



EUROPEAN MEDICINES AGENCY
SCIENCE MEDICINES HEALTH

25 April 2024
EMA/255632/2024
Human Medicines Division

List of nationally authorised medicinal products

Active substance(s): naloxone / oxycodone

Procedure No. PSUSA/00002114/202308



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
Armoneve 10 mg/5 mg depottabletter	DE/H/4793/001-007/DC	55127	MUNDIPHARMA AB	SE
Armoneve 20 mg/10 mg depottabletter	DE/H/4793/001-007/DC	55129	MUNDIPHARMA AB	SE
Armoneve 40 mg/20 mg depottabletter	DE/H/4793/001-007/DC	55131	MUNDIPHARMA AB	SE
Armoneve 5 mg/2,5 mg depottabletter	DE/H/4793/001-007/DC	55126	MUNDIPHARMA AB	SE
OXSYNIA 10 mg/5 mg comprimé à libération prolongée	DE/H/4793/001-007/DC	34009 301 323 2 6	MUNDIPHARMA SAS	FR
OXSYNIA 10 mg/5 mg comprimé à libération prolongée	DE/H/4793/001-007/DC	34009 301 323 1 9	MUNDIPHARMA SAS	FR
OXSYNIA 10 mg/5 mg comprimé à libération prolongée	DE/H/4793/001-007/DC	34009 301 323 0 2	MUNDIPHARMA SAS	FR
OXSYNIA 10 mg/5 mg comprimé à libération prolongée	DE/H/4793/001-007/DC	34009 301 322 3 4	MUNDIPHARMA SAS	FR
OXSYNIA 10 mg/5 mg comprimé à libération prolongée	DE/H/4793/001-007/DC	34009 550 491 6 6	MUNDIPHARMA SAS	FR
OXSYNIA 10 mg/5 mg comprimé à libération prolongée	DE/H/4793/001-007/DC	34009 301 322 9 6	MUNDIPHARMA SAS	FR
OXSYNIA 10 mg/5 mg comprimé à libération prolongée	DE/H/4793/001-007/DC	34009 301 322 8 9	MUNDIPHARMA SAS	FR
OXSYNIA 10 mg/5 mg comprimé à libération prolongée	DE/H/4793/001-007/DC	34009 301 322 6 5	MUNDIPHARMA SAS	FR
OXSYNIA 10 mg/5 mg comprimé à libération prolongée	DE/H/4793/001-007/DC	34009 301 322 5 8	MUNDIPHARMA SAS	FR
OXSYNIA 10 mg/5 mg	DE/H/4793/001-	34009 301 322 4 1	MUNDIPHARMA SAS	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
comprimé à libération prolongée	007/DC			
OXSYNIA 10 mg/5 mg comprimé à libération prolongée	DE/H/4793/001-007/DC	34009 550 491 8 0	MUNDIPHARMA SAS	FR
OXSYNIA 15 mg/7,5 mg, comprimé à libération prolongée	DE/H/4793/001-007/DC	34009 301 321 4 2	MUNDIPHARMA SAS	FR
OXSYNIA 15 mg/7,5 mg, comprimé à libération prolongée	DE/H/4793/001-007/DC	34009 301 321 6 6	MUNDIPHARMA SAS	FR
OXSYNIA 15 mg/7,5 mg, comprimé à libération prolongée	DE/H/4793/001-007/DC	34009 301 321 7 3	MUNDIPHARMA SAS	FR
OXSYNIA 15 mg/7,5 mg, comprimé à libération prolongée	DE/H/4793/001-007/DC	34009 301 321 8 0	MUNDIPHARMA SAS	FR
OXSYNIA 15 mg/7,5 mg, comprimé à libération prolongée	DE/H/4793/001-007/DC	34009 301 321 9 7	MUNDIPHARMA SAS	FR
OXSYNIA 15 mg/7,5 mg, comprimé à libération prolongée	DE/H/4793/001-007/DC	34009 301 322 0 3	MUNDIPHARMA SAS	FR
OXSYNIA 15 mg/7,5 mg, comprimé à libération prolongée	DE/H/4793/001-007/DC	34009 301 322 2 7	MUNDIPHARMA SAS	FR
OXSYNIA 15 mg/7,5 mg, comprimé à libération prolongée	DE/H/4793/001-007/DC	34009 550 491 5 9	MUNDIPHARMA SAS	FR
OXSYNIA 2,5 mg/1,25 mg, comprimé à libération prolongée	DE/H/4793/001-007/DC	34009 301 324 4 9	MUNDIPHARMA SAS	FR
OXSYNIA 2,5 mg/1,25 mg, comprimé à libération prolongée	DE/H/4793/001-007/DC	34009 301 324 9 4	MUNDIPHARMA SAS	FR
OXSYNIA 2,5 mg/1,25 mg, comprimé à libération prolongée	DE/H/4793/001-007/DC	34009 301 324 7 0	MUNDIPHARMA SAS	FR
OXSYNIA 2,5 mg/1,25	DE/H/4793/001-	34009 301 324 6 3	MUNDIPHARMA SAS	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
mg, comprimé à libération prolongée	007/DC			
OXSYNIA 2,5 mg/1,25 mg, comprimé à libération prolongée	DE/H/4793/001-007/DC	34009 301 324 5 6	MUNDIPHARMA SAS	FR
OXSYNIA 2,5 mg/1,25 mg, comprimé à libération prolongée	DE/H/4793/001-007/DC	34009 301 325 2 4	MUNDIPHARMA SAS	FR
OXSYNIA 2,5 mg/1,25 mg, comprimé à libération prolongée	DE/H/4793/001-007/DC	34009 301 325 1 7	MUNDIPHARMA SAS	FR
OXSYNIA 2,5 mg/1,25 mg, comprimé à libération prolongée	DE/H/4793/001-007/DC	34009 301 325 0 0	MUNDIPHARMA SAS	FR
OXSYNIA 2,5 mg/1,25 mg, comprimé à libération prolongée	DE/H/4793/001-007/DC	34009 550 492 1 0	MUNDIPHARMA SAS	FR
OXSYNIA 2,5 mg/1,25 mg, comprimé à libération prolongée	DE/H/4793/001-007/DC	34009 550 492 2 7	MUNDIPHARMA SAS	FR
OXSYNIA 20 mg/10 mg comprimé à libération prolongée	DE/H/4793/001-007/DC	34009 301 320 3 6	MUNDIPHARMA SAS	FR
OXSYNIA 20 mg/10 mg comprimé à libération prolongée	DE/H/4793/001-007/DC	34009 550 491 3 5	MUNDIPHARMA SAS	FR
OXSYNIA 20 mg/10 mg comprimé à libération prolongée	DE/H/4793/001-007/DC	34009 301 321 2 8	MUNDIPHARMA SAS	FR
OXSYNIA 20 mg/10 mg comprimé à libération prolongée	DE/H/4793/001-007/DC	34009 301 321 1 1	MUNDIPHARMA SAS	FR
OXSYNIA 20 mg/10 mg comprimé à libération prolongée	DE/H/4793/001-007/DC	34009 550 491 2 8	MUNDIPHARMA SAS	FR
OXSYNIA 20 mg/10 mg comprimé à libération prolongée	DE/H/4793/001-007/DC	34009 301 320 7 4	MUNDIPHARMA SAS	FR
OXSYNIA 20 mg/10 mg	DE/H/4793/001-	34009 301 321 3 5	MUNDIPHARMA SAS	FR

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comprimé à libération prolongée	007/DC			
OXSYNIA 20 mg/10 mg comprimé à libération prolongée	DE/H/4793/001-007/DC	34009 301 320 6 7	MUNDIPHARMA SAS	FR
OXSYNIA 20 mg/10 mg comprimé à libération prolongée	DE/H/4793/001-007/DC	34009 301 320 9 8	MUNDIPHARMA SAS	FR
OXSYNIA 20 mg/10 mg comprimé à libération prolongée	DE/H/4793/001-007/DC	34009 301 320 5 0	MUNDIPHARMA SAS	FR
OXSYNIA 30 mg/15 mg, comprimé à libération prolongée	DE/H/4793/001-007/DC	34009 301 319 3 0	MUNDIPHARMA SAS	FR
OXSYNIA 30 mg/15 mg, comprimé à libération prolongée	DE/H/4793/001-007/DC	34009 550 491 1 1	MUNDIPHARMA SAS	FR
OXSYNIA 30 mg/15 mg, comprimé à libération prolongée	DE/H/4793/001-007/DC	34009 550 491 0 4	MUNDIPHARMA SAS	FR
OXSYNIA 30 mg/15 mg, comprimé à libération prolongée	DE/H/4793/001-007/DC	34009 301 320 2 9	MUNDIPHARMA SAS	FR
OXSYNIA 30 mg/15 mg, comprimé à libération prolongée	DE/H/4793/001-007/DC	34009 301 320 1 2	MUNDIPHARMA SAS	FR
OXSYNIA 30 mg/15 mg, comprimé à libération prolongée	DE/H/4793/001-007/DC	34009 301 320 0 5	MUNDIPHARMA SAS	FR
OXSYNIA 30 mg/15 mg, comprimé à libération prolongée	DE/H/4793/001-007/DC	34009 301 319 7 8	MUNDIPHARMA SAS	FR
OXSYNIA 30 mg/15 mg, comprimé à libération prolongée	DE/H/4793/001-007/DC	34009 301 319 8 5	MUNDIPHARMA SAS	FR
OXSYNIA 30 mg/15 mg, comprimé à libération prolongée	DE/H/4793/001-007/DC	34009 301 319 5 4	MUNDIPHARMA SAS	FR
OXSYNIA 30 mg/15 mg,	DE/H/4793/001-	34009 301 319 4 7	MUNDIPHARMA SAS	FR

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comprimé à libération prolongée	007/DC			
OXYNIA 30 mg/15 mg, comprimé à libération prolongée	DE/H/4793/001-007/DC	34009 301 319 9 2	MUNDIPHARMA SAS	FR
OXYNIA 40 mg/20 mg comprimé à libération prolongée	DE/H/4793/001-007/DC	34009 301 318 6 2	MUNDIPHARMA SAS	FR
OXYNIA 40 mg/20 mg comprimé à libération prolongée	DE/H/4793/001-007/DC	34009 550 490 9 8	MUNDIPHARMA SAS	FR
OXYNIA 40 mg/20 mg comprimé à libération prolongée	DE/H/4793/001-007/DC	34009 550 490 7 4	MUNDIPHARMA SAS	FR
OXYNIA 40 mg/20 mg comprimé à libération prolongée	DE/H/4793/001-007/DC	34009 550 490 6 7	MUNDIPHARMA SAS	FR
OXYNIA 40 mg/20 mg comprimé à libération prolongée	DE/H/4793/001-007/DC	34009 550 490 8 1	MUNDIPHARMA SAS	FR
OXYNIA 40 mg/20 mg comprimé à libération prolongée	DE/H/4793/001-007/DC	34009 301 318 7 9	MUNDIPHARMA SAS	FR
OXYNIA 40 mg/20 mg comprimé à libération prolongée	DE/H/4793/001-007/DC	34009 301 319 2 3	MUNDIPHARMA SAS	FR
OXYNIA 40 mg/20 mg comprimé à libération prolongée	DE/H/4793/001-007/DC	34009 301 318 9 3	MUNDIPHARMA SAS	FR
OXYNIA 5 mg/2,5 mg comprimé à libération prolongée	DE/H/4793/001-007/DC	34009 301 323 4 0	MUNDIPHARMA SAS	FR
OXYNIA 5 mg/2,5 mg comprimé à libération prolongée	DE/H/4793/001-007/DC	34009 301 323 9 5	MUNDIPHARMA SAS	FR
OXYNIA 5 mg/2,5 mg comprimé à libération prolongée	DE/H/4793/001-007/DC	34009 301 323 8 8	MUNDIPHARMA SAS	FR
OXYNIA 5 mg/2,5 mg	DE/H/4793/001-	34009 301 323 6 4	MUNDIPHARMA SAS	FR

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comprimé à libération prolongée	007/DC			
OXSYNIA 5 mg/2,5 mg comprimé à libération prolongée	DE/H/4793/001-007/DC	34009 301 323 7 1	MUNDIPHARMA SAS	FR
OXSYNIA 5 mg/2,5 mg comprimé à libération prolongée	DE/H/4793/001-007/DC	34009 301 323 5 7	MUNDIPHARMA SAS	FR
OXSYNIA 5 mg/2,5 mg comprimé à libération prolongée	DE/H/4793/001-007/DC	34009 301 324 1 8	MUNDIPHARMA SAS	FR
OXSYNIA 5 mg/2,5 mg comprimé à libération prolongée	DE/H/4793/001-007/DC	34009 301 324 2 5	MUNDIPHARMA SAS	FR
OXSYNIA 5 mg/2,5 mg comprimé à libération prolongée	DE/H/4793/001-007/DC	34009 550 491 9 7	MUNDIPHARMA SAS	FR
OXSYNIA 5 mg/2,5 mg comprimé à libération prolongée	DE/H/4793/001-007/DC	34009 550 492 0 3	MUNDIPHARMA SAS	FR
OXSYNIA 5 mg/2,5 mg comprimé à libération prolongée	DE/H/4793/001-007/DC	34009 301 324 0 1	MUNDIPHARMA SAS	FR
OXSYNIA LP 15 mg/7,5 mg, comprimé à libération prolongée	DE/H/4793/004	34009 301 322 1 0	MUNDIPHARMA SAS	FR
OXSYNIA LP 15 mg/7,5 mg, comprimé à libération prolongée	DE/H/4793/004	34009 550 491 4 2	MUNDIPHARMA SAS	FR
OXSYNIA LP 15 mg/7,5 mg, comprimé à libération prolongée	DE/H/4793/004	34009 301 321 5 9	MUNDIPHARMA SAS	FR
OXSYNIA LP 2,5 mg/1,25 mg, comprimé à libération prolongée	DE/H/4793/001	34009 301 324 8 7	MUNDIPHARMA SAS	FR
OXSYNIA LP 20 mg/10 mg comprimé à libération prolongée	DE/H/4793/005	34009 301 320 8 1	MUNDIPHARMA SAS	FR
OXSYNIA LP 40 mg/20	DE/H/4793/007	34009 301 319 1 6	MUNDIPHARMA SAS	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
mg comprimé à libération prolongée				
OXYYNIA LP 40 mg/20 mg comprimé à libération prolongée	DE/H/4793/007	34009 301 319 0 9	MUNDIPHARMA SAS	FR
OXYYNIA LP 40 mg/20 mg comprimé à libération prolongée	DE/H/4793/007	34009 301 318 8 6	MUNDIPHARMA SAS	FR
Oxycodon / Naloxon Krugmann 10 mg/5 mg Retardtabletten	DE/H/4793/001-007/DC	98165.00.00	KRUGMANN GMBH (GERMANY)	DE
Oxycodon / Naloxon KRUGMANN 10 mg/5 mg Retardtabletten	not available	91850.00.00	KRUGMANN GMBH (GERMANY)	DE
Oxycodon / Naloxon Krugmann 15 mg/7,5 mg Retardtabletten	DE/H/4793/004	98166.00.00	KRUGMANN GMBH (GERMANY)	DE
Oxycodon / Naloxon Krugmann 2,5 mg/1,25 mg Retardtabletten	DE/H/4793/001-007/DC	98163.00.00	KRUGMANN GMBH (GERMANY)	DE
Oxycodon / Naloxon Krugmann 20 mg/10 mg Retardtabletten	DE/H/4793/001-007/DC	98167.00.00	KRUGMANN GMBH (GERMANY)	DE
Oxycodon / Naloxon KRUGMANN 20 mg/10 mg Retardtabletten	not available	91851.00.00	KRUGMANN GMBH (GERMANY)	DE
Oxycodon / Naloxon Krugmann 30 mg/15 mg Retardtabletten	DE/H/4793/001-007/DC	98168.00.00	KRUGMANN GMBH (GERMANY)	DE
Oxycodon / Naloxon Krugmann 40 mg/20 mg Retardtabletten	DE/H/4793/001-007/DC	98169.00.00	KRUGMANN GMBH (GERMANY)	DE
Oxycodon / Naloxon KRUGMANN 40 mg/20 mg Retardtabletten	not available	91852.00.00	KRUGMANN GMBH (GERMANY)	DE
Oxycodon / Naloxon Krugmann 5 mg/2,5 mg Retardtabletten	DE/H/4793/001-007/DC	98164.00.00	KRUGMANN GMBH (GERMANY)	DE
Oxycodon / Naloxon	not available	91849.00.00	KRUGMANN 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
KRUGMANN 5 mg/2,5 mg Retardtabletten			(GERMANY)	
Oxycodon / Naloxon KRUGMANN 60 mg / 30 mg Retardtabletten	not available	2200024.00.00	KRUGMANN GMBH (GERMANY)	DE
Oxycodon / Naloxon KRUGMANN 80 mg / 40 mg Retardtabletten	not available	2200025.00.00	KRUGMANN GMBH (GERMANY)	DE
Targin 10 mg/ 5 mg comprimidos de liberación prolongada	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	71124	MUNDIPHARMA PHARMACEUTICALS SL	ES
Targin 10 mg/5 mg compresse a rilascio prolungato	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	39586223	MUNDIPHARMA PHARMACEUTICALS SRL	IT
Targin 10 mg/5 mg compresse a rilascio prolungato	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	39586159	MUNDIPHARMA PHARMACEUTICALS SRL	IT
Targin 10 mg/5 mg compresse a rilascio prolungato	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	39586173	MUNDIPHARMA PHARMACEUTICALS SRL	IT
Targin 10 mg/5 mg compresse a rilascio prolungato	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	39586209	MUNDIPHARMA PHARMACEUTICALS SRL	IT
Targin 10 mg/5 mg compresse a rilascio prolungato	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	39586134	MUNDIPHARMA PHARMACEUTICALS SRL	IT
Targin 10 mg/5 mg compresse a rilascio prolungato	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	39586185	MUNDIPHARMA PHARMACEUTICALS SRL	IT
Targin 10 mg/5 mg compresse a rilascio prolungato	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	39586211	MUNDIPHARMA PHARMACEUTICALS SRL	IT
Targin 10 mg/5 mg compresse a rilascio prolungato	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	39586197	MUNDIPHARMA PHARMACEUTICALS SRL	IT
Targin 10 mg/5 mg compresse a rilascio prolungato	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	39586146	MUNDIPHARMA PHARMACEUTICALS SRL	IT
Targin 10 mg/5 mg	DE/H/1612/004;	39586161	MUNDIPHARMA	IT

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comprese a rilascio prolungato	DE/H/1612/003; DE/H/1612/002; DE/H/		PHARMACEUTICALS SRL	
Targin 10 mg/5 mg compresse a rilascio prolungato	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	39586122	MUNDIPHARMA PHARMACEUTICALS SRL	IT
Targin 10 mg/5 mg comprimidos de libertação prolongada_001	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	5219043	MUNDIPHARMA FARMACÊUTICA, LDA.	PT
Targin 10 mg/5 mg comprimidos de libertação prolongada_002	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	5260153	MUNDIPHARMA FARMACÊUTICA, LDA.	PT
Targin 10 mg/5 mg comprimidos de libertação prolongada_003	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	5219035	MUNDIPHARMA FARMACÊUTICA, LDA.	PT
Targin 10 mg/5 mg forðatöflur	DE/H/1545/001	IS/1/08/096/01	MUNDIPHARMA A/S	IS
Targin 10 mg/5 mg prolonged-release tablets	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	PA 1688/010/002	MUNDIPHARMA PHARMACEUTICALS LIMITED	IE
Targin 10 mg/5 mg Retardtabletten	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	1-27955	MUNDIPHARMA GES.M.B.H	AT
Targin 10 mg/5 mg Retardtabletten	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	73083.00.00	MUNDIPHARMA GMBH	DE
Targin 10 mg/5 mg tablety s prodlouženým uvolňováním	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	65/468/09-C	MUNDIPHARMA GES.M.B.H	CZ
Targin 10/5 mg tablety s predĺženým uvoľňovaním	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	65/0625/10-S	MUNDIPHARMA GESELLSCHAFT M.B.H.	SK
Targin 15 mg/7,5 mg compresse a rilascio prolungato_001	DE/H/1612/005;DE/H/1612/006;DE/H/1612/007	39586565	MUNDIPHARMA PHARMACEUTICALS SRL	IT
Targin 15 mg/7,5 mg compresse a rilascio	DE/H/1612/005;DE/H/1612/006;DE/H/1612/	39586577	MUNDIPHARMA PHARMACEUTICALS SRL	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
prolungato_002	007			
Targin 15 mg/7,5 mg compresse a rilascio prolungato_003	DE/H/1612/005;DE/H/1612/006;DE/H/1612/007	39586589	MUNDIPHARMA PHARMACEUTICALS SRL	IT
Targin 15 mg/7,5 mg compresse a rilascio prolungato_004	DE/H/1612/005;DE/H/1612/006;DE/H/1612/007	39586591	MUNDIPHARMA PHARMACEUTICALS SRL	IT
Targin 15 mg/7,5 mg compresse a rilascio prolungato_005	DE/H/1612/005;DE/H/1612/006;DE/H/1612/007	39586603	MUNDIPHARMA PHARMACEUTICALS SRL	IT
Targin 15 mg/7,5 mg compresse a rilascio prolungato_006	DE/H/1612/005;DE/H/1612/006;DE/H/1612/007	39586615	MUNDIPHARMA PHARMACEUTICALS SRL	IT
Targin 15 mg/7,5 mg compresse a rilascio prolungato_007	DE/H/1612/005;DE/H/1612/006;DE/H/1612/007	39586627	MUNDIPHARMA PHARMACEUTICALS SRL	IT
Targin 15 mg/7,5 mg compresse a rilascio prolungato_008	DE/H/1612/005;DE/H/1612/006;DE/H/1612/007	39586639	MUNDIPHARMA PHARMACEUTICALS SRL	IT
Targin 15 mg/7,5 mg compresse a rilascio prolungato_009	DE/H/1612/005;DE/H/1612/006;DE/H/1612/007	39586641	MUNDIPHARMA PHARMACEUTICALS SRL	IT
Targin 15 mg/7,5 mg compresse a rilascio prolungato_010	DE/H/1612/005;DE/H/1612/006;DE/H/1612/007	39586654	MUNDIPHARMA PHARMACEUTICALS SRL	IT
Targin 15 mg/7,5 mg compresse a rilascio prolungato_011	DE/H/1612/005;DE/H/1612/006;DE/H/1612/007	39586666	MUNDIPHARMA PHARMACEUTICALS SRL	IT
Targin 15 mg/7,5 mg comprimidos de libertação prolongada	DE/H/1612/006	5609508	MUNDIPHARMA PHARMACEUTICALS LTD	PT
Targin 15 mg/7,5 mg Retardtabletten	DE/H/1612/005;DE/H/1612/006;DE/H/1612/007	135732	MUNDIPHARMA GES.M.B.H	AT
Targin 15 mg/7,5 mg Retardtabletten	DE/H/1612/005;DE/H/1612/006;DE/H/1612/007	88178.00.00	MUNDIPHARMA GMBH	DE
Targin 15 mg/7.5 mg prolonged-release	DE/H/1612/005;DE/H/1612/006;DE/H/1612/	PA 1688/010/006	MUNDIPHARMA PHARMACEUTICALS	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
tablets	007		LIMITED	
Targin 2,5 mg/1,25 mg compresse a rilascio prolungato_001	DE/H/1612/005;DE/H/1612/006;DE/H/1612/007	39586452	MUNDIPHARMA PHARMACEUTICALS SRL	IT
Targin 2,5 mg/1,25 mg compresse a rilascio prolungato_002	DE/H/1612/005;DE/H/1612/006;DE/H/1612/007	39586464	MUNDIPHARMA PHARMACEUTICALS SRL	IT
Targin 2,5 mg/1,25 mg compresse a rilascio prolungato_003	DE/H/1612/005;DE/H/1612/006;DE/H/1612/007	39586476	MUNDIPHARMA PHARMACEUTICALS SRL	IT
Targin 2,5 mg/1,25 mg compresse a rilascio prolungato_004	DE/H/1612/005;DE/H/1612/006;DE/H/1612/007	39586488	MUNDIPHARMA PHARMACEUTICALS SRL	IT
Targin 2,5 mg/1,25 mg compresse a rilascio prolungato_005	DE/H/1612/005;DE/H/1612/006;DE/H/1612/007	39586490	MUNDIPHARMA PHARMACEUTICALS SRL	IT
Targin 2,5 mg/1,25 mg compresse a rilascio prolungato_006	DE/H/1612/005;DE/H/1612/006;DE/H/1612/007	39586502	MUNDIPHARMA PHARMACEUTICALS SRL	IT
Targin 2,5 mg/1,25 mg compresse a rilascio prolungato_007	DE/H/1612/005;DE/H/1612/006;DE/H/1612/007	39586514	MUNDIPHARMA PHARMACEUTICALS SRL	IT
Targin 2,5 mg/1,25 mg compresse a rilascio prolungato_008	DE/H/1612/005;DE/H/1612/006;DE/H/1612/007	39586526	MUNDIPHARMA PHARMACEUTICALS SRL	IT
Targin 2,5 mg/1,25 mg compresse a rilascio prolungato_009	DE/H/1612/005;DE/H/1612/006;DE/H/1612/007	39586538	MUNDIPHARMA PHARMACEUTICALS SRL	IT
Targin 2,5 mg/1,25 mg compresse a rilascio prolungato_010	DE/H/1612/005;DE/H/1612/006;DE/H/1612/007	39586540	MUNDIPHARMA PHARMACEUTICALS SRL	IT
Targin 2,5 mg/1,25 mg compresse a rilascio prolungato_011	DE/H/1612/005;DE/H/1612/006;DE/H/1612/007	39586553	MUNDIPHARMA PHARMACEUTICALS SRL	IT
Targin 2,5 mg/1,25 mg comprimidos de libertação prolongada	DE/H/1612/005	5609516	MUNDIPHARMA PHARMACEUTICALS LTD	PT
Targin 2,5mg/1,25 mg Retardtabletten	DE/H/1612/005;DE/H/1612/006;DE/H/1612/	135730	MUNDIPHARMA GES.M.B.H	AT

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	007			
Targin 2,5mg/1,25 mg Retardtabletten	DE/H/1612/005;DE/H/1612/006;DE/H/1612/007	88177.00.00	MUNDIPHARMA GMBH	DE
Targin 20 mg/10 mg compresse a rilascio prolungato	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	39586250	MUNDIPHARMA PHARMACEUTICALS SRL	IT
Targin 20 mg/10 mg compresse a rilascio prolungato	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	39586274	MUNDIPHARMA PHARMACEUTICALS SRL	IT
Targin 20 mg/10 mg compresse a rilascio prolungato	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	39586298	MUNDIPHARMA PHARMACEUTICALS SRL	IT
Targin 20 mg/10 mg compresse a rilascio prolungato	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	39586324	MUNDIPHARMA PHARMACEUTICALS SRL	IT
Targin 20 mg/10 mg compresse a rilascio prolungato	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	39586235	MUNDIPHARMA PHARMACEUTICALS SRL	IT
Targin 20 mg/10 mg compresse a rilascio prolungato	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	39586336	MUNDIPHARMA PHARMACEUTICALS SRL	IT
Targin 20 mg/10 mg compresse a rilascio prolungato	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	39586312	MUNDIPHARMA PHARMACEUTICALS SRL	IT
Targin 20 mg/10 mg compresse a rilascio prolungato	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	39586262	MUNDIPHARMA PHARMACEUTICALS SRL	IT
Targin 20 mg/10 mg compresse a rilascio prolungato	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	39586286	MUNDIPHARMA PHARMACEUTICALS SRL	IT
Targin 20 mg/10 mg compresse a rilascio prolungato	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	39586247	MUNDIPHARMA PHARMACEUTICALS SRL	IT
Targin 20 mg/10 mg compresse a rilascio prolungato	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	39586300	MUNDIPHARMA PHARMACEUTICALS SRL	IT
Targin 20 mg/10 mg comprimidos de	DE/H/1612/004; DE/H/1612/003;	71125	MUNDIPHARMA PHARMACEUTICALS SL	ES

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liberación prolongada	DE/H/1612/002; DE/H/			
Targin 20 mg/10 mg comprimidos de libertação prolongada_001	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	5219068	MUNDIPHARMA FARMACÊUTICA, LDA.	PT
Targin 20 mg/10 mg comprimidos de libertação prolongada_002	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	5260161	MUNDIPHARMA FARMACÊUTICA, LDA.	PT
Targin 20 mg/10 mg comprimidos de libertação prolongada_003	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	5219050	MUNDIPHARMA FARMACÊUTICA, LDA.	PT
Targin 20 mg/10 mg forðatöflur	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	IS/1/08/096/02	MUNDIPHARMA A/S	IS
Targin 20 mg/10 mg prolonged-release tablets	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	PA 1688/010/003	MUNDIPHARMA PHARMACEUTICALS LIMITED	IE
Targin 20 mg/10 mg Retardtabletten	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	1-27956	MUNDIPHARMA GES.M.B.H	AT
Targin 20 mg/10 mg Retardtabletten	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	73084.00.00	MUNDIPHARMA GMBH	DE
Targin 20 mg/10 mg tablety s prodlouženým uvolňováním	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	65/469/09-C	MUNDIPHARMA GES.M.B.H	CZ
Targin 20/10 mg tablety s predĺženým uvoľňovaním	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	65/0626/10-S	MUNDIPHARMA GESELLSCHAFT M.B.H.	SK
Targin 30 mg/15 mg compresse a rilascio prolungato	DE/H/1612/005;DE/H/1612/006;DE/H/1612/007	39586716	MUNDIPHARMA PHARMACEUTICALS SRL	IT
Targin 30 mg/15 mg compresse a rilascio prolungato	DE/H/1612/005;DE/H/1612/006;DE/H/1612/007	39586767	MUNDIPHARMA PHARMACEUTICALS SRL	IT
Targin 30 mg/15 mg compresse a rilascio	DE/H/1612/005;DE/H/1612/006;DE/H/1612/	39586704	MUNDIPHARMA PHARMACEUTICALS SRL	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
prolungato	007			
Targin 30 mg/15 mg compresse a rilascio prolungato	DE/H/1612/005;DE/H/1612/006;DE/H/1612/007	39586678	MUNDIPHARMA PHARMACEUTICALS SRL	IT
Targin 30 mg/15 mg compresse a rilascio prolungato	DE/H/1612/005;DE/H/1612/006;DE/H/1612/007	39586742	MUNDIPHARMA PHARMACEUTICALS SRL	IT
Targin 30 mg/15 mg compresse a rilascio prolungato	DE/H/1612/005;DE/H/1612/006;DE/H/1612/007	39586692	MUNDIPHARMA PHARMACEUTICALS SRL	IT
Targin 30 mg/15 mg compresse a rilascio prolungato	DE/H/1612/005;DE/H/1612/006;DE/H/1612/007	39586779	MUNDIPHARMA PHARMACEUTICALS SRL	IT
Targin 30 mg/15 mg compresse a rilascio prolungato	DE/H/1612/005;DE/H/1612/006;DE/H/1612/007	39586680	MUNDIPHARMA PHARMACEUTICALS SRL	IT
Targin 30 mg/15 mg compresse a rilascio prolungato	DE/H/1612/005;DE/H/1612/006;DE/H/1612/007	39586755	MUNDIPHARMA PHARMACEUTICALS SRL	IT
Targin 30 mg/15 mg compresse a rilascio prolungato	DE/H/1612/005;DE/H/1612/006;DE/H/1612/007	39586728	MUNDIPHARMA PHARMACEUTICALS SRL	IT
Targin 30 mg/15 mg compresse a rilascio prolungato	DE/H/1612/005;DE/H/1612/006;DE/H/1612/007	39586730	MUNDIPHARMA PHARMACEUTICALS SRL	IT
Targin 30 mg/15 mg prolonged-release tablets	DE/H/1612/005;DE/H/1612/006;DE/H/1612/007	PA1688/010/007	MUNDIPHARMA PHARMACEUTICALS LIMITED	IE
Targin 30 mg/15 mg Retardtabletten	DE/H/1612/005;DE/H/1612/006;DE/H/1612/007	135733	MUNDIPHARMA GES.M.B.H	AT
Targin 30 mg/15 mg Retardtabletten	DE/H/1612/005;DE/H/1612/006;DE/H/1612/007	88179.00.00	MUNDIPHARMA GMBH	DE
Targin 30 mg/15 mg comprimidos de libertação prolongada	DE/H/1612/007	5609524	MUNDIPHARMA PHARMACEUTICALS LTD	PT
Targin 40 mg/20 mg compresse a rilascio	DE/H/1612/004;DE/H/1612/003;	39586437	MUNDIPHARMA PHARMACEUTICALS SRL	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
prolungato	DE/H/1612/002; DE/H/			
Targin 40 mg/20 mg compresse a rilascio prolungato	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	39586375	MUNDIPHARMA PHARMACEUTICALS SRL	IT
Targin 40 mg/20 mg compresse a rilascio prolungato	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	39586401	MUNDIPHARMA PHARMACEUTICALS SRL	IT
Targin 40 mg/20 mg compresse a rilascio prolungato	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	39586348	MUNDIPHARMA PHARMACEUTICALS SRL	IT
Targin 40 mg/20 mg compresse a rilascio prolungato	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	39586399	MUNDIPHARMA PHARMACEUTICALS SRL	IT
Targin 40 mg/20 mg compresse a rilascio prolungato	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	39586449	MUNDIPHARMA PHARMACEUTICALS SRL	IT
Targin 40 mg/20 mg compresse a rilascio prolungato	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	39586351	MUNDIPHARMA PHARMACEUTICALS SRL	IT
Targin 40 mg/20 mg compresse a rilascio prolungato	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	39586363	MUNDIPHARMA PHARMACEUTICALS SRL	IT
Targin 40 mg/20 mg compresse a rilascio prolungato	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	39586387	MUNDIPHARMA PHARMACEUTICALS SRL	IT
Targin 40 mg/20 mg compresse a rilascio prolungato	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	39586413	MUNDIPHARMA PHARMACEUTICALS SRL	IT
Targin 40 mg/20 mg compresse a rilascio prolungato	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	39586425	MUNDIPHARMA PHARMACEUTICALS SRL	IT
Targin 40 mg/20 mg comprimidos de liberación prolongada	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	71126	MUNDIPHARMA PHARMACEUTICALS SL	ES
Targin 40 mg/20 mg comprimidos de libertação prolongada_001	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	5219100	MUNDIPHARMA FARMACÊUTICA, LDA.	PT
Targin 40 mg/20 mg	DE/H/1612/004;	5260179	MUNDIPHARMA	PT

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comprimidos de libertação prolongada_002	DE/H/1612/003; DE/H/1612/002; DE/H/		FARMACÊUTICA, LDA.	
Targin 40 mg/20 mg comprimidos de libertação prolongada_003	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	5219076	MUNDIPHARMA FARMACÊUTICA, LDA.	PT
Targin 40 mg/20 mg forðatöflur	DE/H/1612/003	IS/1/09/006/02	MUNDIPHARMA A/S	IS
Targin 40 mg/20 mg prolonged-release tablets	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	PA 1688/010/004	MUNDIPHARMA PHARMACEUTICALS LIMITED	IE
Targin 40 mg/20 mg Retardtabletten	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	1-28503	MUNDIPHARMA GES.M.B.H	AT
Targin 40 mg/20 mg Retardtabletten	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	73085.00.00	MUNDIPHARMA GMBH	DE
Targin 40 mg/20 mg tablety s prodĺouŕeným uvolňováním	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	65/470/09-C	MUNDIPHARMA GES.M.B.H	CZ
Targin 40/20 mg tablety s predĺženým uvolnovaním	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	65/0627/10-S	MUNDIPHARMA GESELLSCHAFT M.B.H.	SK
Targin 5 mg/2,5 mg compresse a rilascio prolungato	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	39586058	MUNDIPHARMA PHARMACEUTICALS SRL	IT
Targin 5 mg/2,5 mg compresse a rilascio prolungato	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	39586060	MUNDIPHARMA PHARMACEUTICALS SRL	IT
Targin 5 mg/2,5 mg compresse a rilascio prolungato	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	39586045	MUNDIPHARMA PHARMACEUTICALS SRL	IT
Targin 5 mg/2,5 mg compresse a rilascio prolungato	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	39586084	MUNDIPHARMA PHARMACEUTICALS SRL	IT
Targin 5 mg/2,5 mg compresse a rilascio prolungato	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	39586110	MUNDIPHARMA PHARMACEUTICALS SRL	IT

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Targin 5 mg/2,5 mg compresse a rilascio prolungato	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	39586021	MUNDIPHARMA PHARMACEUTICALS SRL	IT
Targin 5 mg/2,5 mg compresse a rilascio prolungato	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	39586108	MUNDIPHARMA PHARMACEUTICALS SRL	IT
Targin 5 mg/2,5 mg compresse a rilascio prolungato	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	39586096	MUNDIPHARMA PHARMACEUTICALS SRL	IT
Targin 5 mg/2,5 mg compresse a rilascio prolungato	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	39586019	MUNDIPHARMA PHARMACEUTICALS SRL	IT
Targin 5 mg/2,5 mg compresse a rilascio prolungato	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	39586033	MUNDIPHARMA PHARMACEUTICALS SRL	IT
Targin 5 mg/2,5 mg compresse a rilascio prolungato	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	39586072	MUNDIPHARMA PHARMACEUTICALS SRL	IT
Targin 5 mg/2,5 mg comprimidos de liberación prolongada	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	71127	MUNDIPHARMA PHARMACEUTICALS SL	ES
Targin 5 mg/2,5 mg comprimidos de libertação prolongada	DE/H/1612/004	5219118	MUNDIPHARMA FARMACÊUTICA, LDA.	PT
Targin 5 mg/2,5 mg comprimidos de libertação prolongada	DE/H/1612/004	5260203	MUNDIPHARMA FARMACÊUTICA, LDA.	PT
Targin 5 mg/2,5 mg comprimidos de libertação prolongada	DE/H/1612/004	5219126	MUNDIPHARMA FARMACÊUTICA, LDA.	PT
targin 5 mg/2,5 mg forðatöflur	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	IS/1/09/006/01	MUNDIPHARMA A/S	IS
Targin 5 mg/2,5 mg Retardtabletten	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	73086.00.00	MUNDIPHARMA GMBH	DE
Targin 5 mg/2.5 mg prolonged-release tablets	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	PA1688/010/001	MUNDIPHARMA PHARMACEUTICALS 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
Targin 5 mg/2.5 mg Retardtabletten	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	1-28502	MUNDIPHARMA GES.M.B.H	AT
Targin 5/2,5 mg tablety s predĺženým uvoľňovaním	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	65/0624/10-S	MUNDIPHARMA GESELLSCHAFT M.B.H.	SK
Targin 60 mg/30 mg prolonged release tablets	DE/H/1612/008/DC;DE /H/1612/009/DC	PA 1688/010/008	MUNDIPHARMA PHARMACEUTICALS LIMITED	IE
Targin 60 mg/30 mg Retardtabletten	DE/H/1612/008/DC;DE /H/1612/009/DC	137663	MUNDIPHARMA GES.M.B.H	AT
Targin 60 mg/30 mg Retardtabletten	DE/H/1612/008/DC;DE /H/1612/009/DC	94173.00.00	MUNDIPHARMA GMBH	DE
Targin 80 mg/40 mg prolonged release tablets	DE/H/1612/008/DC;DE /H/1612/009/DC	PA 1688/010/009	MUNDIPHARMA PHARMACEUTICALS LIMITED	IE
Targin 80 mg/40 mg Retardtabletten	DE/H/1612/008/DC;DE /H/1612/009/DC	137664	MUNDIPHARMA GES.M.B.H	AT
Targin 80 mg/40 mg Retardtabletten	DE/H/1612/008/DC;DE /H/1612/009/DC	94174.00.00	MUNDIPHARMA GMBH	DE
Targin, 10 mg + 5 mg, tabletki o przedłużonym uwalnianiu	DE/H/1612/001	16106	MUNDIPHARMA A/S	PL
Targin, 20 mg + 10 mg, tabletki o przedłużonym uwalnianiu	DE/H/1612/002	16107	MUNDIPHARMA A/S	PL
Targin, 40 mg + 20 mg, tabletki o przedłużonym uwalnianiu	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	16108	MUNDIPHARMA A/S	PL
Targin, 5 mg + 2,5 mg, tabletki o przedłużonym uwalnianiu	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	16109	MUNDIPHARMA A/S	PL
Targin, depottabletter 10 mg/5 mg	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	43673	MUNDIPHARMA A/S	DK
Targin, depottabletter 15 mg/7.5 mg	DE/H/1612/005;DE/H/1612/006;DE/H/1612/007	51503	MUNDIPHARMA A/S	DK

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Targin, depottabletter 20 mg/10 mg	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	43674	MUNDIPHARMA A/S	DK
Targin, depottabletter 30 mg/15 mg	DE/H/1612/005;DE/H/ 1612/006;DE/H/1612/ 007	51504	MUNDIPHARMA A/S	DK
Targin, depottabletter 40 mg/20 mg	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	43427	MUNDIPHARMA A/S	DK
Targin, depottabletter 5 mg/2,5 mg	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	43426	MUNDIPHARMA A/S	DK
Targinact 10 mg /5 mg, tabletten met verlengde afgifte	DE/H/1612/001	BE330626	MUNDIPHARMA COMM VA	BE
Targinact 10 mg, 5 mg, tabletten met verlengde afgifte	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	BE434271	MUNDIPHARMA COMM VA	BE
Targinact 10 mg/5 mg tablete s podaljšanim sproščanjem	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	H/10/01492/026	MUNDIPHARMA GES.M.B.H	SI
Targinact 10 mg/5 mg tablete s podaljšanim sproščanjem	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	H/10/01492/033	MUNDIPHARMA GES.M.B.H	SI
Targinact 10 mg/5 mg tablete s podaljšanim sproščanjem	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	H/10/01492/028	MUNDIPHARMA GES.M.B.H	SI
Targinact 10 mg/5 mg tablete s podaljšanim sproščanjem	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	H/10/01492/030	MUNDIPHARMA GES.M.B.H	SI
Targinact 10 mg/5 mg tablete s podaljšanim sproščanjem	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	H/10/01492/029	MUNDIPHARMA GES.M.B.H	SI
Targinact 10 mg/5 mg tablete s podaljšanim sproščanjem	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	H/10/01492/027	MUNDIPHARMA GES.M.B.H	SI
Targinact 10 mg/5 mg tablete s podaljšanim sproščanjem	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	H/10/01492/031	MUNDIPHARMA GES.M.B.H	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
Targinact 10 mg/5 mg tablete s podaljšanim sproščanjem	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	H/10/01492/034	MUNDIPHARMA GES.M.B.H	SI
Targinact 10 mg/5 mg tablete s podaljšanim sproščanjem	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	H/10/01492/035	MUNDIPHARMA GES.M.B.H	SI
Targinact 10 mg/5 mg tablete s podaljšanim sproščanjem	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	H/10/01492/032	MUNDIPHARMA GES.M.B.H	SI
Targinact 10 mg/5 mg tablete s podaljšanim sproščanjem	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	H/10/01492/024	MUNDIPHARMA GES.M.B.H	SI
Targinact 10 mg/5 mg tablete s podaljšanim sproščanjem	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	H/10/01492/025	MUNDIPHARMA GES.M.B.H	SI
Targinact 10 mg/5 mg tablete s produljenim oslobađanjem	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	HR-H-347037202	MUNDIPHARMA GES.M.B.H	HR
Targinact 10 mg/5 mg tablete s produljenim oslobađanjem	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	HR-H-347037202-01	MUNDIPHARMA GES.M.B.H	HR
Targinact 10 mg/5 mg tablete s produljenim oslobađanjem	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	HR-H-347037202-02	MUNDIPHARMA GES.M.B.H	HR
Targinact 10 mg/5 mg tablete s produljenim oslobađanjem	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	HR-H-347037202-03	MUNDIPHARMA GES.M.B.H	HR
Targinact 10 mg/5 mg tablete s produljenim oslobađanjem	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	HR-H-347037202-04	MUNDIPHARMA GES.M.B.H	HR
Targinact 10 mg/5 mg tablete s produljenim oslobađanjem	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	HR-H-347037202-08	MUNDIPHARMA GES.M.B.H	HR
Targinact 10 mg/5 mg tablete s produljenim oslobađanjem	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	HR-H-347037202-07	MUNDIPHARMA GES.M.B.H	HR
Targinact 10 mg/5 mg tablete s produljenim oslobađanjem	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	HR-H-347037202-06	MUNDIPHARMA GES.M.B.H	HR

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Targinact 10 mg/5 mg tablete s produljenim oslobađanjem	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	HR-H-347037202-12	MUNDIPHARMA GES.M.B.H	HR
Targinact 10 mg/5 mg tablete s produljenim oslobađanjem	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	HR-H-347037202-11	MUNDIPHARMA GES.M.B.H	HR
Targinact 10 mg/5 mg tablete s produljenim oslobađanjem	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	HR-H-347037202-10	MUNDIPHARMA GES.M.B.H	HR
Targinact 10 mg/5 mg tablete s produljenim oslobađanjem	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	HR-H-347037202-09	MUNDIPHARMA GES.M.B.H	HR
Targinact 10 mg/5 mg tablete s produljenim oslobađanjem	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	HR-H-347037202-05	MUNDIPHARMA GES.M.B.H	HR
Targinact 10 mg/5 mg δισκία παρατεταμένης αποδέσμευσης	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	20570	MUNDIPHARMA PHARMACEUTICALS LTD	CY
Targinact 10 mg/5 mg, comprimés à libération prolongée	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	BE434271	MUNDIPHARMA COMM VA	BE
Targinact 10 mg/5 mg, comprimés à libération prolongée	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	BE330626	MUNDIPHARMA COMM VA	BE
Targinact 10 mg/5 mg, comprimés à libération prolongée	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	2009030016	MUNDIPHARMA COMM VA	LU
Targinact 20 mg/10 mg tablete s podaljšanim sproščanjem	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	H/10/01492/055	MUNDIPHARMA GES.M.B.H	SI
Targinact 20 mg/10 mg tablete s podaljšanim sproščanjem	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	H/10/01492/050	MUNDIPHARMA GES.M.B.H	SI
Targinact 20 mg/10 mg tablete s podaljšanim sproščanjem	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	H/10/01492/058	MUNDIPHARMA GES.M.B.H	SI
Targinact 20 mg/10 mg tablete s podaljšanim sproščanjem	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	H/10/01492/051	MUNDIPHARMA GES.M.B.H	SI

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Targinact 20 mg/10 mg tablete s podaljšanim sproščanjem	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	H/10/01492/056	MUNDIPHARMA GES.M.B.H	SI
Targinact 20 mg/10 mg tablete s podaljšanim sproščanjem	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	H/10/01492/057	MUNDIPHARMA GES.M.B.H	SI
Targinact 20 mg/10 mg tablete s podaljšanim sproščanjem	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	H/10/01492/049	MUNDIPHARMA GES.M.B.H	SI
Targinact 20 mg/10 mg tablete s podaljšanim sproščanjem	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	H/10/01492/054	MUNDIPHARMA GES.M.B.H	SI
Targinact 20 mg/10 mg tablete s podaljšanim sproščanjem	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	H/10/01492/053	MUNDIPHARMA GES.M.B.H	SI
Targinact 20 mg/10 mg tablete s podaljšanim sproščanjem	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	H/10/01492/052	MUNDIPHARMA GES.M.B.H	SI
Targinact 20 mg/10 mg tablete s podaljšanim sproščanjem	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	H/10/01492/047	MUNDIPHARMA GES.M.B.H	SI
Targinact 20 mg/10 mg tablete s podaljšanim sproščanjem	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	H/10/01492/048	MUNDIPHARMA GES.M.B.H	SI
Targinact 20 mg/10 mg tablete s produljenim oslobađanjem	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	HR-H-639349110	MUNDIPHARMA GES.M.B.H	HR
Targinact 20 mg/10 mg tablete s produljenim oslobađanjem	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	HR-H-639349110-12	MUNDIPHARMA GES.M.B.H	HR
Targinact 20 mg/10 mg tablete s produljenim oslobađanjem	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	HR-H-639349110-05	MUNDIPHARMA GES.M.B.H	HR
Targinact 20 mg/10 mg tablete s produljenim oslobađanjem	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	HR-H-639349110-10	MUNDIPHARMA GES.M.B.H	HR
Targinact 20 mg/10 mg tablete s produljenim oslobađanjem	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	HR-H-639349110-01	MUNDIPHARMA GES.M.B.H	HR

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Targinact 20 mg/10 mg tablete s produljenim oslobađanjem	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	HR-H-639349110-03	MUNDIPHARMA GES.M.B.H	HR
Targinact 20 mg/10 mg tablete s produljenim oslobađanjem	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	HR-H-639349110-11	MUNDIPHARMA GES.M.B.H	HR
Targinact 20 mg/10 mg tablete s produljenim oslobađanjem	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	HR-H-639349110-09	MUNDIPHARMA GES.M.B.H	HR
Targinact 20 mg/10 mg tablete s produljenim oslobađanjem	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	HR-H-639349110-08	MUNDIPHARMA GES.M.B.H	HR
Targinact 20 mg/10 mg tablete s produljenim oslobađanjem	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	HR-H-639349110-06	MUNDIPHARMA GES.M.B.H	HR
Targinact 20 mg/10 mg tablete s produljenim oslobađanjem	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	HR-H-639349110-07	MUNDIPHARMA GES.M.B.H	HR
Targinact 20 mg/10 mg tablete s produljenim oslobađanjem	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	HR-H-639349110-04	MUNDIPHARMA GES.M.B.H	HR
Targinact 20 mg/10 mg tablete s produljenim oslobađanjem	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	HR-H-639349110-02	MUNDIPHARMA GES.M.B.H	HR
Targinact 20 mg/10 mg δισκία παρατεταμένης αποδέσμευσης	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	20571	MUNDIPHARMA PHARMACEUTICALS LTD	CY
Targinact 20 mg/10 mg, comprimés à libération prolongée	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	BE330635	MUNDIPHARMA COMM VA	BE
Targinact 20 mg/10 mg, comprimés à libération prolongée	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	BE434287	MUNDIPHARMA COMM VA	BE
Targinact 20 mg/10 mg, comprimés à libération prolongée	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	2009030017	MUNDIPHARMA COMM VA	LU
Targinact 40 mg/ 20 mg, comprimés à libération prolongée	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	BE341923	MUNDIPHARMA COMM VA	BE

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Targinact 40 mg/ 20 mg, comprimés à libération prolongée	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	BE434296	MUNDIPHARMA COMM VA	BE
Targinact 40 mg/ 20 mg, comprimés à libération prolongée	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	2009110049	MUNDIPHARMA COMM VA	LU
Targinact 40 mg/20 mg tablete s podaljšanim sproščanjem	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	H/10/01492/075	MUNDIPHARMA GES.M.B.H	SI
Targinact 40 mg/20 mg tablete s podaljšanim sproščanjem	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	H/10/01492/076	MUNDIPHARMA GES.M.B.H	SI
Targinact 40 mg/20 mg tablete s podaljšanim sproščanjem	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	H/10/01492/074	MUNDIPHARMA GES.M.B.H	SI
Targinact 40 mg/20 mg tablete s podaljšanim sproščanjem	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	H/10/01492/078	MUNDIPHARMA GES.M.B.H	SI
Targinact 40 mg/20 mg tablete s podaljšanim sproščanjem	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	H/10/01492/079	MUNDIPHARMA GES.M.B.H	SI
Targinact 40 mg/20 mg tablete s podaljšanim sproščanjem	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	H/10/01492/081	MUNDIPHARMA GES.M.B.H	SI
Targinact 40 mg/20 mg tablete s podaljšanim sproščanjem	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	H/10/01492/072	MUNDIPHARMA GES.M.B.H	SI
Targinact 40 mg/20 mg tablete s podaljšanim sproščanjem	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	H/10/01492/080	MUNDIPHARMA GES.M.B.H	SI
Targinact 40 mg/20 mg tablete s podaljšanim sproščanjem	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	H/10/01492/077	MUNDIPHARMA GES.M.B.H	SI
Targinact 40 mg/20 mg tablete s podaljšanim sproščanjem	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	H/10/01492/073	MUNDIPHARMA GES.M.B.H	SI
Targinact 40 mg/20 mg tablete s podaljšanim sproščanjem	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	H/10/01492/070	MUNDIPHARMA GES.M.B.H	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
Targinact 40 mg/20 mg tablete s podaljšanim sproščanjem	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	H/10/01492/071	MUNDIPHARMA GES.M.B.H	SI
Targinact 40 mg/20 mg tablete s produljenim oslobađanjem	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	HR-H-875883221	MUNDIPHARMA GES.M.B.H	HR
Targinact 40 mg/20 mg tablete s produljenim oslobađanjem	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	HR-H-875883221-10	MUNDIPHARMA GES.M.B.H	HR
Targinact 40 mg/20 mg tablete s produljenim oslobađanjem	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	HR-H-875883221-12	MUNDIPHARMA GES.M.B.H	HR
Targinact 40 mg/20 mg tablete s produljenim oslobađanjem	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	HR-H-875883221-11	MUNDIPHARMA GES.M.B.H	HR
Targinact 40 mg/20 mg tablete s produljenim oslobađanjem	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	HR-H-875883221-09	MUNDIPHARMA GES.M.B.H	HR
Targinact 40 mg/20 mg tablete s produljenim oslobađanjem	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	HR-H-875883221-06	MUNDIPHARMA GES.M.B.H	HR
Targinact 40 mg/20 mg tablete s produljenim oslobađanjem	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	HR-H-875883221-08	MUNDIPHARMA GES.M.B.H	HR
Targinact 40 mg/20 mg tablete s produljenim oslobađanjem	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	HR-H-875883221-07	MUNDIPHARMA GES.M.B.H	HR
Targinact 40 mg/20 mg tablete s produljenim oslobađanjem	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	HR-H-875883221-05	MUNDIPHARMA GES.M.B.H	HR
Targinact 40 mg/20 mg tablete s produljenim oslobađanjem	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	HR-H-875883221-01	MUNDIPHARMA GES.M.B.H	HR
Targinact 40 mg/20 mg tablete s produljenim oslobađanjem	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	HR-H-875883221-04	MUNDIPHARMA GES.M.B.H	HR
Targinact 40 mg/20 mg tablete s produljenim oslobađanjem	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	HR-H-875883221-03	MUNDIPHARMA GES.M.B.H	HR

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Targinact 40 mg/20 mg tablete s produljenim oslobađanjem	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	HR-H-875883221-02	MUNDIPHARMA GES.M.B.H	HR
Targinact 40 mg/20 mg δισκία παρατεταμένης αποδέσμευσης	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	20572	MUNDIPHARMA PHARMACEUTICALS LTD	CY
Targinact 40 mg/20 mg, tabletten met verlengde afgifte	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	BE434296	MUNDIPHARMA COMM VA	BE
Targinact 40 mg/20 mg, tabletten met verlengde afgifte	DE/H/1612/003	BE434296	MUNDIPHARMA COMM VA	BE
Targinact 5 mg/ 2,5 mg, comprimés à libération prolongée	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	BE341914	MUNDIPHARMA COMM VA	BE
Targinact 5 mg/ 2,5 mg, comprimés à libération prolongée	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	BE434305	MUNDIPHARMA COMM VA	BE
Targinact 5 mg/ 2,5 mg, comprimés à libération prolongée	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	2009110048	MUNDIPHARMA COMM VA	LU
Targinact 5 mg/2,5 mg tablete s podaljšanim sproščanjem	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	H/10/01492/015	MUNDIPHARMA GES.M.B.H	SI
Targinact 5 mg/2,5 mg tablete s podaljšanim sproščanjem	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	H/10/01492/021	MUNDIPHARMA GES.M.B.H	SI
Targinact 5 mg/2,5 mg tablete s podaljšanim sproščanjem	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	H/10/01492/020	MUNDIPHARMA GES.M.B.H	SI
Targinact 5 mg/2,5 mg tablete s podaljšanim sproščanjem	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	H/10/01492/023	MUNDIPHARMA GES.M.B.H	SI
Targinact 5 mg/2,5 mg tablete s podaljšanim sproščanjem	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	H/10/01492/014	MUNDIPHARMA GES.M.B.H	SI
Targinact 5 mg/2,5 mg tablete s podaljšanim sproščanjem	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	H/10/01492/022	MUNDIPHARMA GES.M.B.H	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
Targinact 5 mg/2,5 mg tablete s podaljšanim sproščanjem	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	H/10/01492/017	MUNDIPHARMA GES.M.B.H	SI
Targinact 5 mg/2,5 mg tablete s podaljšanim sproščanjem	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	H/10/01492/018	MUNDIPHARMA GES.M.B.H	SI
Targinact 5 mg/2,5 mg tablete s podaljšanim sproščanjem	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	H/10/01492/019	MUNDIPHARMA GES.M.B.H	SI
Targinact 5 mg/2,5 mg tablete s podaljšanim sproščanjem	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	H/10/01492/016	MUNDIPHARMA GES.M.B.H	SI
Targinact 5 mg/2,5 mg tablete s podaljšanim sproščanjem	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	H/10/01492/012	MUNDIPHARMA GES.M.B.H	SI
Targinact 5 mg/2,5 mg tablete s podaljšanim sproščanjem	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	H/10/01492/013	MUNDIPHARMA GES.M.B.H	SI
Targinact 5 mg/2,5 mg tablete s produljenim oslobađanjem	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	HR-H-961831038	MUNDIPHARMA GES.M.B.H	HR
Targinact 5 mg/2,5 mg tablete s produljenim oslobađanjem	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	HR-H-961831038-06	MUNDIPHARMA GES.M.B.H	HR
Targinact 5 mg/2,5 mg tablete s produljenim oslobađanjem	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	HR-H-961831038-05	MUNDIPHARMA GES.M.B.H	HR
Targinact 5 mg/2,5 mg tablete s produljenim oslobađanjem	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	HR-H-961831038-03	MUNDIPHARMA GES.M.B.H	HR
Targinact 5 mg/2,5 mg tablete s produljenim oslobađanjem	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	HR-H-961831038-10	MUNDIPHARMA GES.M.B.H	HR
Targinact 5 mg/2,5 mg tablete s produljenim oslobađanjem	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	HR-H-961831038-12	MUNDIPHARMA GES.M.B.H	HR
Targinact 5 mg/2,5 mg tablete s produljenim oslobađanjem	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	HR-H-961831038-11	MUNDIPHARMA GES.M.B.H	HR

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Targinact 5 mg/2,5 mg tablete s produljenim oslobađanjem	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	HR-H-961831038-08	MUNDIPHARMA GES.M.B.H	HR
Targinact 5 mg/2,5 mg tablete s produljenim oslobađanjem	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	HR-H-961831038-09	MUNDIPHARMA GES.M.B.H	HR
Targinact 5 mg/2,5 mg tablete s produljenim oslobađanjem	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	HR-H-961831038-07	MUNDIPHARMA GES.M.B.H	HR
Targinact 5 mg/2,5 mg tablete s produljenim oslobađanjem	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	HR-H-961831038-04	MUNDIPHARMA GES.M.B.H	HR
Targinact 5 mg/2,5 mg tablete s produljenim oslobađanjem	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	HR-H-961831038-02	MUNDIPHARMA GES.M.B.H	HR
Targinact 5 mg/2,5 mg tablete s produljenim oslobađanjem	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	HR-H-961831038-01	MUNDIPHARMA GES.M.B.H	HR
Targinact 5 mg/2,5 mg, tabletten met verlengde afgifte	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	BE434305	MUNDIPHARMA COMM VA	BE
Targinact 5 mg/2,5 mg, tabletten met verlengde afgifte	DE/H/1612/004	BE434305	MUNDIPHARMA COMM VA	BE
Targinact 5 mg/2.5 mg δισκία παρατεταμένης αποδέσμευσης	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	20573	MUNDIPHARMA PHARMACEUTICALS LTD	CY
Targinact Retard 10 mg/5 mg, tabletten met verlengde afgifte	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	RVG 102951	MUNDIPHARMA PHARMACEUTICALS BV	NL
Targinact Retard 20 mg/10 mg, tabletten met verlengde afgifte	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	BE330635	MUNDIPHARMA COMM VA	BE
Targinact Retard 20 mg/10 mg, tabletten met verlengde afgifte	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	BE434287	MUNDIPHARMA COMM VA	BE
Targinact Retard 20 mg/10 mg, tabletten met verlengde afgifte	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	RVG 102961	MUNDIPHARMA PHARMACEUTICALS BV	NL

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Targinact Retard 40 mg /20 mg, tabletten met verlengde afgifte	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	RVG 102647	MUNDIPHARMA PHARMACEUTICALS BV	NL
Targinact Retard 5 mg /2,5 mg tabletten met verlengde afgifte	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	RVG 102645	MUNDIPHARMA PHARMACEUTICALS BV	NL
Targinact, 10 mg/5 mg toimeainet prolongeeritult vabastavad tabletid	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	716210	MUNDIPHARMA GES.M.B.H	EE
Targinact, 20 mg/10 mg toimeainet prolongeeritult vabastavad tabletid	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	716110	MUNDIPHARMA GES.M.B.H	EE
Targinact, 40 mg/20 mg toimeainet prolongeeritult vabastavad tabletid	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	716010	MUNDIPHARMA GES.M.B.H	EE
Targinact, 5 mg/2,5 mg toimeainet prolongeeritult vabastavad tabletid	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	716310	MUNDIPHARMA GES.M.B.H	EE
Targiniq 10 mg/5 mg depottabletter	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	08-6097	MUNDIPHARMA AS	NO
Targiniq 10 mg/5 mg depottabletter	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	27745	MUNDIPHARMA AB	SE
Targiniq 10 mg/5 mg depottabletti	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	25238	MUNDIPHARMA OY	FI
Targiniq 15 mg/7,5 mg depottabletter	DE/H/1612/005;DE/H/1612/006;DE/H/1612/007	12-9215	MUNDIPHARMA AS	NO
Targiniq 15 mg/7,5 mg depottabletti	DE/H/1612/005;DE/H/1612/006;DE/H/1612/007	31037	MUNDIPHARMA OY	FI
Targiniq 20 mg/10 mg depottabletter	DE/H/1612/004; DE/H/1612/003;	08-6098	MUNDIPHARMA AS	NO

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
	DE/H/1612/002; DE/H/			
Targiniq 20 mg/10 mg depottabletter	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	27746	MUNDIPHARMA AB	SE
Targiniq 20 mg/10 mg depottabletti	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	25239	MUNDIPHARMA OY	FI
Targiniq 30 mg/15 mg depottabletter	DE/H/1612/005;DE/H/ 1612/006;DE/H/1612/ 007	12-9216	MUNDIPHARMA AS	NO
Targiniq 30 mg/15 mg depottabletti	DE/H/1612/005;DE/H/ 1612/006;DE/H/1612/ 007	31038	MUNDIPHARMA OY	FI
Targiniq 40 mg/20 mg depottabletter	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	08-5980	MUNDIPHARMA AS	NO
Targiniq 40 mg/20 mg depottabletter	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	27082	MUNDIPHARMA AB	SE
Targiniq 40 mg/20 mg depottabletti	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	25068	MUNDIPHARMA OY	FI
Targiniq 5 mg/2,5 mg depottabletter	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	08-5981	MUNDIPHARMA AS	NO
Targiniq 5 mg/2.5 mg depottabletter	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	27083	MUNDIPHARMA AB	SE
Targiniq 5 mg/2.5 mg depottabletti	DE/H/1612/004; DE/H/1612/003; DE/H/1612/002; DE/H/	25069	MUNDIPHARMA OY	FI