 600 mg Filmomhulde tabletten	not available	BE193252	PFIZER S.A.	BE
ZITROMAX 250 mg Filmomhulde tabletten	not available	BE193261	PFIZER S.A.	BE
ZITROMAX 200 mg/5 ml Poeder voor orale suspensie	not available	BE165961	PFIZER S.A.	BE
Zitromax 500 mg Pulver für ein Konzentrat zur Herstellung einer Infusionslösung	not available	BE562017	PFIZER S.A.	BE
СУМАМЕД сироп 100 mg/5ml прах за перорална суспензия	not available	20000867	TEVA B.V	BG
СУМАМЕД 500 mg филмирани таблетки	not available	20010175	TEVA B.V	BG
СУМАМЕД 500 mg прах за инфузионен разтвор	not available	20010799	TEVA B.V	BG
СУМАМЕД 250 mg твърди капсули	not available	20000866	TEVA B.V	BG
Сумамед форте сироп 200 mg/5 ml прах за перорална суспензия	not available	20010176	TEVA B.V	BG

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
ZITHROMAX, 500mg/φιαλίδιο, κόνις για διάλυμα προς έγχυση.	not available	19592	PFIZER HELLAS A.E.	CY
ZITHROMAX 200 mg/5 ml κόνις για πόσιμο εναιώρημα	not available	16336	PFIZER HELLAS A.E.	CY
SUMAMED tvrdé tobolky	not available	15/226/90-C	TEVA PHARMACEUTICALS CR, S.R.O.	CZ
SUMAMED forte sirup prášek pro perorální suspenzi	not available	15/352/92-B/C	TEVA PHARMACEUTICALS CR, S.R.O.	CZ
SUMAMED sirup prášek pro perorální suspenzi	not available	15/352/92-A/C	TEVA PHARMACEUTICALS CR, S.R.O.	CZ
SUMAMED 500 mg infuze prášek pro infuzní roztok	not available	15-223-03-C	TEVA PHARMACEUTICALS CR, S.R.O.	CZ
SUMAMED 500 mg potahované tablety	not available	15/351/92-B/C	TEVA PHARMACEUTICALS CR, S.R.O.	CZ
SUMAMED 125 mg potahované tablety	not available	15/351/92-A/C	TEVA PHARMACEUTICALS CR, S.R.O.	CZ
Azithromycin Aurovitas 500 mg potahované tablety	PT/H/1475/002	15/482/16-C	AUROVITAS, SPOL. S R.O.	CZ
Zithromax® 500 mg Filmdabletten	not available	37174.01.00	PFIZER PHARMA PFE GMBH	DE
Zithromax® Trockensaft 200 mg/5 ml Pulver zur Herstellung einer Suspension zum Einnehmen	not available	25156.00.01	PFIZER PHARMA PFE GMBH	DE
Zithromax® 250 mg Filmdabletten	not available	37174.00.00	PFIZER PHARMA PFE GMBH	DE
Ultreon® 600 mg Filmdabletten	not available	37344.00.00	PFIZER PHARMA PFE GMBH	DE
Zitromax, pulver til koncentrat til infusionsvæske,	not available	32123	PFIZER APS	DK

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opløsning				
Zitromax, pulver til oral suspension	not available	14083	PFIZER APS	DK
Zitromax, filmovertrukne tabletter	not available	19118	PFIZER APS	DK
Zitromax, filmovertrukne tabletter	not available	19117	PFIZER APS	DK
SUMAMED, 250 mg kõvakapslid	not available	292599	TEVA PHARMA B.V.	EE
SUMAMED, 500 mg õhukese polümeerikattega tabletid	not available	292499	TEVA PHARMA B.V.	EE
SUMAMED FORTE, 200 mg/5 ml suukaudse suspensiooni pulber	not available	357201	TEVA PHARMA B.V.	EE
Azitromicina Teva 250 mg comprimidos recubiertos con película EFG.	NL/H/0614/001	67335	TEVA PHARMA S.L.U.,	ES
Azitromicina Teva 250 mg comprimidos dispersables	PL/H/0423/001	82363	TEVA PHARMA S.L.U.,	ES
Zitromax 250 mg cápsulas duras	not available	59.616	PFIZER, S.L.	ES
Zitromax 200 mg/5 ml polvo para suspensión oral en frasco	not available	59.615	PFIZER, S.L.	ES
Zitromax 1000 mg polvo para suspensión oral en sobre	not available	60.065	PFIZER, S.L.	ES
Zitromax 500 mg polvo para suspensión oral en sobre	not available	60.066	PFIZER, S.L.	ES
Zitromax 250 mg polvo para suspensión oral en sobre	not available	59.620	PFIZER, S.L.	ES
Zitromax 500 mg	not available	61.272	PFIZER, S.L.	ES

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comprimidos recubiertos con película				
Azitromicina ratiopharm 250 mg comprimidos dispersables	PL/H/0424/001	82365	TEVA PHARMA S.L.U.,	ES
Zitromax 500 mg polvo para solución para perfusión.	not available	64.834	PFIZER, S.L.	ES
ZITHROMAX 40 mg/ml pulver till oral suspension	not available	11615	PFIZER OY	FI
ZITHROMAX 250 mg tablett, filmdragerad	not available	12016	PFIZER OY	FI
ZITHROMAX 500 mg tablett, filmdragerad	not available	12017	PFIZER OY	FI
ZITHROMAX IV 500 mg pulver till infusionsvätska, lösning	not available	16152	PFIZER OY	FI
AZITHROMYCINE EG 250 MG, COMPRIME PELLICULE	not available	NL31030	EG LABO LABORATOIRES EUROGENERICS - DO NOT USE	FR
ORDIPHA 500 mg, comprimé dispersible sécable	not available	34009 395 347 9 4	TONIPHARM	FR
AZITHROMYCINE MONODOSE SANDOZ 250 mg, comprimé pelliculé	not available	34009 387 628 2 2	SANDOZ	FR
AZITHROMYCINE SANDOZ 250 mg, comprimé pelliculé	not available	34009 379 352 1 0	SANDOZ	FR
AZITHROMYCINE SANDOZ 250 mg, comprimé pelliculé	not available	34009 379 353 8 8	SANDOZ	FR
AZITHROMYCINE SANDOZ 250 mg, comprimé pelliculé	not available	34009 570 753 7 8	SANDOZ	FR
AZITHROMYCINE	not available	34009 570 754 3 9	SANDOZ	FR

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SANDOZ 250 mg, comprimé pelliculé				
AZADOSE 600 mg, comprimé pelliculé	not available	34009 343 362 7 0	PFIZER HOLDING FRANCE	FR
AZADOSE 600 mg, comprimé pelliculé	not available	34009 343 361 0 2	PFIZER HOLDING FRANCE	FR
AZADOSE 600 mg, comprimé pelliculé	not available	34009 343 336 6 8	PFIZER HOLDING FRANCE	FR
AZADOSE 600 mg, comprimé pelliculé	not available	34009 343 338 9 7	PFIZER HOLDING FRANCE	FR
AZADOSE 600 mg, comprimé pelliculé	not available	34009 343 363 3 1	PFIZER HOLDING FRANCE	FR
AZADOSE 600 mg, comprimé pelliculé	not available	34009 343 337 2 9	PFIZER HOLDING FRANCE	FR
AZITHROMYCINE VIATRIS 250 mg, comprimé pelliculé	not available	NL 31034	VIATRIS SANTE	FR
ZITHROMAX 250 mg, comprimé pelliculé	not available	34009 351 774 9 0	PFIZER HOLDING FRANCE	FR
ZITHROMAX 40 mg/ml ENFANTS, poudre pour suspension buvable	not available	34009 356 564 2 1	PFIZER HOLDING FRANCE	FR
ZITHROMAX 40 mg/ml ENFANTS, poudre pour suspension buvable	not available	34009 356 565 9 9	PFIZER HOLDING FRANCE	FR
ZITHROMAX 250 mg, comprimé pelliculé	not available	34009 351 773 2 2	PFIZER HOLDING FRANCE	FR
ZITHROMAX MONODOSE 250 mg, comprimé pelliculé	not available	34009 351 777 8 0	PFIZER HOLDING FRANCE	FR
ZITHROMAX MONODOSE 250 mg, comprimé pelliculé	not available	34009 351 778 4 1	PFIZER HOLDING FRANCE	FR
ZITHROMAX 40 mg/ml ENFANTS, poudre pour suspension buvable	not available	34009 356 561 3 1	PFIZER HOLDING FRANCE	FR
ZITHROMAX 40 mg/ml ENFANTS, poudre pour	not available	34009 356 563 6 0	PFIZER HOLDING FRANCE	FR

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suspension buvable				
AZITHROMYCINE ARROW LAB 500 mg, comprimé pelliculé sécable	not available	NL 46280	ARROW GENERIQUES	FR
ZITHROMAX 500 mg επικαλυμμένα με λεπτό υμένιο δισκία.	not available	44409/5-11-2009	PFIZER HELLAS A.E.	GR
ZITHROMAX 200 mg/5 ml κόνις για πόσιμο εναιώρημα.	not available	41167/10/31-5-2011	PFIZER HELLAS A.E.	GR
ZITHROMAX 250 mg επικαλυμμένα με λεπτό υμένιο δισκία.	not available	44407/5-11-2009	PFIZER HELLAS A.E.	GR
ZITHROMAX 250 mg καψάκια σκληρά.	not available	44405/5-11-2009	PFIZER HELLAS A.E.	GR
ZITHROMAX, 500mg/φιαλίδιο, κόνις για διάλυμα προς έγχυση.	not available	44411/5-11-2009	PFIZER HELLAS A.E.	GR
Sumamed 500 mg filmom obložene tablete	not available	HR-H-425760378	PLIVA HRVATSKA D.O.O.	HR
Sumamed 125 mg filmom obložene tablete	not available	HR-H-844869851	PLIVA HRVATSKA D.O.O.	HR
Sumamed 500 mg prašak za koncentrat za otopinu za infuziju	not available	HR-H-261024118	PLIVA HRVATSKA D.O.O.	HR
Sumamed forte 200 mg/5 ml prašak za oralnu suspenziju	not available	HR-H-096865641	PLIVA HRVATSKA D.O.O.	HR
Sumamed 1500, 200 mg/5 ml prašak za oralnu suspenziju	not available	HR-H-096496463	PLIVA HRVATSKA D.O.O.	HR
Sumamed 1200, 200 mg/5 ml prašak za oralnu suspenziju	not available	HR-H-406709782	PLIVA HRVATSKA D.O.O.	HR
Sumamed 100 mg/5 ml prašak za oralnu	not available	HR-H-129408559	PLIVA HRVATSKA D.O.O.	HR

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suspenziju				
Sumamed 500 mg tablete za oralnu suspenziju	not available	HR-H-120748759	PLIVA HRVATSKA D.O.O.	HR
Sumamed 250 mg tvrde kapsule	not available	HR-H-304657205	PLIVA HRVATSKA D.O.O.	HR
Sumamed 100 mg/5 ml por sziruphoz	not available	OGYI-T-5272/03	TEVA GYÓGYSZERGYÁR ZRT	HU
Sumamed forte 200 mg/5 ml por sziruphoz	not available	OGYI-T-5272/02	TEVA GYÓGYSZERGYÁR ZRT	HU
Sumamed forte 200 mg/5 ml por sziruphoz	not available	OGYI-T-5272/08	TEVA GYÓGYSZERGYÁR ZRT	HU
Sumamed 100 mg/ml por oldatos infúzióhoz való koncentrátumhoz	not available	OGYI-T-5272/07	TEVA GYÓGYSZERGYÁR ZRT	HU
Sumamed S 500 mg filmdoboz	not available	OGYI-T-5272/06	TEVA GYÓGYSZERGYÁR ZRT	HU
Sumamed 500 mg filmdoboz	not available	OGYI-T-5272/05	TEVA GYÓGYSZERGYÁR ZRT	HU
Zitrocin 500 mg filmdoboz	not available	OGYI-T-9027/05	TEVA GYÓGYSZERGYÁR ZRT	HU
Sumamed 250 mg kemény kapszula	not available	OGYI-T-5272/04	TEVA GYÓGYSZERGYÁR ZRT	HU
Zithromax Capsules 250 mg.	not available	PA 0822/191/001	PFIZER HEALTHCARE IRELAND	IE
Zithromax Powder for Oral Suspension 200 mg/5 ml.	not available	PA 0822/191/002	PFIZER HEALTHCARE IRELAND	IE
Zitromax 500 mg stofn fyrir innrennslisþykki, lausn.	not available	IS/1/02/106/01	PFIZER APS	IS
Zitromax 500 mg filmhúðaðar töflur.	not available	970282	PFIZER APS	IS
Zitromax 250 mg filmhúðaðar töflur.	not available	970281	PFIZER APS	IS
Zitromax 40 mg/ml mixtúrudoft, dreifa.	not available	920013	PFIZER APS	IS
Azitromicina Teva 250	NL/H/0614/001	037555315	TEVA ITALIA S.R.L.	IT

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mg compresse rivestite con film				
Azitromicina Teva 250 mg compresse rivestite con film	NL/H/0614/001	037555291	TEVA ITALIA S.R.L.	IT
Azitromicina Teva 250 mg compresse rivestite con film	NL/H/0614/001	037555303	TEVA ITALIA S.R.L.	IT
Azitromicina Teva 250 mg compresse rivestite con film	NL/H/0614/001	037555289	TEVA ITALIA S.R.L.	IT
ZITROMAX 200 mg/5 ml polvere per sospensione orale	not available	027860055	PFIZER ITALIA S.R.L.	IT
ZITROMAX 500 mg compresse rivestite con film	not available	027860042	PFIZER ITALIA S.R.L.	IT
ZITROMAX 250 mg capsule rigide	not available	027860016	PFIZER ITALIA S.R.L.	IT
ZITROMAX Avium 600 mg compresse rivestite con film	not available	027860143	PFIZER ITALIA S.R.L.	IT
ZITROMAX 500 mg polvere per soluzione per infusione	not available	027860156	PFIZER ITALIA S.R.L.	IT
ZITROMAX 200 mg/5 ml polvere per sospensione orale	not available	027860079	PFIZER ITALIA S.R.L.	IT
ZITROMAX 100 mg polvere per sospensione orale	not available	027860081	PFIZER ITALIA S.R.L.	IT
ZITROMAX 400 mg polvere per sospensione orale	not available	027860129	PFIZER ITALIA S.R.L.	IT
ZITROMAX 200 mg polvere per sospensione orale	not available	027860105	PFIZER ITALIA S.R.L.	IT
ZITROMAX 150 mg	not available	027860093	PFIZER ITALIA S.R.L.	IT

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
polvere per sospensione orale				
ZITROMAX 300 mg polvere per sospensione orale	not available	027860117	PFIZER ITALIA S.R.L.	IT
ZITROMAX 200 mg/5 ml polvere per sospensione orale	not available	027860067	PFIZER ITALIA S.R.L.	IT
ZITROMAX 200 mg/5 ml polvere per sospensione orale	not available	027860028	PFIZER ITALIA S.R.L.	IT
TROZOCINA 2 g granulato per sospensione orale a rilascio prolungato	not available	027948102	ALFASIGMA S.P.A.	IT
TROZOCINA 200 mg/5 ml polvere per sospensione orale	not available	027948076	ALFASIGMA S.P.A.	IT
TROZOCINA 250 mg capsule rigide	not available	027948049	ALFASIGMA S.P.A.	IT
TROZOCINA 500 mg polvere per soluzione per infusione	not available	027948090	ALFASIGMA S.P.A.	IT
TROZOCINA Avium 600 mg compresse rivestite con film	not available	027948088	ALFASIGMA S.P.A.	IT
TROZOCINA 200 mg/5 ml polvere per sospensione orale	not available	027948052	ALFASIGMA S.P.A.	IT
TROZOCINA 500 mg compresse rivestite con film	not available	027948064	ALFASIGMA S.P.A.	IT
AZITROMICINA PFIZER 200 mg/5 ml polvere per sospensione orale	not available	027897026	PFIZER ITALIA S.R.L.	IT
AZITROMICINA PFIZER 500 mg compresse rivestite con film	not available	027897040	PFIZER ITALIA S.R.L.	IT

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Sumamed forte 200 mg/5 ml milteliai geriamajai suspensijai	not available	LT/1/94/0529/007	TEVA PHARMA B.V.	LT
Sumamed forte 200 mg/5 ml milteliai geriamajai suspensijai	not available	LT/1/94/0529/006	TEVA PHARMA B.V.	LT
Sumamed forte 200 mg/5 ml milteliai geriamajai suspensijai	not available	LT/1/94/0529/008	TEVA PHARMA B.V.	LT
Azibiot 20 mg/ml milteliai geriamajai suspensijai	SK/H/0148/003	LT/1/14/3600/003	KRKA, D.D., NOVO MESTO	LT
Azithromycin Teva 500 mg disperguojamosios tabletės	PL/H/0423/002	LT/1/17/4093/010	TEVA PHARMA B.V.	LT
Azithromycin Teva 500 mg disperguojamosios tabletės	PL/H/0423/002	LT/1/17/4093/009	TEVA PHARMA B.V.	LT
Azithromycin Teva 500 mg disperguojamosios tabletės	PL/H/0423/002	LT/1/17/4093/012	TEVA PHARMA B.V.	LT
Azithromycin Teva 500 mg disperguojamosios tabletės	PL/H/0423/002	LT/1/17/4093/011	TEVA PHARMA B.V.	LT
Sumamed 250 mg kietosios kapsulės	not available	LT/1/94/0529/002	TEVA PHARMA B.V.	LT
Sumamed 500 mg plėvele dengtos tabletės	not available	LT/1/94/0529/004	TEVA PHARMA B.V.	LT
Sumamed 500 mg plėvele dengtos tabletės	not available	LT/1/94/0529/003	TEVA PHARMA B.V.	LT
Azithromycin Sandoz 500 mg plėvele dengtos tabletės	NL/H/4670/002	LT/1/05/0323/007	SANDOZ GMBH	LT
Azithromycin Sandoz 500 mg plėvele dengtos tabletės	NL/H/4670/002	LT/1/05/0323/010	SANDOZ GMBH	LT
Azithromycin Sandoz 500 mg plėvele dengtos tabletės	NL/H/4670/002	LT/1/05/0323/008	SANDOZ GMBH	LT

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
tabletės				
Azithromycin Sandoz 500 mg plėvele dengtos tabletės	NL/H/4670/002	LT/1/05/0323/009	SANDOZ GMBH	LT
Azithromycin Sandoz 500 mg plėvele dengtos tabletės	NL/H/4670/002	LT/1/05/0323/011	SANDOZ GMBH	LT
Azithromycin Sandoz 500 mg plėvele dengtos tabletės	NL/H/4670/002	LT/1/05/0323/012	SANDOZ GMBH	LT
Azithromycin Sandoz 500 mg plėvele dengtos tabletės	NL/H/4670/002	LT/1/05/0323/013	SANDOZ GMBH	LT
Azithromycin Sandoz 500 mg plėvele dengtos tabletės	NL/H/4670/002	LT/1/05/0323/014	SANDOZ GMBH	LT
Zitromax 600 mg Filmtabletten	not available	2011111334	PFIZER S.A.	LU
ZITROMAX 250 mg Comprimés pelliculés	not available	1998080024	PFIZER S.A.	LU
ZITROMAX 200 mg/5 ml Poudre pour suspension buvable	not available	2011111331	PFIZER S.A.	LU
ZITROMAX 500 mg Comprimés pelliculés	not available	2011111333	PFIZER S.A.	LU
Sumamed 250 mg cietās kapsulas	not available	94-0231	TEVA PHARMA B.V.	LV
Sumamed 500 mg apvalkotas tabletes	not available	00-0863	TEVA PHARMA B.V.	LV
Zithromax Capsules 250 mg.	not available	MA505/03902	PFIZER HELLAS, A.E.	MT
Zithromax Powder for Oral Suspension 200 mg/5 ml.	not available	MA505/03901	PFIZER HELLAS, A.E.	MT
Azitromycine Sandoz 100 mg/5 ml, poeder voor orale suspensie	NL/H/0886/001	RVG 32016	SANDOZ B.V.	NL
Zithromax 250, tabletten	not available	RVG 19432	PFIZER B.V.	NL

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250 mg				
Zithromax 500, tabletten 500 mg	not available	RVG 19433	PFIZER B.V.	NL
Zithromax, suspensie (poeder voor) 200 mg/5 ml	not available	RVG 14999	PFIZER B.V.	NL
Azitromax 500 mg tabletter.	not available	1995-00245	PFIZER AS	NO
Azitromax 40 mg/ml, pulver til mikstur, suspensjon.	not available	0000-08025	PFIZER AS	NO
Sumamed 500 mg, tabletki do sporządzania zawiesiny doustnej	PL/H/0423/002	24162	TEVA PHARMACEUTICALS POLSKA SP. Z O.O.	PL
Sumamed 1000 mg, tabletki do sporządzania zawiesiny doustnej	PL/H/0423/003	24163	TEVA PHARMACEUTICALS POLSKA SP. Z O.O.	PL
Sumamed 250 mg, tabletki do sporządzania zawiesiny doustnej	PL/H/0423/001	24161	TEVA PHARMACEUTICALS POLSKA SP. Z O.O.	PL
Sumamed, 500 mg, tabletki powlekane	not available	7424	TEVA PHARMACEUTICALS POLSKA SP. Z O.O.	PL
SUMAMED forte, 200 mg/5 ml, proszek do sporządzania zawiesiny doustnej	not available	7422	TEVA PHARMACEUTICALS POLSKA SP. Z O.O.	PL
Sumamed, 250 mg, kapsułki twarde	not available	7423	TEVA PHARMACEUTICALS POLSKA SP. Z O.O.	PL
SUMAMED, 100 mg/5 ml, proszek do sporządzania zawiesiny doustnej	not available	7421	TEVA PHARMACEUTICALS POLSKA SP. Z O.O.	PL
Azithromycin Teva 250 mg, tabletki do sporządzania zawiesiny doustnej	PL/H/0424/001	24159	TEVA PHARMACEUTICALS POLSKA SP. Z O.O.	PL
Azithromycin Teva, 500 mg, tabletki do	PL/H/0424/002	24160	TEVA PHARMACEUTICALS POLSKA SP. Z O.O.	PL

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
sporządzania zawiesiny doustnej				
Sumamed, 125 mg, tabletki powlekane	not available	8462	TEVA PHARMACEUTICALS POLSKA SP. Z O.O.	PL
Azithromycin Aurovitas, 250 mg, tabletki powlekane	PT/H/1475/001/DC	23591	AUROVITAS PHARMA POLSKA SP. Z O.O	PL
Azithromycin Aurovitas, 500 mg, tabletki powlekane	PT/H/1475/002/DC	23592	AUROVITAS PHARMA POLSKA SP. Z O.O	PL
Sumamed 1000 mg comprimidos dispersíveis	PL/H/0423/003	5722541	TEVA B.V	PT
Sumamed 1000 mg comprimidos dispersíveis	PL/H/0423/003	5722533	TEVA B.V	PT
Zithromax 500 mg comprimido revestido por película	not available	4576385	LABORATÓRIOS PFIZER LDA.	PT
Zithromax 500 mg comprimido revestido por película	not available	2521284	LABORATÓRIOS PFIZER LDA.	PT
Zithromax 40 mg/ml pó para suspensão oral	not available	3969581	LABORATÓRIOS PFIZER LDA.	PT
Zithromax 40 mg/ml pó para suspensão oral	not available	4576286	LABORATÓRIOS PFIZER LDA.	PT
Zithromax 40 mg/ml pó para suspensão oral	not available	2248284	LABORATÓRIOS PFIZER LDA.	PT
Zithromax 40 mg/ml pó para suspensão oral	not available	2248383	LABORATÓRIOS PFIZER LDA.	PT
Zithromax IV 500 mg pó para solução para perfusão	not available	3486982	LABORATÓRIOS PFIZER LDA.	PT
Zithromax IV 500 mg pó para solução para perfusão	not available	5462569	LABORATÓRIOS PFIZER LDA.	PT
Sumamed 500 mg comprimato filmate	not available	8766/2016/01	TEVA B.V	RO
Sumamed 125 mg comprimato filmate	not available	8765/2016/01	TEVA B.V	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
Sumamed Forte 200 mg/5 ml pulbere pentru suspensie orală	not available	8767/2016/02	TEVA B.V	RO
Sumamed Forte 200 mg/5 ml pulbere pentru suspensie orală	not available	8767/2016/03	TEVA B.V	RO
Sumamed Forte 200 mg/5 ml pulbere pentru suspensie orală	not available	8767/2016/01	TEVA B.V	RO
Sumamed 250 mg capsule	not available	10608/2018/01	TEVA B.V	RO
Sumamed Forte 200 mg/5 ml pulbere pentru suspensie orală	not available	8767/2016/02	TEVA B.V	RO
Sumamed Forte 200 mg/5 ml pulbere pentru suspensie orală	not available	8767/2016/03	TEVA B.V	RO
Sumamed Forte 200 mg/5 ml pulbere pentru suspensie orală	not available	8767/2016/01	TEVA B.V	RO
Azitromax 250 mg filmdragerade tabletter	not available	12582	PFIZER AB	SE
Azitromax 500 mg, pulver till koncentrat till infusionsvätska, lösning	not available	16698	PFIZER AB	SE
Azitromax 40 mg/ml pulver till oral suspension.	not available	11609	PFIZER AB	SE
Azitromicin Krka 20 mg/ml prašek za peroralno suspenzijo	SK/H/0148/003	H/14/00246/006	KRKA, D.D., NOVO MESTO	SI
SUMAMED 250 mg trde kapsule	not available	H/92/01472/002	PLIVA LJUBLJANA D.O.O.;	SI
SUMAMED 40 mg/ml prašek za peroralno suspenzijo	not available	H/92/01472/006	PLIVA LJUBLJANA D.O.O.;	SI
SUMAMED 125 mg filmsko obložene tablete	not available	H/92/01472/001	PLIVA LJUBLJANA D.O.O.;	SI

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
SUMAMED 500 mg filmsko obložene tablete	not available	H/92/01472/003	PLIVA LJUBLJANA D.O.O.;	SI
SUMAMED 500 mg prašek za raztopino za infundiranje	not available	H/92/01472/007	PLIVA LJUBLJANA D.O.O.;	SI
SUMAMED za otroke 20 mg/ml prašek za peroralno suspenzijo	not available	H/92/01472/005	PLIVA LJUBLJANA D.O.O.;	SI
Azibiot NEO 20 mg/ml prašek na peroralnu suspenziju	SK/H/0148/003	15/0001/17-S	KRKA, D.D., NOVO MESTO	SK
Sumamed tvrdé kapsuly	not available	15/0226/90-CS	TEVA PHARMACEUTICALS SLOVAKIA S.R.O	SK
Sumamed 500 mg filmom obalené tablety	not available	15/0154/19-S	TEVA PHARMACEUTICALS SLOVAKIA S.R.O	SK
Sumamed 125 mg filmom obalené tablety	not available	15/0351/92-C/S	TEVA PHARMACEUTICALS SLOVAKIA S.R.O	SK
Sumamed forte prašek na sirup	not available	15/0155/19-S	TEVA PHARMACEUTICALS SLOVAKIA S.R.O	SK
Sumamed prašek na sirup	not available	15/0352/92-C/S	TEVA PHARMACEUTICALS SLOVAKIA S.R.O	SK
ZITHROMAX 250 mg Capsules	not available	PL 00057/0335	PFIZER LIMITED	XI
Zithromax 200 mg in 5 ml suspension	not available	PL 00057/0336	PFIZER LIMITED	XI
Azithromycin 250mg Tablets	AT/H/0266/001	PL 04416/1235	SANDOZ LTD	XI
Azithromycin 500 mg Tablets	NL/H/4670/002	PL 04416/0668	SANDOZ LTD	XI
Azithromycin 250mg Capsules	not available	PL 20416/0548	CRESCENT PHARMA LIMITED	XI
Azithromycin 250 mg Capsules	not available	PL 17780/0873	ZENTIVA PHARMA UK LIMITED	XI
Azithromycin 250 mg Film-coated tablets	not available	PL 20416/0482	CRESCENT PHARMA LIMITED	XI
Azithromycin 500 mg Film-coated tablets	not available	PL 20416/0483	CRESCENT PHARMA LIMITED	XI

