



EUROPEAN MEDICINES AGENCY  
SCIENCE MEDICINES HEALTH

25 March 2021  
EMA/146362/2021  
Human Medicines Evaluation Division

## List of nationally authorised medicinal products

Active substance: quetiapine

Procedure no.: PSUSA/00002589/202007

---

**Official address** Domenico Scarlattilaan 6 • 1083 HS Amsterdam • The Netherlands

**Address for visits and deliveries** Refer to [www.ema.europa.eu/how-to-find-us](http://www.ema.europa.eu/how-to-find-us)

**Send us a question** Go to [www.ema.europa.eu/contact](http://www.ema.europa.eu/contact) **Telephone** +31 (0)88 781 6000

An agency of the European Union



<b>Product Name (in authorisation country)</b>	<b>MRP/DCP Authorisation number</b>	<b>National Authorisation Number</b>	<b>MAH of product in the member state</b>	<b>Member State where product is authorised</b>
Alcreno, 150 mg, tabletki powlekane	NL/H/4782/003	18456	ACCORD HEALTHCARE POLSKA SP. Z O.O.	PL
Alcreno, 300 mg, tabletki powlekane	NL/H/4782/005	18458	ACCORD HEALTHCARE POLSKA SP. Z O.O.	PL
Alzen SR 150 mg comprimidos de libertação prolongada	PT/H/1439/002	5176953	ASTRAZENECA PRODUTOS FARMACEUTICOS LDA	PT
Alzen SR 150 mg comprimidos de libertação prolongada	PT/H/1439/002	5176938	ASTRAZENECA PRODUTOS FARMACEUTICOS LDA	PT
Alzen SR 150 mg comprimidos de libertação prolongada	PT/H/1439/002	5176961	ASTRAZENECA PRODUTOS FARMACEUTICOS LDA	PT
Alzen SR 150 mg comprimidos de libertação prolongada	PT/H/1439/002	5176979	ASTRAZENECA PRODUTOS FARMACEUTICOS LDA	PT
Alzen SR 150 mg comprimidos de libertação prolongada	PT/H/1439/002	5176946	ASTRAZENECA PRODUTOS FARMACEUTICOS LDA	PT
Alzen SR 200 mg comprimidos de libertação prolongada	PT/H/1439/003	5090832	ASTRAZENECA PRODUTOS FARMACEUTICOS LDA	PT
Alzen SR 200 mg comprimidos de libertação prolongada	PT/H/1439/003	5090857	ASTRAZENECA PRODUTOS FARMACEUTICOS LDA	PT
Alzen SR 200 mg comprimidos de libertação prolongada	PT/H/1439/003	5090865	ASTRAZENECA PRODUTOS FARMACEUTICOS LDA	PT
Alzen SR 200 mg comprimidos de libertação prolongada	PT/H/1439/003	5090873	ASTRAZENECA PRODUTOS FARMACEUTICOS LDA	PT
Alzen SR 200 mg comprimidos de libertação prolongada	PT/H/1439/003	5090840	ASTRAZENECA PRODUTOS FARMACEUTICOS LDA	PT

<b>Product Name (in authorisation country)</b>	<b>MRP/DCP Authorisation number</b>	<b>National Authorisation Number</b>	<b>MAH of product in the member state</b>	<b>Member State where product is authorised</b>
Alzen SR 300 mg comprimidos de libertação prolongada	PT/H/1439/004	5090915	ASTRAZENECA PRODUTOS FARMACEUTICOS LDA	PT
Alzen SR 300 mg comprimidos de libertação prolongada	PT/H/1439/004	5090949	ASTRAZENECA PRODUTOS FARMACEUTICOS LDA	PT
Alzen SR 300 mg comprimidos de libertação prolongada	PT/H/1439/004	5090907	ASTRAZENECA PRODUTOS FARMACEUTICOS LDA	PT
Alzen SR 300 mg comprimidos de libertação prolongada	PT/H/1439/004	5090923	ASTRAZENECA PRODUTOS FARMACEUTICOS LDA	PT
Alzen SR 300 mg comprimidos de libertação prolongada	PT/H/1439/004	5090931	ASTRAZENECA PRODUTOS FARMACEUTICOS LDA	PT
Alzen SR 400 mg comprimidos de libertação prolongada	PT/H/1439/005	5090964	ASTRAZENECA PRODUTOS FARMACEUTICOS LDA	PT
Alzen SR 400 mg comprimidos de libertação prolongada	PT/H/1439/005	5090972	ASTRAZENECA PRODUTOS FARMACEUTICOS LDA	PT
Alzen SR 400 mg comprimidos de libertação prolongada	PT/H/1439/005	5091012	ASTRAZENECA PRODUTOS FARMACEUTICOS LDA	PT
Alzen SR 400 mg comprimidos de libertação prolongada	PT/H/1439/005	5090956	ASTRAZENECA PRODUTOS FARMACEUTICOS LDA	PT
Alzen SR 400 mg comprimidos de libertação prolongada	PT/H/1439/005	5091004	ASTRAZENECA PRODUTOS FARMACEUTICOS LDA	PT
Alzen SR 50 mg comprimidos de libertação prolongada	PT/H/1439/001	5090808	ASTRAZENECA PRODUTOS FARMACEUTICOS LDA	PT
Alzen SR 50 mg	PT/H/1439/001	5090824	ASTRAZENECA PRODUTOS	PT

<b>Product Name (in authorisation country)</b>	<b>MRP/DCP Authorisation number</b>	<b>National Authorisation Number</b>	<b>MAH of product in the member state</b>	<b>Member State where product is authorised</b>
comprimidos de libertação prolongada			FARMACEUTICOS LDA	
Alzen SR 50 mg comprimidos de libertação prolongada	PT/H/1439/001	5090774	ASTRAZENECA PRODUTOS FARMACEUTICOS LDA	PT
Alzen SR 50 mg comprimidos de libertação prolongada	PT/H/1439/001	5090816	ASTRAZENECA PRODUTOS FARMACEUTICOS LDA	PT
Alzen SR 50 mg comprimidos de libertação prolongada	PT/H/1439/001	5090766	ASTRAZENECA PRODUTOS FARMACEUTICOS LDA	PT
Biquelle XL 150 mg prolonged-release tablets	not available	PL 35533/0052	ASPIRE PHARMA LIMITED	UK
Biquelle XL 200 mg prolonged-release tablets	not available	PL 35533/0053	ASPIRE PHARMA LIMITED	UK
Biquelle XL 300 mg prolonged-release tablets	not available	PL 35533/0054	ASPIRE PHARMA LIMITED	UK
Biquelle XL 400 mg prolonged-release tablets	not available	PL 35533/0055	ASPIRE PHARMA LIMITED	UK
Biquelle XL 50 mg prolonged-release tablets	not available	PL 35533/0051	ASPIRE PHARMA LIMITED	UK
Biquelle XL 600 mg prolonged-release tablets	not available	PL 35533/0155	ASPIRE PHARMA LIMITED	UK
Desiquet 20 mg/ml Suspension zum Einnehmen	IE/H/1009/001	93617.00.00	ROSEMONT PHARMACEUTICALS LIMITED	DE
Ketilept 150 mg filmtabletta	HU/H/0126/001-005/MR	OGYI-T-20056/09; OGYI-T-20056/11; OGYI-T-20056/56-60	EGIS PHARMACEUTICALS PLC	HU
Ketilept 150 mg apvalkotās tabletes	HU/H/0126/001-005/MR	07-0333	EGIS PHARMACEUTICALS PLC	LV
Ketilept 150 mg comprimato filmate	HU/H/0126/001-005/MR	4214/2012/03-04	EGIS PHARMACEUTICALS PLC	RO
Ketilept 150 mg filmom obalené tablety	HU/H/0126/001-005/MR	68/0143/07-S	EGIS PHARMACEUTICALS PLC	SK

<b>Product Name (in authorisation country)</b>	<b>MRP/DCP Authorisation number</b>	<b>National Authorisation Number</b>	<b>MAH of product in the member state</b>	<b>Member State where product is authorised</b>
Ketilept 150 mg filmtableta	HU/H/0126/001-005/MR	OGYI-T-20056/10; OGYI-T-20056/12	EGIS PHARMACEUTICALS PLC	HU
Ketilept 150 mg filmtableta	HU/H/0126/001-005/MR	OGYI-T-20056/09; OGYI-T-20056/11; OGYI-T-20056/56-60	EGIS PHARMACEUTICALS PLC	HU
Ketilept 300 mg apvalkotās tabletes	HU/H/0126/001-005/MR	07-0335	EGIS PHARMACEUTICALS PLC	LV
Ketilept 300 mg potahované tablety	HU/H/0126/001-005/MR	68/255/07-C	EGIS PHARMACEUTICALS PLC	CZ
Ketilept 300 mg potahované tablety	HU/H/0126/001-005/MR	68/255/07-C	EGIS PHARMACEUTICALS PLC	CZ
Ketilept Prolong 150 mg tablety s predĺženým uvoľňovaním	HU/H/0579/001-005/DC	68/0033/15-S	EGIS PHARMACEUTICALS PLC	SK
Kvelux 100 mg filmsko obložene tablete	DK/H/1527/001	H/09/01997/004	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 100 mg filmsko obložene tablete	DK/H/1527/001	H/09/01997/007	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 100 mg filmsko obložene tablete	DK/H/1527/001	H/09/01997/014	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 100 mg filmsko obložene tablete	DK/H/1527/001	H/09/01997/015	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 100 mg filmsko obložene tablete	DK/H/1527/001	H/09/01997/001	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 100 mg filmsko obložene tablete	DK/H/1527/001	H/09/01997/002	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 100 mg filmsko obložene tablete	DK/H/1527/001	H/09/01997/003	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 100 mg filmsko obložene tablete	DK/H/1527/001	H/09/01997/006	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 100 mg filmsko obložene tablete	DK/H/1527/001	H/09/01997/005	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 100 mg filmsko obložene tablete	DK/H/1527/001	H/09/01997/008	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 100 mg filmsko obložene tablete	DK/H/1527/001	H/09/01997/009	LEK PHARMACEUTICALS	SI

<b>Product Name (in authorisation country)</b>	<b>MRP/DCP Authorisation number</b>	<b>National Authorisation Number</b>	<b>MAH of product in the member state</b>	<b>Member State where product is authorised</b>
obložene tablete			D.D. LJUBLJANA	
Kvelux 100 mg filmsko obložene tablete	DK/H/1527/001	H/09/01997/010	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 100 mg filmsko obložene tablete	DK/H/1527/001	H/09/01997/011	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 100 mg filmsko obložene tablete	DK/H/1527/001	H/09/01997/012	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 100 mg filmsko obložene tablete	DK/H/1527/001	H/09/01997/013	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 100 mg filmsko obložene tablete	DK/H/1527/001	H/09/01997/045	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 100 mg filmsko obložene tablete	DK/H/1527/001	H/09/01997/046	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 100 mg filmsko obložene tablete	DK/H/1527/001	H/09/01997/047	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 100 mg filmsko obložene tablete	DK/H/1527/001	H/09/01997/049	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 100 mg filmsko obložene tablete	DK/H/1527/001	H/09/01997/050	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 100 mg filmsko obložene tablete	DK/H/1527/001	H/09/01997/051	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 100 mg filmsko obložene tablete	DK/H/1527/001	H/09/01997/052	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 100 mg filmsko obložene tablete	DK/H/1527/001	H/09/01997/053	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 100 mg filmsko obložene tablete	DK/H/1527/001	H/09/01997/054	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 100 mg filmsko obložene tablete	DK/H/1527/001	H/09/01997/055	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 100 mg filmsko obložene tablete	DK/H/1527/001	H/09/01997/056	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 100 mg filmsko obložene tablete	DK/H/1527/001	H/09/01997/057	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 100 mg filmsko obložene tablete	DK/H/1527/001	H/09/01997/058	LEK PHARMACEUTICALS	SI

<b>Product Name (in authorisation country)</b>	<b>MRP/DCP Authorisation number</b>	<b>National Authorisation Number</b>	<b>MAH of product in the member state</b>	<b>Member State where product is authorised</b>
obložene tablete			D.D. LJUBLJANA	
Kvelux 100 mg filmsko obložene tablete	DK/H/1527/001	H/09/01997/059	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 100 mg filmsko obložene tablete	DK/H/1527/001	H/09/01997/060	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 100 mg filmsko obložene tablete	DK/H/1527/001	H/09/01997/061	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 100 mg filmsko obložene tablete	DK/H/1527/001	H/09/01997/062	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 100 mg filmsko obložene tablete	DK/H/1527/001	H/09/01997/063	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 100 mg filmsko obložene tablete	DK/H/1527/001	H/09/01997/064	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 100 mg filmsko obložene tablete	DK/H/1527/001	H/09/01997/065	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 100 mg filmsko obložene tablete	DK/H/1527/001	H/09/01997/066	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 100 mg filmsko obložene tablete	DK/H/1527/001	H/09/01997/048	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 200 mg filmsko obložene tablete	DK/H/1527/003	H/09/01997/016	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 200 mg filmsko obložene tablete	DK/H/1527/003	H/09/01997/017	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 200 mg filmsko obložene tablete	DK/H/1527/003	H/09/01997/018	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 200 mg filmsko obložene tablete	DK/H/1527/003	H/09/01997/019	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 200 mg filmsko obložene tablete	DK/H/1527/003	H/09/01997/021	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 200 mg filmsko obložene tablete	DK/H/1527/003	H/09/01997/020	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 200 mg filmsko obložene tablete	DK/H/1527/003	H/09/01997/022	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 200 mg filmsko obložene tablete	DK/H/1527/003	H/09/01997/023	LEK PHARMACEUTICALS	SI

<b>Product Name (in authorisation country)</b>	<b>MRP/DCP Authorisation number</b>	<b>National Authorisation Number</b>	<b>MAH of product in the member state</b>	<b>Member State where product is authorised</b>
obložene tablete			D.D. LJUBLJANA	
Kvelux 200 mg filmsko obložene tablete	DK/H/1527/003	H/09/01997/024	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 200 mg filmsko obložene tablete	DK/H/1527/003	H/09/01997/025	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 200 mg filmsko obložene tablete	DK/H/1527/003	H/09/01997/026	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 200 mg filmsko obložene tablete	DK/H/1527/003	H/09/01997/027	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 200 mg filmsko obložene tablete	DK/H/1527/003	H/09/01997/028	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 200 mg filmsko obložene tablete	DK/H/1527/003	H/09/01997/029	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 200 mg filmsko obložene tablete	DK/H/1527/003	H/09/01997/067	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 200 mg filmsko obložene tablete	DK/H/1527/003	H/09/01997/068	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 200 mg filmsko obložene tablete	DK/H/1527/003	H/09/01997/069	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 200 mg filmsko obložene tablete	DK/H/1527/003	H/09/01997/070	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 200 mg filmsko obložene tablete	DK/H/1527/003	H/09/01997/071	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 200 mg filmsko obložene tablete	DK/H/1527/003	H/09/01997/072	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 200 mg filmsko obložene tablete	DK/H/1527/003	H/09/01997/073	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 200 mg filmsko obložene tablete	DK/H/1527/003	H/09/01997/074	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 200 mg filmsko obložene tablete	DK/H/1527/003	H/09/01997/075	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 200 mg filmsko obložene tablete	DK/H/1527/003	H/09/01997/077	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 200 mg filmsko obložene tablete	DK/H/1527/003	H/09/01997/078	LEK PHARMACEUTICALS	SI



<b>Product Name (in authorisation country)</b>	<b>MRP/DCP Authorisation number</b>	<b>National Authorisation Number</b>	<b>MAH of product in the member state</b>	<b>Member State where product is authorised</b>
obložene tablete			D.D. LJUBLJANA	
Kvelux 200 mg filmsko obložene tablete	DK/H/1527/003	H/09/01997/079	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 200 mg filmsko obložene tablete	DK/H/1527/003	H/09/01997/080	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 200 mg filmsko obložene tablete	DK/H/1527/003	H/09/01997/081	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 200 mg filmsko obložene tablete	DK/H/1527/003	H/09/01997/082	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 200 mg filmsko obložene tablete	DK/H/1527/003	H/09/01997/083	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 200 mg filmsko obložene tablete	DK/H/1527/003	H/09/01997/084	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 200 mg filmsko obložene tablete	DK/H/1527/003	H/09/01997/086	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 200 mg filmsko obložene tablete	DK/H/1527/003	H/09/01997/087	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 200 mg filmsko obložene tablete	DK/H/1527/003	H/09/01997/088	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 200 mg filmsko obložene tablete	DK/H/1527/003	H/09/01997/076	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 200 mg filmsko obložene tablete	DK/H/1527/003	H/09/01997/085	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 300 mg filmsko obložene tablete	DK/H/1527/004	H/09/01997/030	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 300 mg filmsko obložene tablete	DK/H/1527/004	H/09/01997/031	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 300 mg filmsko obložene tablete	DK/H/1527/004	H/09/01997/032	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 300 mg filmsko obložene tablete	DK/H/1527/004	H/09/01997/033	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 300 mg filmsko obložene tablete	DK/H/1527/004	H/09/01997/034	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 300 mg filmsko obložene tablete	DK/H/1527/004	H/09/01997/035	LEK PHARMACEUTICALS	SI

<b>Product Name (in authorisation country)</b>	<b>MRP/DCP Authorisation number</b>	<b>National Authorisation Number</b>	<b>MAH of product in the member state</b>	<b>Member State where product is authorised</b>
obložene tablete			D.D. LJUBLJANA	
Kvelux 300 mg filmsko obložene tablete	DK/H/1527/004	H/09/01997/036	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 300 mg filmsko obložene tablete	DK/H/1527/004	H/09/01997/037	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 300 mg filmsko obložene tablete	DK/H/1527/004	H/09/01997/038	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 300 mg filmsko obložene tablete	DK/H/1527/004	H/09/01997/039	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 300 mg filmsko obložene tablete	DK/H/1527/004	H/09/01997/040	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 300 mg filmsko obložene tablete	DK/H/1527/004	H/09/01997/041	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 300 mg filmsko obložene tablete	DK/H/1527/004	H/09/01997/042	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 300 mg filmsko obložene tablete	DK/H/1527/004	H/09/01997/043	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 300 mg filmsko obložene tablete	DK/H/1527/004	H/09/01997/044	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 300 mg filmsko obložene tablete	DK/H/1527/004	H/09/01997/090	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 300 mg filmsko obložene tablete	DK/H/1527/004	H/09/01997/101	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 300 mg filmsko obložene tablete	DK/H/1527/004	H/09/01997/097	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 300 mg filmsko obložene tablete	DK/H/1527/004	H/09/01997/102	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 300 mg filmsko obložene tablete	DK/H/1527/004	H/09/01997/104	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 300 mg filmsko obložene tablete	DK/H/1527/004	H/09/01997/107	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 300 mg filmsko obložene tablete	DK/H/1527/004	H/09/01997/089	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 300 mg filmsko	DK/H/1527/004	H/09/01997/091	LEK PHARMACEUTICALS	SI

<b>Product Name (in authorisation country)</b>	<b>MRP/DCP Authorisation number</b>	<b>National Authorisation Number</b>	<b>MAH of product in the member state</b>	<b>Member State where product is authorised</b>
obložene tablete			D.D. LJUBLJANA	
Kvelux 300 mg filmsko obložene tablete	DK/H/1527/004	H/09/01997/092	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 300 mg filmsko obložene tablete	DK/H/1527/004	H/09/01997/093	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 300 mg filmsko obložene tablete	DK/H/1527/004	H/09/01997/094	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 300 mg filmsko obložene tablete	DK/H/1527/004	H/09/01997/096	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 300 mg filmsko obložene tablete	DK/H/1527/004	H/09/01997/100	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 300 mg filmsko obložene tablete	DK/H/1527/004	H/09/01997/105	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 300 mg filmsko obložene tablete	DK/H/1527/004	H/09/01997/106	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 300 mg filmsko obložene tablete	DK/H/1527/004	H/09/01997/108	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 300 mg filmsko obložene tablete	DK/H/1527/004	H/09/01997/109	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 300 mg filmsko obložene tablete	DK/H/1527/004	H/09/01997/095	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 300 mg filmsko obložene tablete	DK/H/1527/004	H/09/01997/098	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 300 mg filmsko obložene tablete	DK/H/1527/004	H/09/01997/099	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 300 mg filmsko obložene tablete	DK/H/1527/004	H/09/01997/103	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 300 mg filmsko obložene tablete	DK/H/1527/004	H/09/01997/110	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 50 mg filmsko obložene tablete	DK/H/1430/002	H/09/01933/017	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 50 mg filmsko obložene tablete	DK/H/1430/002	H/09/01933/018	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 50 mg filmsko obložene tablete	DK/H/1430/002	H/09/01933/019	LEK PHARMACEUTICALS	SI

<b>Product Name (in authorisation country)</b>	<b>MRP/DCP Authorisation number</b>	<b>National Authorisation Number</b>	<b>MAH of product in the member state</b>	<b>Member State where product is authorised</b>
obložene tablete			D.D. LJUBLJANA	
Kvelux 50 mg filmsko obložene tablete	DK/H/1430/002	H/09/01933/020	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 50 mg filmsko obložene tablete	DK/H/1430/002	H/09/01933/021	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 50 mg filmsko obložene tablete	DK/H/1430/002	H/09/01933/022	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 50 mg filmsko obložene tablete	DK/H/1430/002	H/09/01933/023	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 50 mg filmsko obložene tablete	DK/H/1430/002	H/09/01933/024	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 50 mg filmsko obložene tablete	DK/H/1430/002	H/09/01933/025	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 50 mg filmsko obložene tablete	DK/H/1430/002	H/09/01933/026	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 50 mg filmsko obložene tablete	DK/H/1430/002	H/09/01933/027	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 50 mg filmsko obložene tablete	DK/H/1430/002	H/09/01933/028	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 50 mg filmsko obložene tablete	DK/H/1430/002	H/09/01933/029	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 50 mg filmsko obložene tablete	DK/H/1430/002	H/09/01933/030	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 50 mg filmsko obložene tablete	DK/H/1430/002	H/09/01933/031	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 50 mg filmsko obložene tablete	DK/H/1430/002	H/09/01933/032	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 50 mg filmsko obložene tablete	DK/H/1430/002	H/09/01933/033	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 50 mg filmsko obložene tablete	DK/H/1430/002	H/09/01933/034	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 50 mg filmsko obložene tablete	DK/H/1430/002	H/09/01933/036	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 50 mg filmsko obložene tablete	DK/H/1430/002	H/09/01933/035	LEK PHARMACEUTICALS	SI

<b>Product Name (in authorisation country)</b>	<b>MRP/DCP Authorisation number</b>	<b>National Authorisation Number</b>	<b>MAH of product in the member state</b>	<b>Member State where product is authorised</b>
obložene tablete			D.D. LJUBLJANA	
Kvelux 50 mg filmsko obložene tablete	DK/H/1430/002	H/09/01933/059	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 50 mg filmsko obložene tablete	DK/H/1430/002	H/09/01933/060	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 50 mg filmsko obložene tablete	DK/H/1430/002	H/09/01933/061	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 50 mg filmsko obložene tablete	DK/H/1430/002	H/09/01933/062	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 50 mg filmsko obložene tablete	DK/H/1430/002	H/09/01933/063	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 50 mg filmsko obložene tablete	DK/H/1430/002	H/09/01933/064	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 50 mg filmsko obložene tablete	DK/H/1430/002	H/09/01933/065	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 50 mg filmsko obložene tablete	DK/H/1430/002	H/09/01933/066	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 50 mg filmsko obložene tablete	DK/H/1430/002	H/09/01933/067	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 50 mg filmsko obložene tablete	DK/H/1430/002	H/09/01933/068	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 50 mg filmsko obložene tablete	DK/H/1430/002	H/09/01933/069	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 50 mg filmsko obložene tablete	DK/H/1430/002	H/09/01933/070	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 50 mg filmsko obložene tablete	DK/H/1430/002	H/09/01933/071	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 50 mg filmsko obložene tablete	DK/H/1430/002	H/09/01933/072	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 50 mg filmsko obložene tablete	DK/H/1430/002	H/09/01933/073	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 50 mg filmsko obložene tablete	DK/H/1430/002	H/09/01933/074	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 50 mg filmsko obložene tablete	DK/H/1430/002	H/09/01933/075	LEK PHARMACEUTICALS	SI

<b>Product Name (in authorisation country)</b>	<b>MRP/DCP Authorisation number</b>	<b>National Authorisation Number</b>	<b>MAH of product in the member state</b>	<b>Member State where product is authorised</b>
obložene tablete			D.D. LJUBLJANA	
Kvelux 50 mg filmsko obložene tablete	DK/H/1430/002	H/09/01933/076	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 50 mg filmsko obložene tablete	DK/H/1430/002	H/09/01933/077	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 50 mg filmsko obložene tablete	DK/H/1430/002	H/09/01933/078	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 50 mg filmsko obložene tablete	DK/H/1430/002	H/09/01933/079	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 50 mg filmsko obložene tablete	DK/H/1430/002	H/09/01933/080	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 50 mg filmsko obložene tablete	DK/H/1430/002	H/09/01933/081	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 50 mg filmsko obložene tablete	DK/H/1430/002	H/09/01933/082	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 50 mg filmsko obložene tablete	DK/H/1430/002	H/09/01933/083	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kvelux 50 mg filmsko obložene tablete	DK/H/1430/002	H/09/01933/084	LEK PHARMACEUTICALS D.D. LJUBLJANA	SI
Kventiax 150 mg comprimate filmate	DK/H/1059/003	5524/2013/13	KRKA, D.D., NOVO MESTO	RO
Kventiax 150 mg comprimate filmate	DK/H/1059/003	5524/2013/01	KRKA, D.D., NOVO MESTO	RO
Kventiax 150 mg comprimate filmate	DK/H/1059/003	5524/2013/02	KRKA, D.D., NOVO MESTO	RO
Kventiax 150 mg comprimate filmate	DK/H/1059/003	5524/2013/03	KRKA, D.D., NOVO MESTO	RO
Kventiax 150 mg comprimate filmate	DK/H/1059/003	5524/2013/04	KRKA, D.D., NOVO MESTO	RO
Kventiax 150 mg comprimate filmate	DK/H/1059/003	5524/2013/05	KRKA, D.D., NOVO MESTO	RO
Kventiax 150 mg comprimate filmate	DK/H/1059/003	5524/2013/06	KRKA, D.D., NOVO MESTO	RO
Kventiax 150 mg	DK/H/1059/003	5524/2013/07	KRKA, D.D., NOVO MESTO	RO

<b>Product Name (in authorisation country)</b>	<b>MRP/DCP Authorisation number</b>	<b>National Authorisation Number</b>	<b>MAH of product in the member state</b>	<b>Member State where product is authorised</b>
comprimatate filmate				
Kventiax 150 mg comprimatate filmate	DK/H/1059/003	5524/2013/08	KRKA, D.D., NOVO MESTO	RO
Kventiax 150 mg comprimatate filmate	DK/H/1059/003	5524/2013/09	KRKA, D.D., NOVO MESTO	RO
Kventiax 150 mg comprimatate filmate	DK/H/1059/003	5524/2013/10	KRKA, D.D., NOVO MESTO	RO
Kventiax 150 mg comprimatate filmate	DK/H/1059/003	5524/2013/11	KRKA, D.D., NOVO MESTO	RO
Kventiax 150 mg comprimatate filmate	DK/H/1059/003	5524/2013/12	KRKA, D.D., NOVO MESTO	RO
Kventiax 150 mg filmom obalené tablety	DK/H/1059/003	68/0266/07-S	KRKA, D.D., NOVO MESTO	SK
Kventiax 200 mg apvalkotās tabletes	DK/H/1059/004	07-0232	KRKA, D.D., NOVO MESTO	LV
Kventiax 25 mg apvalkotās tabletes	DK/H/1059/001	07-0229	KRKA, D.D., NOVO MESTO	LV
Kventiax 25 mg comprimatate filmate	DK/H/1059/001	5522/2013/11	KRKA, D.D., NOVO MESTO	RO
Kventiax 25 mg comprimatate filmate	DK/H/1059/001	5522/2013/01	KRKA, D.D., NOVO MESTO	RO
Kventiax 25 mg comprimatate filmate	DK/H/1059/001	5522/2013/02	KRKA, D.D., NOVO MESTO	RO
Kventiax 25 mg comprimatate filmate	DK/H/1059/001	5522/2013/03	KRKA, D.D., NOVO MESTO	RO
Kventiax 25 mg comprimatate filmate	DK/H/1059/001	5522/2013/04	KRKA, D.D., NOVO MESTO	RO
Kventiax 25 mg comprimatate filmate	DK/H/1059/001	5522/2013/05	KRKA, D.D., NOVO MESTO	RO
Kventiax 25 mg comprimatate filmate	DK/H/1059/001	5522/2013/06	KRKA, D.D., NOVO MESTO	RO
Kventiax 25 mg comprimatate filmate	DK/H/1059/001	5522/2013/07	KRKA, D.D., NOVO MESTO	RO
Kventiax 25 mg	DK/H/1059/001	5522/2013/08	KRKA, D.D., NOVO MESTO	RO

<b>Product Name (in authorisation country)</b>	<b>MRP/DCP Authorisation number</b>	<b>National Authorisation Number</b>	<b>MAH of product in the member state</b>	<b>Member State where product is authorised</b>
comprimate filmate				
Kventiax 25 mg comprimate filmate	DK/H/1059/001	5522/2013/09	KRKA, D.D., NOVO MESTO	RO
Kventiax 25 mg comprimate filmate	DK/H/1059/001	5522/2013/10	KRKA, D.D., NOVO MESTO	RO
Kventiax 300 mg apvalkotās tabletes	DK/H/1059/005	07-0233	KRKA, D.D., NOVO MESTO	LV
Kventiax 300 mg potahované tablety	DK/H/1059/005	68/476/07-C	KRKA, D.D., NOVO MESTO	CZ
Kventiax 300 mg tabletki powlekane	DK/H/1059/005	14113	KRKA, D.D., NOVO MESTO	PL
Kventiax, 25 mg õhukese polümeerikattega tabletid	DK/H/1059/001	558307	KRKA, D.D., NOVO MESTO	EE
Kventiax, 300 mg õhukese polümeerikattega tabletid	DK/H/1059/005	558007	KRKA, D.D., NOVO MESTO	EE
Psicotric 400 mg comprimidos recubiertos con película	not available	84359	NEURAXPHARM SPAIN, S.L.U.	ES
Psicotric 50 mg comprimidos recubiertos con película	not available	80459	NEURAXPHARM SPAIN, S.L.U.	ES
Psicotric Retard 600 mg comprimidos de liberación prolongada	DK/H/3051/001	84969	NEURAXPHARM SPAIN, S.L.U.	ES
Questax Prolong 600 mg tablety s prodlouženým uvolňováním	DK/H/3051/001	68/019/19-C	NEURAXPHARM BOHEMIA S.R.O.	CZ
Questax XR 600 mg retard tableta	DK/H/3051/001	OGYI-T-23570/31	NEURAXPHARM BOHEMIA S.R.O.	HU
Questax XR 600 mg retard tableta	DK/H/3051/001	OGYI-T-23570/32	NEURAXPHARM BOHEMIA S.R.O.	HU



<b>Product Name (in authorisation country)</b>	<b>MRP/DCP Authorisation number</b>	<b>National Authorisation Number</b>	<b>MAH of product in the member state</b>	<b>Member State where product is authorised</b>
Questax XR 600 mg retard tabletta	DK/H/3051/001	OGYI-T-23570/33	NEURAXPHARM BOHEMIA S.R.O.	HU
Questax XR 600 mg retard tabletta	DK/H/3051/001	OGYI-T-23570/34	NEURAXPHARM BOHEMIA S.R.O.	HU
Questax XR 600 mg retard tabletta	DK/H/3051/001	OGYI-T-23570/35	NEURAXPHARM BOHEMIA S.R.O.	HU
Questax XR 600 mg retard tabletta	DK/H/3051/001	OGYI-T-23570/36	NEURAXPHARM BOHEMIA S.R.O.	HU
Questax XR 600 mg tablety s predĺženým uvoľňovaním	DK/H/3051/001	68/033/20-S	NEURAXPHARM BOHEMIA S.R.O.	SK
Questax, 400 mg, tabletki powlekane	PT/H/2158/007	25617	NEURAXPHARM ARZNEIMITTEL GMBH	PL
Questax, 50 mg, tabletki powlekane	PT/H/2158/002	25612	NEURAXPHARM ARZNEIMITTEL GMBH	PL
Quetamed 400 mg comprimidos revestidos por película	PT /H/2158/007	5772702	NEURAXPHARM SPAIN, S.L.U.	PT
Quetamed 50 mg comprimidos revestidos por película	PT /H/2158/002	5772637	NEURAXPHARM SPAIN, S.L.U.	PT
Quetamed 600 mg depottabletter	DK/H/3051/001	62219-2019	NEURAXPHARM ARZNEIMITTEL GMBH	DK
Quetex 100 mg Film-Coated Tablets	DK/H/1528/001	PA0711/156/003	ROWEX LTD	IE
Quetex 200mg Film-Coated Tablets	DK/H/1528/003	PA 0711/156/004	ROWEX LTD	IE
Quetex 300mg Film-Coated Tablets	DK/H/1528/004	PA0711/156/005	ROWEX LTD	IE
Quetiapin - 1 A Pharma 100 mg Filmtabletten	DK/H/1529/001	73497.00.00	1 A PHARMA GMBH	DE
Quetiapin - 1 A Pharma 150 mg Filmtabletten	DK/H/1529/002	73498.00.00	1 A PHARMA GMBH	DE
Quetiapin - 1 A Pharma	DK/H/1529/003	73499.00.00	1 A PHARMA GMBH	DE

<b>Product Name (in authorisation country)</b>	<b>MRP/DCP Authorisation number</b>	<b>National Authorisation Number</b>	<b>MAH of product in the member state</b>	<b>Member State where product is authorised</b>
200 mg Filmtabletten				
Quetiapin - 1 A Pharma 300 mg Filmtabletten	DK/H/1529/004	73500.00.00	1 A PHARMA GMBH	DE
Quetiapin - 1 A Pharma 400 mg Filmtabletten	DK/H/1529/005	73501.00.00	1 A PHARMA GMBH	DE
Quetiapin "1A Farma", filmovertrukne tabletter 100 mg	DK/H/1529/001	43582	1A FARMA A/S	DK
Quetiapin "1A Farma", filmovertrukne tabletter 150 mg	DK/H/1529/002	43583	1A FARMA A/S	DK
Quetiapin "1A Farma", filmovertrukne tabletter 200 mg	DK/H/1529/003	43584	1A FARMA A/S	DK
Quetiapin "1A Farma", filmovertrukne tabletter 300 mg	DK/H/1529/004	43585	1A FARMA A/S	DK
Quetiapin "1A Farma", filmovertrukne tabletter 400 mg	DK/H/1529/005	43586	1A FARMA A/S	DK
Quetiapin "1A Farma", filmovertrukne tabletter 50 mg	DK/H/1432/002	42923	1A FARMA A/S	DK
Quetiapin "Sandoz", filmovertrukne tabletter 100 mg	DK/H/1527/001	43568	SANDOZ A/S	DK
Quetiapin "Sandoz", filmovertrukne tabletter 150 mg	DK/H/1527/002	43569	SANDOZ A/S	DK
Quetiapin "Sandoz", filmovertrukne tabletter 200 mg	DK/H/1527/003	43570	SANDOZ A/S	DK
Quetiapin "Sandoz", filmovertrukne tabletter	DK/H/1527/004	43571	SANDOZ A/S	DK

<b>Product Name (in authorisation country)</b>	<b>MRP/DCP Authorisation number</b>	<b>National Authorisation Number</b>	<b>MAH of product in the member state</b>	<b>Member State where product is authorised</b>
300 mg				
Quetiapin "Sandoz", filmovertrukne tabletter 50 mg	DK/H/1430/002	42919	SANDOZ A/S	DK
Quetiapin 1A Pharma 100 mg - Filmtabletten	DK/H/1529/001	1-29169	1A PHARMA GMBH	AT
Quetiapin 1A Pharma 200 mg - Filmtabletten	DK/H/1529/003	1-29171	1A PHARMA GMBH	AT
Quetiapin 1A Pharma 300 mg - Filmtabletten	DK/H/1529/004	1-29172	1A PHARMA GMBH	AT
Quetiapin 1A Pharma 400 mg - Filmtabletten	DK/H/1529/005	1-29173	1A PHARMA GMBH	AT
Quetiapin 1A Pharma 50 mg - Filmtabletten	DK/H/1432/002	1-28720	1A PHARMA GMBH	AT
Quetiapin Gerot 200 mg-Retardtabletten	not available	1-31088	G.L. PHARMA GMBH	AT
Quetiapin Gerot 300 mg-Retardtabletten	not available	1-31089	G.L. PHARMA GMBH	AT
Quetiapin Gerot 400 mg-Retardtabletten	not available	1-31090	G.L. PHARMA GMBH	AT
Quetiapin Gerot 50 mg-Retardtabletten	not available	1-31085	G.L. PHARMA GMBH	AT
Quetiapin Heumann 150 mg Filmtabletten	DE/H/2610/004	79953.00.00	HEUMANN PHARMA GMBH & CO. GENERICA KG	DE
Quetiapin Heumann 400 mg Filmtabletten	DE/H/2610/007	79956.00.00	HEUMANN PHARMA GMBH & CO. GENERICA KG	DE
Quetiapin Heumann 50 mg Filmtabletten	DE/H/2610/002	79951.00.00	HEUMANN PHARMA GMBH & CO. GENERICA KG	DE
Quetiapin HEXAL 100 mg Filmtabletten	DK/H/1528/001	73558.00.00	HEXAL AG	DE
Quetiapin HEXAL 150 mg Filmtabletten	DK/H/1528/002	73559.00.00	HEXAL AG	DE
Quetiapin HEXAL 200 mg Filmtabletten	DK/H/1528/003	73560.00.00	HEXAL AG	DE

<b>Product Name (in authorisation country)</b>	<b>MRP/DCP Authorisation number</b>	<b>National Authorisation Number</b>	<b>MAH of product in the member state</b>	<b>Member State where product is authorised</b>
Quetiapin HEXAL 300 mg Filmtabletten	DK/H/1528/004	73561.00.00	HEXAL AG	DE
Quetiapin HEXAL 400 mg Filmtabletten	DK/H/1528/005	73562.00.00	HEXAL AG	DE
Quetiapin HEXAL 50 mg Filmtabletten	DK/H/1431/002	71239.00.00	HEXAL AG	DE
Quetiapin Lannacher 200 mg-Retardtabletten	not available	1-31092	G.L. PHARMA GMBH	AT
Quetiapin Lannacher 300 mg-Retardtabletten	not available	1-31096	G.L. PHARMA GMBH	AT
Quetiapin Lannacher 400 mg-Retardtabletten	not available	1-31097	G.L. PHARMA GMBH	AT
Quetiapin Lannacher 50 mg-Retardtabletten	not available	1-31091	G.L. PHARMA GMBH	AT
Quetiapin Neuraxpharm 400 mg filmom obalené tablety	DK/H/2628/007	68/0140/19-S	NEURAXPHARM BOHEMIA S.R.O.	SK
Quetiapin Neuraxpharm 400 mg potahované tablety	PT/H/2158/007	68/092/18-C	NEURAXPHARM BOHEMIA S.R.O.	CZ
Quetiapin Neuraxpharm 50 mg filmom obalené tablety	DK/H/2628/002	68/0135/19-S	NEURAXPHARM BOHEMIA S.R.O.	SK
Quetiapin Neuraxpharm 50 mg potahované tablety	PT/H/2158/002	68/087/18-C	NEURAXPHARM BOHEMIA S.R.O.	CZ
Quetiapin Passauer 400 mg Filmtabletten	PT/H/2158/007	2202160.00.00	NEURAXPHARM ARZNEIMITTEL GMBH	DE
Quetiapin Passauer 50 mg Filmtabletten	PT/H/2158/002	2202155.00.00	NEURAXPHARM ARZNEIMITTEL GMBH	DE
Quetiapin Sandoz 100 mg – Filmtabletten	DK/H/1527/001	1-29174	SANDOZ GMBH	AT
Quetiapin Sandoz 100 mg filmdragerade	DK/H/1527/001	27752	SANDOZ A/S	SE

<b>Product Name (in authorisation country)</b>	<b>MRP/DCP Authorisation number</b>	<b>National Authorisation Number</b>	<b>MAH of product in the member state</b>	<b>Member State where product is authorised</b>
tabletter				
Quetiapin Sandoz 200 mg – Filmtabletten	DK/H/1527/003	1-29176	SANDOZ GMBH	AT
Quetiapin Sandoz 200 mg filmdragerade tabletter	DK/H/1527/003	27754	SANDOZ A/S	SE
Quetiapin Sandoz 300 mg – Filmtabletten	DK/H/1527/004	1-29177	SANDOZ GMBH	AT
Quetiapin Sandoz 50 mg – Filmtabletten	DK/H/1430/002	1-28698	SANDOZ GMBH	AT
Quetiapin TAD® 150 mg Filmtabletten	DK/H/1059/003	78988.00.00	TAD PHARMA GMBH	DE
Quetiapina Qualigen 400 mg comprimidos recubiertos con película	not available	84364	NEURAXPHARM SPAIN, S.L.U.	ES
Quetiapina Qualigen 50 mg comprimidos recubiertos con película	not available	84363	NEURAXPHARM SPAIN, S.L.U.	ES
Quetiapina Qualigen Medica 50 mg comprimidos recubiertos con película	not available	84.366	NEURAXPHARM SPAIN, S.L.U.	ES
QUETIAPINA SANDOZ GMBH 100 mg compresse rivestite con film	DK/H/1527/001	040968012	SANDOZ GMBH	IT
QUETIAPINA SANDOZ GMBH 100 mg compresse rivestite con film	DK/H/1527/001	040968048	SANDOZ GMBH	IT
QUETIAPINA SANDOZ GMBH 100 mg compresse rivestite con film	DK/H/1527/001	040968075	SANDOZ GMBH	IT

<b>Product Name (in authorisation country)</b>	<b>MRP/DCP Authorisation number</b>	<b>National Authorisation Number</b>	<b>MAH of product in the member state</b>	<b>Member State where product is authorised</b>
QUETIAPINA SANDOZ GMBH 100 mg compresse rivestite con film	DK/H/1527/001	040968099	SANDOZ GMBH	IT
QUETIAPINA SANDOZ GMBH 100 mg compresse rivestite con film	DK/H/1527/001	040968101	SANDOZ GMBH	IT
QUETIAPINA SANDOZ GMBH 100 mg compresse rivestite con film	DK/H/1527/001	040968024	SANDOZ GMBH	IT
QUETIAPINA SANDOZ GMBH 100 mg compresse rivestite con film	DK/H/1527/001	040968051	SANDOZ GMBH	IT
QUETIAPINA SANDOZ GMBH 100 mg compresse rivestite con film	DK/H/1527/001	040968063	SANDOZ GMBH	IT
QUETIAPINA SANDOZ GMBH 100 mg compresse rivestite con film	DK/H/1527/001	040968087	SANDOZ GMBH	IT
QUETIAPINA SANDOZ GMBH 100 mg compresse rivestite con film	DK/H/1527/001	040968113	SANDOZ GMBH	IT
QUETIAPINA SANDOZ GMBH 100 mg compresse rivestite con film	DK/H/1527/001	040968125	SANDOZ GMBH	IT
QUETIAPINA SANDOZ GMBH 100 mg	DK/H/1527/001	040968137	SANDOZ GMBH	IT

<b>Product Name (in authorisation country)</b>	<b>MRP/DCP Authorisation number</b>	<b>National Authorisation Number</b>	<b>MAH of product in the member state</b>	<b>Member State where product is authorised</b>
compresse rivestite con film				
QUETIAPINA SANDOZ GMBH 100 mg compresse rivestite con film	DK/H/1527/001	040968036	SANDOZ GMBH	IT
QUETIAPINA SANDOZ GMBH 200 mg compresse rivestite con film	DK/H/1527/003	040968152	SANDOZ GMBH	IT
QUETIAPINA SANDOZ GMBH 200 mg compresse rivestite con film	DK/H/1527/003	040968176	SANDOZ GMBH	IT
QUETIAPINA SANDOZ GMBH 200 mg compresse rivestite con film	DK/H/1527/003	040968190	SANDOZ GMBH	IT
QUETIAPINA SANDOZ GMBH 200 mg compresse rivestite con film	DK/H/1527/003	040968226	SANDOZ GMBH	IT
QUETIAPINA SANDOZ GMBH 200 mg compresse rivestite con film	DK/H/1527/003	040968240	SANDOZ GMBH	IT
QUETIAPINA SANDOZ GMBH 200 mg compresse rivestite con film	DK/H/1527/003	040968253	SANDOZ GMBH	IT
QUETIAPINA SANDOZ GMBH 200 mg compresse rivestite con film	DK/H/1527/003	040968149	SANDOZ GMBH	IT

<b>Product Name (in authorisation country)</b>	<b>MRP/DCP Authorisation number</b>	<b>National Authorisation Number</b>	<b>MAH of product in the member state</b>	<b>Member State where product is authorised</b>
QUETIAPINA SANDOZ GMBH 200 mg compresse rivestite con film	DK/H/1527/003	040968164	SANDOZ GMBH	IT
QUETIAPINA SANDOZ GMBH 200 mg compresse rivestite con film	DK/H/1527/003	040968188	SANDOZ GMBH	IT
QUETIAPINA SANDOZ GMBH 200 mg compresse rivestite con film	DK/H/1527/003	040968238	SANDOZ GMBH	IT
QUETIAPINA SANDOZ GMBH 200 mg compresse rivestite con film	DK/H/1527/003	040968202	SANDOZ GMBH	IT
QUETIAPINA SANDOZ GMBH 200 mg compresse rivestite con film	DK/H/1527/003	040968214	SANDOZ GMBH	IT
QUETIAPINA SANDOZ GMBH 200 mg compresse rivestite con film	DK/H/1527/003	040968265	SANDOZ GMBH	IT
QUETIAPINA SANDOZ GMBH 300 mg compresse rivestite con film	DK/H/1527/004	040968315	SANDOZ GMBH	IT
QUETIAPINA SANDOZ GMBH 300 mg compresse rivestite con film	DK/H/1527/004	040968327	SANDOZ GMBH	IT
QUETIAPINA SANDOZ GMBH 300 mg	DK/H/1527/004	040968339	SANDOZ GMBH	IT



<b>Product Name (in authorisation country)</b>	<b>MRP/DCP Authorisation number</b>	<b>National Authorisation Number</b>	<b>MAH of product in the member state</b>	<b>Member State where product is authorised</b>
compresse rivestite con film				
QUETIAPINA SANDOZ GMBH 300 mg compresse rivestite con film	DK/H/1527/004	040968378	SANDOZ GMBH	IT
QUETIAPINA SANDOZ GMBH 300 mg compresse rivestite con film	DK/H/1527/004	040968277	SANDOZ GMBH	IT
QUETIAPINA SANDOZ GMBH 300 mg compresse rivestite con film	DK/H/1527/004	040968291	SANDOZ GMBH	IT
QUETIAPINA SANDOZ GMBH 300 mg compresse rivestite con film	DK/H/1527/004	040968303	SANDOZ GMBH	IT
QUETIAPINA SANDOZ GMBH 300 mg compresse rivestite con film	DK/H/1527/004	040968341	SANDOZ GMBH	IT
QUETIAPINA SANDOZ GMBH 300 mg compresse rivestite con film	DK/H/1527/004	040968354	SANDOZ GMBH	IT
QUETIAPINA SANDOZ GMBH 300 mg compresse rivestite con film	DK/H/1527/004	040968366	SANDOZ GMBH	IT
QUETIAPINA SANDOZ GMBH 300 mg compresse rivestite con film	DK/H/1527/004	040968380	SANDOZ GMBH	IT

<b>Product Name (in authorisation country)</b>	<b>MRP/DCP Authorisation number</b>	<b>National Authorisation Number</b>	<b>MAH of product in the member state</b>	<b>Member State where product is authorised</b>
QUETIAPINA SANDOZ GMBH 300 mg compresse rivestite con film	DK/H/1527/004	040968392	SANDOZ GMBH	IT
QUETIAPINA SANDOZ GMBH 300 mg compresse rivestite con film	DK/H/1527/004	040968289	SANDOZ GMBH	IT
Quetiapine 100 mg film-coated tablets	not available	PL 20416/0601	CRESCENT PHARMA LIMITED	UK
Quetiapine 100 mg Film-coated Tablets	DK/H/1527/001	PL 04416/0962	SANDOZ LTD	UK
Quetiapine 100 mg Film-coated Tablets	not available	PL 30322/0035	ALISSA HEALTHCARE RESEARCH LIMITED	UK
Quetiapine 150 mg film-coated tablets	PT/H/0604/003	MA807/03008	AUROBINDO PHARMA (MALTA) LIMITED	MT
Quetiapine 150 mg film-coated tablets	not available	PL 20416/0602	CRESCENT PHARMA LIMITED	UK
Quetiapine 150 mg Film-coated Tablets	DK/H/1527/002	PL 04416/0963	SANDOZ LTD	UK
Quetiapine 150 mg Film-coated Tablets	not available	PL 30322/0036	ALISSA HEALTHCARE RESEARCH LIMITED	UK
Quetiapine 200 mg film-coated tablets	not available	PL 20416/0603	CRESCENT PHARMA LIMITED	UK
Quetiapine 200 mg Film-coated Tablets	DK/H/1527/003	PL 04416/0964	SANDOZ LTD	UK
Quetiapine 200 mg Film-coated Tablets	not available	PL 30322/0037	ALISSA HEALTHCARE RESEARCH LIMITED	UK
Quetiapine 25 mg film-coated tablets	not available	PL 20416/0600	CRESCENT PHARMA LIMITED	UK
Quetiapine 25 mg Film-coated Tablets	not available	PL 30322/0034	ALISSA HEALTHCARE RESEARCH LIMITED	UK
Quetiapine 300 mg film-coated tablets	PT/H/0604/005	MA807/03010	AUROBINDO PHARMA (MALTA) LIMITED	MT

<b>Product Name (in authorisation country)</b>	<b>MRP/DCP Authorisation number</b>	<b>National Authorisation Number</b>	<b>MAH of product in the member state</b>	<b>Member State where product is authorised</b>
Quetiapine 300 mg film-coated tablets	not available	PL 20416/0604	CRESCENT PHARMA LIMITED	UK
Quetiapine 300 mg Film-coated Tablets	DK/H/1527/004	PL 04416/0965	SANDOZ LTD	UK
Quetiapine 300 mg Film-coated Tablets	not available	PL 30322/0038	ALISSA HEALTHCARE RESEARCH LIMITED	UK
Quetiapine Accord 150 mg comprimate filmate	NL/H/4782/003	13243/2020/07	ACCORD HEALTHCARE POLSKA SP. Z O.O.	RO
Quetiapine Accord 150 mg comprimate filmate	NL/H/4782/003	13243/2020/06	ACCORD HEALTHCARE POLSKA SP. Z O.O.	RO
Quetiapine Accord 150 mg comprimate filmate	NL/H/4782/003	13243/2020/05	ACCORD HEALTHCARE POLSKA SP. Z O.O.	RO
Quetiapine Accord 150 mg comprimate filmate	NL/H/4782/003	13243/2020/04	ACCORD HEALTHCARE POLSKA SP. Z O.O.	RO
Quetiapine Accord 150 mg comprimate filmate	NL/H/4782/003	13243/2020/03	ACCORD HEALTHCARE POLSKA SP. Z O.O.	RO
Quetiapine Accord 150 mg comprimate filmate	NL/H/4782/003	13243/2020/02	ACCORD HEALTHCARE POLSKA SP. Z O.O.	RO
Quetiapine Accord 150 mg comprimate filmate	NL/H/4782/003	13243/2020/12	ACCORD HEALTHCARE POLSKA SP. Z O.O.	RO
Quetiapine Accord 150 mg comprimate filmate	NL/H/4782/003	13243/2020/11	ACCORD HEALTHCARE POLSKA SP. Z O.O.	RO
Quetiapine Accord 150 mg comprimate filmate	NL/H/4782/003	13243/2020/10	ACCORD HEALTHCARE POLSKA SP. Z O.O.	RO
Quetiapine Accord 150 mg comprimate filmate	NL/H/4782/003	13243/2020/09	ACCORD HEALTHCARE POLSKA SP. Z O.O.	RO
Quetiapine Accord 150 mg comprimate filmate	NL/H/4782/003	13243/2020/08	ACCORD HEALTHCARE POLSKA SP. Z O.O.	RO
Quetiapine Accord 150 mg comprimate filmate	NL/H/4782/003	13243/2020/14	ACCORD HEALTHCARE POLSKA SP. Z O.O.	RO
Quetiapine Accord 150 mg comprimate filmate	NL/H/4782/003	13243/2020/01	ACCORD HEALTHCARE POLSKA SP. Z O.O.	RO
Quetiapine Accord 150 mg comprimate filmate	NL/H/4782/003	13243/2020/13	ACCORD HEALTHCARE POLSKA SP. Z O.O.	RO

<b>Product Name (in authorisation country)</b>	<b>MRP/DCP Authorisation number</b>	<b>National Authorisation Number</b>	<b>MAH of product in the member state</b>	<b>Member State where product is authorised</b>
Quetiapine Accord 150 mg filmom obalené tablety	NL/H/4782/003	68/0051/11-S	ACCORD HEALTHCARE POLSKA SP. Z O.O.	SK
Quetiapine Accord 150 mg plèvele dengtos tabletés	NL/H/4782/003	LT/1/11/2387/030	ACCORD HEALTHCARE B.V.	LT
Quetiapine Accord 150 mg plèvele dengtos tabletés	NL/H/4782/003	LT/1/11/2387/031	ACCORD HEALTHCARE B.V.	LT
Quetiapine Accord 150 mg plèvele dengtos tabletés	NL/H/4782/003	LT/1/11/2387/032	ACCORD HEALTHCARE B.V.	LT
Quetiapine Accord 150 mg plèvele dengtos tabletés	NL/H/4782/003	LT/1/11/2387/033	ACCORD HEALTHCARE B.V.	LT
Quetiapine Accord 150 mg plèvele dengtos tabletés	NL/H/4782/003	LT/1/11/2387/034	ACCORD HEALTHCARE B.V.	LT
Quetiapine Accord 150 mg plèvele dengtos tabletés	NL/H/4782/003	LT/1/11/2387/035	ACCORD HEALTHCARE B.V.	LT
Quetiapine Accord 150 mg plèvele dengtos tabletés	NL/H/4782/003	LT/1/11/2387/036	ACCORD HEALTHCARE B.V.	LT
Quetiapine Accord 150 mg plèvele dengtos tabletés	NL/H/4782/003	LT/1/11/2387/037	ACCORD HEALTHCARE B.V.	LT
Quetiapine Accord 150 mg plèvele dengtos tabletés	NL/H/4782/003	LT/1/11/2387/038	ACCORD HEALTHCARE B.V.	LT
Quetiapine Accord 150 mg plèvele dengtos tabletés	NL/H/4782/003	LT/1/11/2387/039	ACCORD HEALTHCARE B.V.	LT
Quetiapine Accord 150	NL/H/4782/003	LT/1/11/2387/040	ACCORD HEALTHCARE B.V.	LT

<b>Product Name (in authorisation country)</b>	<b>MRP/DCP Authorisation number</b>	<b>National Authorisation Number</b>	<b>MAH of product in the member state</b>	<b>Member State where product is authorised</b>
mg plèvele dengtos tabletés				
Quetiapine Accord 150 mg plèvele dengtos tabletés	NL/H/4782/003	LT/1/11/2387/041	ACCORD HEALTHCARE B.V.	LT
Quetiapine Accord 150 mg plèvele dengtos tabletés	NL/H/4782/003	LT/1/11/2387/042	ACCORD HEALTHCARE B.V.	LT
Quetiapine Accord 150 mg plèvele dengtos tabletés	NL/H/4782/003	LT/1/11/2387/030	ACCORD HEALTHCARE B.V.	LT
Quetiapine Accord 150 mg plèvele dengtos tabletés	NL/H/4782/003	LT/1/11/2387/031	ACCORD HEALTHCARE B.V.	LT
Quetiapine Accord 150 mg plèvele dengtos tabletés	NL/H/4782/003	LT/1/11/2387/032	ACCORD HEALTHCARE B.V.	LT
Quetiapine Accord 150 mg plèvele dengtos tabletés	NL/H/4782/003	LT/1/11/2387/033	ACCORD HEALTHCARE B.V.	LT
Quetiapine Accord 150 mg plèvele dengtos tabletés	NL/H/4782/003	LT/1/11/2387/034	ACCORD HEALTHCARE B.V.	LT
Quetiapine Accord 150 mg plèvele dengtos tabletés	NL/H/4782/003	LT/1/11/2387/035	ACCORD HEALTHCARE B.V.	LT
Quetiapine Accord 150 mg plèvele dengtos tabletés	NL/H/4782/003	LT/1/11/2387/036	ACCORD HEALTHCARE B.V.	LT
Quetiapine Accord 150 mg plèvele dengtos tabletés	NL/H/4782/003	LT/1/11/2387/037	ACCORD HEALTHCARE B.V.	LT
Quetiapine Accord 150 mg plèvele dengtos	NL/H/4782/003	LT/1/11/2387/038	ACCORD HEALTHCARE B.V.	LT

<b>Product Name (in authorisation country)</b>	<b>MRP/DCP Authorisation number</b>	<b>National Authorisation Number</b>	<b>MAH of product in the member state</b>	<b>Member State where product is authorised</b>
tabletés				
Quetiapine Accord 150 mg plèvele dengtos tabletés	NL/H/4782/003	LT/1/11/2387/039	ACCORD HEALTHCARE B.V.	LT
Quetiapine Accord 150 mg plèvele dengtos tabletés	NL/H/4782/003	LT/1/11/2387/040	ACCORD HEALTHCARE B.V.	LT
Quetiapine Accord 150 mg plèvele dengtos tabletés	NL/H/4782/003	LT/1/11/2387/041	ACCORD HEALTHCARE B.V.	LT
Quetiapine Accord 150 mg plèvele dengtos tabletés	NL/H/4782/003	LT/1/11/2387/042	ACCORD HEALTHCARE B.V.	LT
Quetiapine Accord 150 mg plèvele dengtos tabletés	NL/H/4782/003	LT/1/11/2387/029	ACCORD HEALTHCARE B.V.	LT
Quetiapine Accord 150 mg plèvele dengtos tabletés	NL/H/4782/003	LT/1/11/2387/029	ACCORD HEALTHCARE B.V.	LT
Quetiapine Accord 25 mg comprimate filmate	NL/H/4782/001	13241/2020/10	ACCORD HEALTHCARE POLSKA SP. Z O.O.	RO
Quetiapine Accord 25 mg comprimate filmate	NL/H/4782/001	13241/2020/12	ACCORD HEALTHCARE POLSKA SP. Z O.O.	RO
Quetiapine Accord 25 mg comprimate filmate	NL/H/4782/001	13241/2020/14	ACCORD HEALTHCARE POLSKA SP. Z O.O.	RO
Quetiapine Accord 25 mg comprimate filmate	NL/H/4782/001	13241/2020/08	ACCORD HEALTHCARE POLSKA SP. Z O.O.	RO
Quetiapine Accord 25 mg comprimate filmate	NL/H/4782/001	13241/2020/09	ACCORD HEALTHCARE POLSKA SP. Z O.O.	RO
Quetiapine Accord 25 mg comprimate filmate	NL/H/4782/001	13241/2020/07	ACCORD HEALTHCARE POLSKA SP. Z O.O.	RO
Quetiapine Accord 25 mg comprimate filmate	NL/H/4782/001	13241/2020/06	ACCORD HEALTHCARE POLSKA SP. Z O.O.	RO
Quetiapine Accord 25 mg	NL/H/4782/001	13241/2020/04	ACCORD HEALTHCARE	RO

<b>Product Name (in authorisation country)</b>	<b>MRP/DCP Authorisation number</b>	<b>National Authorisation Number</b>	<b>MAH of product in the member state</b>	<b>Member State where product is authorised</b>
comprimate filmate			POLSKA SP. Z O.O.	
Quetiapine Accord 25 mg comprimate filmate	NL/H/4782/001	13241/2020/05	ACCORD HEALTHCARE POLSKA SP. Z O.O.	RO
Quetiapine Accord 25 mg comprimate filmate	NL/H/4782/001	13241/2020/03	ACCORD HEALTHCARE POLSKA SP. Z O.O.	RO
Quetiapine Accord 25 mg comprimate filmate	NL/H/4782/001	13241/2020/02	ACCORD HEALTHCARE POLSKA SP. Z O.O.	RO
Quetiapine Accord 25 mg comprimate filmate	NL/H/4782/001	13241/2020/01	ACCORD HEALTHCARE POLSKA SP. Z O.O.	RO
Quetiapine Accord 25 mg comprimate filmate	NL/H/4782/001	13241/2020/11	ACCORD HEALTHCARE POLSKA SP. Z O.O.	RO
Quetiapine Accord 25 mg comprimate filmate	NL/H/4782/001	13241/2020/13	ACCORD HEALTHCARE POLSKA SP. Z O.O.	RO
Quetiapine Accord 300 mg plevele dengtos tabletes	NL/H/4782/005	LT/1/11/2387/068	ACCORD HEALTHCARE B.V.	LT
Quetiapine Accord 300 mg plevele dengtos tabletes	NL/H/4782/005	LT/1/11/2387/069	ACCORD HEALTHCARE B.V.	LT
Quetiapine Accord 300 mg plevele dengtos tabletes	NL/H/4782/005	LT/1/11/2387/070	ACCORD HEALTHCARE B.V.	LT
Quetiapine Accord 300 mg plevele dengtos tabletes	NL/H/4782/005	LT/1/11/2387/057	ACCORD HEALTHCARE B.V.	LT
Quetiapine Accord 300 mg plèvele dengtos tabletés	NL/H/4782/005	LT/1/11/2387/058	ACCORD HEALTHCARE B.V.	LT
Quetiapine Accord 300 mg plèvele dengtos tabletés	NL/H/4782/005	LT/1/11/2387/059	ACCORD HEALTHCARE B.V.	LT
Quetiapine Accord 300 mg plèvele dengtos tabletés	NL/H/4782/005	LT/1/11/2387/060	ACCORD HEALTHCARE B.V.	LT

<b>Product Name (in authorisation country)</b>	<b>MRP/DCP Authorisation number</b>	<b>National Authorisation Number</b>	<b>MAH of product in the member state</b>	<b>Member State where product is authorised</b>
Quetiapine Accord 300 mg plèvele dengtos tabletés	NL/H/4782/005	LT/1/11/2387/061	ACCORD HEALTHCARE B.V.	LT
Quetiapine Accord 300 mg plèvele dengtos tabletés	NL/H/4782/005	LT/1/11/2387/062	ACCORD HEALTHCARE B.V.	LT
Quetiapine Accord 300 mg plèvele dengtos tabletés	NL/H/4782/005	LT/1/11/2387/063	ACCORD HEALTHCARE B.V.	LT
Quetiapine Accord 300 mg plèvele dengtos tabletés	NL/H/4782/005	LT/1/11/2387/064	ACCORD HEALTHCARE B.V.	LT
Quetiapine Accord 300 mg plèvele dengtos tabletés	NL/H/4782/005	LT/1/11/2387/065	ACCORD HEALTHCARE B.V.	LT
Quetiapine Accord 300 mg plèvele dengtos tabletés	NL/H/4782/005	LT/1/11/2387/066	ACCORD HEALTHCARE B.V.	LT
Quetiapine Accord 300 mg plèvele dengtos tabletés	NL/H/4782/005	LT/1/11/2387/067	ACCORD HEALTHCARE B.V.	LT
Quetiapine Accord 300 mg plèvele dengtos tabletés	NL/H/4782/005	LT/1/11/2387/068	ACCORD HEALTHCARE B.V.	LT
Quetiapine Accord 300 mg plèvele dengtos tabletés	NL/H/4782/005	LT/1/11/2387/069	ACCORD HEALTHCARE B.V.	LT
Quetiapine Accord 300 mg plèvele dengtos tabletés	NL/H/4782/005	LT/1/11/2387/070	ACCORD HEALTHCARE B.V.	LT
Quetiapine Accord 300 mg plèvele dengtos tabletés	NL/H/4782/005	LT/1/11/2387/058	ACCORD HEALTHCARE B.V.	LT
Quetiapine Accord 300	NL/H/4782/005	LT/1/11/2387/059	ACCORD HEALTHCARE B.V.	LT



<b>Product Name (in authorisation country)</b>	<b>MRP/DCP Authorisation number</b>	<b>National Authorisation Number</b>	<b>MAH of product in the member state</b>	<b>Member State where product is authorised</b>
mg plèvele dengtos tabletés				
Quetiapine Accord 300 mg plèvele dengtos tabletés	NL/H/4782/005	LT/1/11/2387/060	ACCORD HEALTHCARE B.V.	LT
Quetiapine Accord 300 mg plèvele dengtos tabletés	NL/H/4782/005	LT/1/11/2387/061	ACCORD HEALTHCARE B.V.	LT
Quetiapine Accord 300 mg plèvele dengtos tabletés	NL/H/4782/005	LT/1/11/2387/062	ACCORD HEALTHCARE B.V.	LT
Quetiapine Accord 300 mg plèvele dengtos tabletés	NL/H/4782/005	LT/1/11/2387/063	ACCORD HEALTHCARE B.V.	LT
Quetiapine Accord 300 mg plèvele dengtos tabletés	NL/H/4782/005	LT/1/11/2387/064	ACCORD HEALTHCARE B.V.	LT
Quetiapine Accord 300 mg plèvele dengtos tabletés	NL/H/4782/005	LT/1/11/2387/065	ACCORD HEALTHCARE B.V.	LT
Quetiapine Accord 300 mg plèvele dengtos tabletés	NL/H/4782/005	LT/1/11/2387/066	ACCORD HEALTHCARE B.V.	LT
Quetiapine Accord 300 mg plèvele dengtos tabletés	NL/H/4782/005	LT/1/11/2387/067	ACCORD HEALTHCARE B.V.	LT
Quetiapine Accord 300 mg plèvele dengtos tabletés	NL/H/4782/005	LT/1/11/2387/057	ACCORD HEALTHCARE B.V.	LT
Quetiapine Dawa 100 mg film-coated tablets	not available	PL 30684/0287	DAWA LIMITED	UK
Quetiapine Dawa 150 mg film-coated tablets	not available	PL 30684/0288	DAWA LIMITED	UK
Quetiapine Dawa 200 mg	not available	PL 30684/0289	DAWA LIMITED	UK

<b>Product Name (in authorisation country)</b>	<b>MRP/DCP Authorisation number</b>	<b>National Authorisation Number</b>	<b>MAH of product in the member state</b>	<b>Member State where product is authorised</b>
film-coated tablets				
Quetiapine Dawa 25 mg film-coated tablets	not available	PL 30684/0286	DAWA LIMITED	UK
Quetiapine Dawa 300 mg film-coated tablets	not available	PL 30684/0290	DAWA LIMITED	UK
Quetiapine Krka 150 mg filmomhulde tabletten	DK/H/1059/003	BE370596	KRKA, D.D., NOVO MESTO	BE
Quetiapine Neuraxpharm 50 mg filmomhulde tabletten	NL/H/1616/006	RVG 110369	NEURAXPHARM ARZNEIMITTEL GMBH	NL
Quetiapine Passauer 400 mg filmtabletta	PT/H/2158/007	OGYI-T-23543/08	NEURAXPHARM BOHEMIA S.R.O.	HU
Quetiapine Passauer 50 mg filmtabletta	PT/H/2158/002	OGYI-T-23543/03	NEURAXPHARM BOHEMIA S.R.O.	HU
Quetiapine Rosemont 20mg/ml Oral Suspension	UK/H/5869/001	PA312/28/5	ROSEMONT PHARMACEUTICALS LIMITED	IE
Quetiapine Rosemont 20mg/ml Oral Suspension	IE/H/1009/001	PL 00427/0240	ROSEMONT PHARMACEUTICALS LIMITED	UK
QUETIAPINE SANDOZ 100 MG, FILMOMHULDE TABLETTEN	DK/H/1527/001	RVG 102974	SANDOZ B.V.	NL
Quetiapine Sandoz 200 mg, filmomhulde tabletten	DK/H/1527/003	RVG 102992	SANDOZ B.V.	NL
QUETIAPINE SANDOZ 300 MG, FILMOMHULDE TABLETTEN	DK/H/1527/004	RVG 102993	SANDOZ B.V.	NL
Quetiapine Teva 200 mg apvalkotās tabletes	DE/H/5984/004	08-0340	TEVA PHARMA B.V.	LV
Quetiapine Teva 25 mg apvalkotās tabletes	DE/H/5984/001	08-0342	TEVA PHARMA B.V.	LV
Quetiapine XR	NL/H/1983/002	RVG 106978	ASTRAZENECA BV	NL

<b>Product Name (in authorisation country)</b>	<b>MRP/DCP Authorisation number</b>	<b>National Authorisation Number</b>	<b>MAH of product in the member state</b>	<b>Member State where product is authorised</b>
AstraZeneca 150 mg, tabletten met verlengde afgifte				
Quetiapine XR AstraZeneca 200 mg, tabletten met verlengde afgifte	NL/H/1983/003	RVG 106979	ASTRAZENECA BV	NL
Quetiapine XR AstraZeneca 300 mg, tabletten met verlengde afgifte	NL/H/1983/004	RVG 106980	ASTRAZENECA BV	NL
Quetiapine XR AstraZeneca 400 mg, tabletten met verlengde afgifte	NL/H/1983/005	RVG 106981	ASTRAZENECA BV	NL
Quetiapine XR AstraZeneca 50 mg, tabletten met verlengde afgifte	NL/H/1983/001	RVG 106971	ASTRAZENECA BV	NL
Quetiapine/Rosemont 20mg/ml πόσιμο εναιώρημα	IE/H/1009/1/DC	8988 / 02-02/2017	ROSEMONT PHARMACEUTICALS LIMITED	GR
Quetiapin-neuraxpharm 50 mg Filmtabletten	NL/H/1616/006	85926.00.00	NEURAXPHARM ARZNEIMITTEL GMBH	DE
Quetiapin-neuraxpharm 600 mg Retardtabletten	DE/H/4123/001	2203803.00.00	NEURAXPHARM ARZNEIMITTEL GMBH	DE
Seresano	DK/H/1431/002	42921	HEXAL A/S	DK
Seresano, filmovertrukne tabletter	DK/H/1528/001	43573	HEXAL A/S	DK
Seresano, filmovertrukne tabletter 150 mg	DK/H/1528/002	43574	HEXAL A/S	DK
Seresano, filmovertrukne tabletter 200 mg	DK/H/1528/003	43575	HEXAL A/S	DK
Seresano, filmovertrukne	DK/H/1528/004	43576	HEXAL A/S	DK

<b>Product Name (in authorisation country)</b>	<b>MRP/DCP Authorisation number</b>	<b>National Authorisation Number</b>	<b>MAH of product in the member state</b>	<b>Member State where product is authorised</b>
tabletter 300 mg				
Seresano, filmovetrukne tabletter 400 mg	DK/H/1528/005	43577	HEXAL A/S	DK
Seroquel	NL/H/0156/003	32336	ASTRAZENECA A/S	DK
Seroquel	NL/H/0156/002	32335	ASTRAZENECA A/S	DK
Seroquel	NL/H/0156/001	32334	ASTRAZENECA A/S	DK
Seroquel	NL/H/0156/007	32600	ASTRAZENECA A/S	DK
Seroquel	NL/H/0156/006	32599	ASTRAZENECA A/S	DK
SEROQUEL 100 mg compresse rivestite con film	NL/H/0156/002	A.I.C . 32944035	ASTRAZENECA S.P.A.	IT
SEROQUEL 100 mg compresse rivestite con film	NL/H/0156/002	A.I.C 32944023	ASTRAZENECA S.P.A.	IT
SEROQUEL 100 mg compresse rivestite con film	NL/H/0156/002	A.I.C . 032944213	ASTRAZENECA S.P.A.	IT
SEROQUEL 100 mg compresse rivestite con film	NL/H/0156/002	A.I.C . 032944249	ASTRAZENECA S.P.A.	IT
SEROQUEL 100 mg compresse rivestite con film	NL/H/0156/002	A.I.C . 032944237	ASTRAZENECA S.P.A.	IT
SEROQUEL 100 mg compresse rivestite con film	NL/H/0156/002	A.I.C . 032944252	ASTRAZENECA S.P.A.	IT
SEROQUEL 100 mg compresse rivestite con film	NL/H/0156/002	A.I.C . 032944225	ASTRAZENECA S.P.A.	IT
Seroquel 100 mg comprimidos recubiertos con película	NL/H/0156/002	63.055	ASTRAZENECA FARMACÉUTICA SPAIN, S.A.	ES
Seroquel 100 mg comprimidos revestidos	NL/H/0156/002	3074382	ASTRAZENECA PRODUTOS FARMACEUTICOS LDA	PT

<b>Product Name (in authorisation country)</b>	<b>MRP/DCP Authorisation number</b>	<b>National Authorisation Number</b>	<b>MAH of product in the member state</b>	<b>Member State where product is authorised</b>
por película				
Seroquel 100 mg comprimidos revestidos por película	NL/H/0156/002	3074580	ASTRAZENECA PRODUTOS FARMACEUTICOS LDA	PT
Seroquel 100 mg comprimidos revestidos por película	NL/H/0156/002	3074481	ASTRAZENECA PRODUTOS FARMACEUTICOS LDA	PT
Seroquel 100 mg comprimidos revestidos por película	NL/H/0156/002	3074283	ASTRAZENECA PRODUTOS FARMACEUTICOS LDA	PT
Seroquel 100 mg comprimidos revestidos por película	NL/H/0156/002	3074184	ASTRAZENECA PRODUTOS FARMACEUTICOS LDA	PT
Seroquel 100 mg comprimidos revestidos por película	NL/H/0156/002	3074689	ASTRAZENECA PRODUTOS FARMACEUTICOS LDA	PT
Seroquel 100 mg comprimidos revestidos por película	NL/H/0156/002	5575980	ASTRAZENECA PRODUTOS FARMACEUTICOS LDA	PT
Seroquel 100 mg comprimidos revestidos por película	NL/H/0156/002	3074788	ASTRAZENECA PRODUTOS FARMACEUTICOS LDA	PT
Seroquel 100 mg film-coated tablets	NL/H/0156/002	PA 1019/019/002	ASTRAZENECA AB	IE
Seroquel 100 mg film-coated tablets	NL/H/0156/002	MA 046/02302	ASTRAZENECA AB	MT
Seroquel 100 mg film-coated tablets	NL/H/0156/002	PL 50827/0003	LUYE PHARMA LIMITED	UK
Seroquel 100 mg filmdragerade tabletter	NL/H/0156/002	18959	ASTRAZENECA AB	SE
Seroquel 100 mg filmdrasjerte tabletter	NL/H/0156/002	02-1436	ASTRAZENECA AS	NO
SEROQUEL 100 mg filmom obložene tablete	NL/H/0156/002	HR-H-461808155	ASTRAZENECA D.O.O.	HR

<b>Product Name (in authorisation country)</b>	<b>MRP/DCP Authorisation number</b>	<b>National Authorisation Number</b>	<b>MAH of product in the member state</b>	<b>Member State where product is authorised</b>
Seroquel 100 mg filmsko obložene tablete	NL/H/0156/002	H/99/01407/007	ASTRAZENECA AB	SI
Seroquel 100 mg filmsko obložene tablete	NL/H/0156/002	H/99/01407/010	ASTRAZENECA AB	SI
Seroquel 100 mg filmsko obložene tablete	NL/H/0156/002	H/99/01407/009	ASTRAZENECA AB	SI
Seroquel 100 mg filmsko obložene tablete	NL/H/0156/002	H/99/01407/058	ASTRAZENECA AB	SI
Seroquel 100 mg filmsko obložene tablete	NL/H/0156/002	H/99/01407/011	ASTRAZENECA AB	SI
Seroquel 100 mg filmsko obložene tablete	NL/H/0156/002	H/99/01407/012	ASTRAZENECA AB	SI
Seroquel 100 mg filmsko obložene tablete	NL/H/0156/002	H/99/01407/008	ASTRAZENECA AB	SI
Seroquel 100 mg filmsko obložene tablete	NL/H/0156/002	H/99/01407/013	ASTRAZENECA AB	SI
Seroquel 100 mg Filmtabletten	NL/H/0156/002	1-23461	ASTRAZENECA OSTERREICH GMBH	AT
Seroquel 100 mg filmuhúðaðar töflur	NL/H/0156/002	IS/1/03/002/02	ASTRAZENECA AB	IS
Seroquel 100 mg tabletti, kalvopäällysteinen	NL/H/0156/002	14758	ASTRAZENECA OY	FI
Seroquel 100 mg επικαλυμμένα με λεπτό υμένιο δισκία	NL/H/0156/002	17717	ASTRAZENECA AB	CY
Seroquel 100 mg επικαλυμμένα με λεπτό υμένιο δισκία	NL/H/0156/002	54573/15.06.2017	ASTRAZENECA S.A	GR
Seroquel 100 mg, comprimés pelliculés	NL/H/0156/002	BE210366	ASTRAZENECA S.A. / N.V.	BE
Seroquel 100 mg, comprimés pelliculés	NL/H/0156/002	2010010630	ASTRAZENECA S.A. / N.V.	LU
SEROQUEL 100 mg,	NL/H/0156/002	BE210366	ASTRAZENECA S.A. / N.V.	BE

<b>Product Name (in authorisation country)</b>	<b>MRP/DCP Authorisation number</b>	<b>National Authorisation Number</b>	<b>MAH of product in the member state</b>	<b>Member State where product is authorised</b>
filmomhulde tabletten				
Seroquel 100 mg, filmomhulde tabletten	NL/H/0156/002	RVG 20827	ASTRAZENECA BV	NL
Seroquel 150 mg compresse a rilascio prolungato	NL/H/0156/012	A.I.C. 032944163	ASTRAZENECA S.P.A.	IT
Seroquel 150 mg compresse a rilascio prolungato	NL/H/0156/012	A.I.C. 032944530	ASTRAZENECA S.P.A.	IT
Seroquel 150 mg compresse a rilascio prolungato	NL/H/0156/012	A.I.C. 032944542	ASTRAZENECA S.P.A.	IT
Seroquel 150 mg compresse a rilascio prolungato	NL/H/0156/012	A.I.C. 032944579	ASTRAZENECA S.P.A.	IT
Seroquel 150 mg compresse a rilascio prolungato	NL/H/0156/012	A.I.C. 032944567	ASTRAZENECA S.P.A.	IT
Seroquel 150 mg compresse a rilascio prolungato	NL/H/0156/012	A.I.C. 032944555	ASTRAZENECA S.P.A.	IT
SEROQUEL 150 mg compresse rivestite con film	NL/H/0156/006	A.I.C. 32944086	ASTRAZENECA S.P.A.	IT
SEROQUEL 150 mg compresse rivestite con film	NL/H/0156/006	A.I.C. 32944074	ASTRAZENECA S.P.A.	IT
SEROQUEL 150 mg compresse rivestite con film	NL/H/0156/006	A.I.C. 032944314	ASTRAZENECA S.P.A.	IT
SEROQUEL 150 mg compresse rivestite con film	NL/H/0156/006	A.I.C. 032944338	ASTRAZENECA S.P.A.	IT
SEROQUEL 150 mg	NL/H/0156/006	A.I.C. 032944290	ASTRAZENECA S.P.A.	IT

<b>Product Name (in authorisation country)</b>	<b>MRP/DCP Authorisation number</b>	<b>National Authorisation Number</b>	<b>MAH of product in the member state</b>	<b>Member State where product is authorised</b>
compresse rivestite con film				
SEROQUEL 150 mg compresse rivestite con film	NL/H/0156/006	A.I.C. 032944326	ASTRAZENECA S.P.A.	IT
SEROQUEL 150 mg compresse rivestite con film	NL/H/0156/006	A.I.C. 032944288	ASTRAZENECA S.P.A.	IT
SEROQUEL 150 mg compresse rivestite con film	NL/H/0156/006	A.I.C. 032944302	ASTRAZENECA S.P.A.	IT
SEROQUEL 150 mg compresse rivestite con film	NL/H/0156/006	A.I.C. 032944264	ASTRAZENECA S.P.A.	IT
SEROQUEL 150 mg compresse rivestite con film	NL/H/0156/006	A.I.C. 032944276	ASTRAZENECA S.P.A.	IT
Seroquel 150 mg comprimidos revestidos por película	NL/H/0156/006	3734084	ASTRAZENECA PRODUTOS FARMACEUTICOS LDA	PT
Seroquel 150 mg comprimidos revestidos por película	NL/H/0156/006	3733086	ASTRAZENECA PRODUTOS FARMACEUTICOS LDA	PT
Seroquel 150 mg comprimidos revestidos por película	NL/H/0156/006	3733987	ASTRAZENECA PRODUTOS FARMACEUTICOS LDA	PT
Seroquel 150 mg comprimidos revestidos por película	NL/H/0156/006	3733284	ASTRAZENECA PRODUTOS FARMACEUTICOS LDA	PT
Seroquel 150 mg comprimidos revestidos por película	NL/H/0156/006	3733383	ASTRAZENECA PRODUTOS FARMACEUTICOS LDA	PT
Seroquel 150 mg comprimidos revestidos	NL/H/0156/006	3733482	ASTRAZENECA PRODUTOS FARMACEUTICOS LDA	PT



<b>Product Name (in authorisation country)</b>	<b>MRP/DCP Authorisation number</b>	<b>National Authorisation Number</b>	<b>MAH of product in the member state</b>	<b>Member State where product is authorised</b>
por película				
Seroquel 150 mg comprimidos revestidos por película	NL/H/0156/006	3733789	ASTRAZENECA PRODUTOS FARMACEUTICOS LDA	PT
Seroquel 150 mg comprimidos revestidos por película	NL/H/0156/006	3733185	ASTRAZENECA PRODUTOS FARMACEUTICOS LDA	PT
Seroquel 150 mg comprimidos revestidos por película	NL/H/0156/006	3733581	ASTRAZENECA PRODUTOS FARMACEUTICOS LDA	PT
Seroquel 150 mg comprimidos revestidos por película	NL/H/0156/006	3733680	ASTRAZENECA PRODUTOS FARMACEUTICOS LDA	PT
Seroquel 150 mg comprimidos revestidos por película	NL/H/0156/006	3733888	ASTRAZENECA PRODUTOS FARMACEUTICOS LDA	PT
Seroquel 150 mg filmdragerade tabletter	NL/H/0156/006	18964	ASTRAZENECA AB	SE
Seroquel 150 mg filmdrasjerte tabletter	NL/H/0156/006	02-1437	ASTRAZENECA AS	NO
Seroquel 150 mg Filmtabletten	NL/H/0156/006	1-24318	ASTRAZENECA OSTERREICH GMBH	AT
Seroquel 150 mg filmuhúðaðar töflur	NL/H/0156/006	IS/1/03/002/03	ASTRAZENECA AB	IS
Seroquel 150 mg tabletti, kalvopäällysteinen	NL/H/0156/006	16558	ASTRAZENECA OY	FI
Seroquel 150 mg επικαλυμμένα με λεπτό υμένιο δισκία	NL/H/0156/006	54577/15.06.2017	ASTRAZENECA S.A	GR
Seroquel 150 mg, comprimés pelliculés	NL/H/0156/006	BE228891	ASTRAZENECA S.A. / N.V.	BE
Seroquel 150 mg, comprimés pelliculés	NL/H/0156/006	2010010632	ASTRAZENECA S.A. / N.V.	LU

<b>Product Name (in authorisation country)</b>	<b>MRP/DCP Authorisation number</b>	<b>National Authorisation Number</b>	<b>MAH of product in the member state</b>	<b>Member State where product is authorised</b>
SEROQUEL 150 mg, filmomhulde tabletten	NL/H/0156/006	BE228891	ASTRAZENECA S.A. / N.V.	BE
Seroquel 150 mg, filmomhulde tabletten	NL/H/0156/006	RVG 25602	ASTRAZENECA BV	NL
Seroquel 200 mg compresse a rilascio prolungato	NL/H/0156/009	A.I.C 32944136	ASTRAZENECA S.P.A.	IT
Seroquel 200 mg compresse a rilascio prolungato	NL/H/0156/009	A.I.C. 032944593	ASTRAZENECA S.P.A.	IT
Seroquel 200 mg compresse a rilascio prolungato	NL/H/0156/009	A.I.C. 032944581	ASTRAZENECA S.P.A.	IT
Seroquel 200 mg compresse a rilascio prolungato	NL/H/0156/009	A.I.C. 032944617	ASTRAZENECA S.P.A.	IT
Seroquel 200 mg compresse a rilascio prolungato	NL/H/0156/009	A.I.C. 032944629	ASTRAZENECA S.P.A.	IT
Seroquel 200 mg compresse a rilascio prolungato	NL/H/0156/009	A.I.C. 032944605	ASTRAZENECA S.P.A.	IT
SEROQUEL 200 mg compresse rivestite con film	NL/H/0156/003	A.I.C 32944050	ASTRAZENECA S.P.A.	IT
SEROQUEL 200 mg compresse rivestite con film	NL/H/0156/003	A.I.C 32944047	ASTRAZENECA S.P.A.	IT
SEROQUEL 200 mg compresse rivestite con film	NL/H/0156/003	A.I.C. 032944389	ASTRAZENECA S.P.A.	IT
SEROQUEL 200 mg compresse rivestite con film	NL/H/0156/003	A.I.C. 032944340	ASTRAZENECA S.P.A.	IT

<b>Product Name (in authorisation country)</b>	<b>MRP/DCP Authorisation number</b>	<b>National Authorisation Number</b>	<b>MAH of product in the member state</b>	<b>Member State where product is authorised</b>
SEROQUEL 200 mg compresse rivestite con film	NL/H/0156/003	A.I.C. 032944365	ASTRAZENECA S.P.A.	IT
SEROQUEL 200 mg compresse rivestite con film	NL/H/0156/003	A.I.C. 032944353	ASTRAZENECA S.P.A.	IT
SEROQUEL 200 mg compresse rivestite con film	NL/H/0156/003	A.I.C. 032944377	ASTRAZENECA S.P.A.	IT
Seroquel 200 mg comprimidos recubiertos con película	NL/H/0156/003	63.056	ASTRAZENECA FARMACÉUTICA SPAIN, S.A.	ES
Seroquel 200 mg comprimidos revestidos por película	NL/H/0156/003	3074887	ASTRAZENECA PRODUTOS FARMACEUTICOS LDA	PT
Seroquel 200 mg comprimidos revestidos por película	NL/H/0156/003	3075488	ASTRAZENECA PRODUTOS FARMACEUTICOS LDA	PT
Seroquel 200 mg comprimidos revestidos por película	NL/H/0156/003	3074986	ASTRAZENECA PRODUTOS FARMACEUTICOS LDA	PT
Seroquel 200 mg comprimidos revestidos por película	NL/H/0156/003	3075181	ASTRAZENECA PRODUTOS FARMACEUTICOS LDA	PT
Seroquel 200 mg comprimidos revestidos por película	NL/H/0156/003	3075389	ASTRAZENECA PRODUTOS FARMACEUTICOS LDA	PT
Seroquel 200 mg comprimidos revestidos por película	NL/H/0156/003	3075082	ASTRAZENECA PRODUTOS FARMACEUTICOS LDA	PT
Seroquel 200 mg comprimidos revestidos por película	NL/H/0156/003	3075587	ASTRAZENECA PRODUTOS FARMACEUTICOS LDA	PT
Seroquel 200 mg	NL/H/0156/003	3075280	ASTRAZENECA PRODUTOS	PT

<b>Product Name (in authorisation country)</b>	<b>MRP/DCP Authorisation number</b>	<b>National Authorisation Number</b>	<b>MAH of product in the member state</b>	<b>Member State where product is authorised</b>
comprimidos revestidos por película			FARMACEUTICOS LDA	
Seroquel 200 mg film-coated tablets	NL/H/0156/003	PA 1019/019/003	ASTRAZENECA AB	IE
Seroquel 200 mg film-coated tablets	NL/H/0156/003	MA 046/02303	ASTRAZENECA AB	MT
Seroquel 200 mg film-coated tablets	NL/H/0156/003	PL 50827/0004	LUYE PHARMA LIMITED	UK
Seroquel 200 mg filmdragerade tabletter	NL/H/0156/003	18960	ASTRAZENECA AB	SE
Seroquel 200 mg filmdrasjerte tabletter	NL/H/0156/003	02-1438	ASTRAZENECA AS	NO
SEROQUEL 200 mg filmom obložene tablete	NL/H/0156/003	HR-H-204008507	ASTRAZENECA D.O.O.	HR
Seroquel 200 mg Filmtabletten	NL/H/0156/003	1-23463	ASTRAZENECA OSTERREICH GMBH	AT
Seroquel 200 mg filmuhúðaðar töflur	NL/H/0156/003	IS/1/03/002/04	ASTRAZENECA AB	IS
Seroquel 200 mg tabletti, kalvopäällysteinen	NL/H/0156/003	14759	ASTRAZENECA OY	FI
Seroquel 200 mg επικαλυμμένα με λεπτό υμένιο δισκία	NL/H/0156/003	17718	ASTRAZENECA AB	CY
Seroquel 200 mg επικαλυμμένα με λεπτό υμένιο δισκία	NL/H/0156/003	54574/15.06.2017	ASTRAZENECA S.A	GR
Seroquel 200 mg, comprimés pelliculés	NL/H/0156/003	BE210375	ASTRAZENECA S.A. / N.V.	BE
Seroquel 200 mg, comprimés pelliculés	NL/H/0156/003	2010010633	ASTRAZENECA S.A. / N.V.	LU
SEROQUEL 200 mg, filmomhulde tabletten	NL/H/0156/003	BE210375	ASTRAZENECA S.A. / N.V.	BE
Seroquel 200 mg,	NL/H/0156/003	RVG 20828	ASTRAZENECA BV	NL

<b>Product Name (in authorisation country)</b>	<b>MRP/DCP Authorisation number</b>	<b>National Authorisation Number</b>	<b>MAH of product in the member state</b>	<b>Member State where product is authorised</b>
filmomhulde tabletten				
SEROQUEL 25 mg compresse rivestite con film	NL/H/0156/001	A.I.C 32944112	ASTRAZENECA S.P.A.	IT
SEROQUEL 25 mg compresse rivestite con film	NL/H/0156/001	A.I.C 32944011	ASTRAZENECA S.P.A.	IT
SEROQUEL 25 mg compresse rivestite con film	NL/H/0156/001	A.I.C. 032944201	ASTRAZENECA S.P.A.	IT
SEROQUEL 25 mg compresse rivestite con film	NL/H/0156/001	A.I.C. 032944175	ASTRAZENECA S.P.A.	IT
SEROQUEL 25 mg compresse rivestite con film	NL/H/0156/001	A.I.C. 032944187	ASTRAZENECA S.P.A.	IT
SEROQUEL 25 mg compresse rivestite con film	NL/H/0156/001	A.I.C. 032944199	ASTRAZENECA S.P.A.	IT
Seroquel 25 mg comprimidos recubiertos con película	NL/H/0156/001	63.054	ASTRAZENECA FARMACÉUTICA SPAIN, S.A.	ES
Seroquel 25 mg comprimidos revestidos por película	NL/H/0156/001	3073681	ASTRAZENECA PRODUTOS FARMACEUTICOS LDA	PT
Seroquel 25 mg comprimidos revestidos por película	NL/H/0156/001	3073582	ASTRAZENECA PRODUTOS FARMACEUTICOS LDA	PT
Seroquel 25 mg comprimidos revestidos por película	NL/H/0156/001	3073988	ASTRAZENECA PRODUTOS FARMACEUTICOS LDA	PT
Seroquel 25 mg comprimidos revestidos por película	NL/H/0156/001	3073780	ASTRAZENECA PRODUTOS FARMACEUTICOS LDA	PT

<b>Product Name (in authorisation country)</b>	<b>MRP/DCP Authorisation number</b>	<b>National Authorisation Number</b>	<b>MAH of product in the member state</b>	<b>Member State where product is authorised</b>
Seroquel 25 mg comprimidos revestidos por película	NL/H/0156/001	3074085	ASTRAZENECA PRODUTOS FARMACEUTICOS LDA	PT
Seroquel 25 mg comprimidos revestidos por película	NL/H/0156/001	3539087	ASTRAZENECA PRODUTOS FARMACEUTICOS LDA	PT
Seroquel 25 mg comprimidos revestidos por película	NL/H/0156/001	3073889	ASTRAZENECA PRODUTOS FARMACEUTICOS LDA	PT
Seroquel 25 mg film-coated tablets	NL/H/0156/001	PA 1019/019/001	ASTRAZENECA AB	IE
Seroquel 25 mg film-coated tablets	NL/H/0156/001	MA 046/02301	ASTRAZENECA AB	MT
Seroquel 25 mg film-coated tablets	NL/H/0156/001	PL 50827/0002	LUYE PHARMA LIMITED	UK
Seroquel 25 mg filmdragerade tabletter	NL/H/0156/001	18958	ASTRAZENECA AB	SE
Seroquel 25 mg filmdrasjerte tabletter	NL/H/0156/001	02-1435	ASTRAZENECA AS	NO
SEROQUEL 25 mg filmom obložene tablete	NL/H/0156/001	HR-H-619090976	ASTRAZENECA D.O.O.	HR
Seroquel 25 mg filmsko obložene tablete	NL/H/0156/001	H/99/01407/001	ASTRAZENECA AB	SI
Seroquel 25 mg filmsko obložene tablete	NL/H/0156/001	H/99/01407/002	ASTRAZENECA AB	SI
Seroquel 25 mg filmsko obložene tablete	NL/H/0156/001	H/99/01407/005	ASTRAZENECA AB	SI
Seroquel 25 mg filmsko obložene tablete	NL/H/0156/001	H/99/01407/003	ASTRAZENECA AB	SI
Seroquel 25 mg filmsko obložene tablete	NL/H/0156/001	H/99/01407/004	ASTRAZENECA AB	SI
Seroquel 25 mg filmsko obložene tablete	NL/H/0156/001	H/99/01407/057	ASTRAZENECA AB	SI
Seroquel 25 mg filmsko obložene tablete	NL/H/0156/001	H/99/01407/056	ASTRAZENECA AB	SI

<b>Product Name (in authorisation country)</b>	<b>MRP/DCP Authorisation number</b>	<b>National Authorisation Number</b>	<b>MAH of product in the member state</b>	<b>Member State where product is authorised</b>
obložene tablete				
Seroquel 25 mg filmsko obložene tablete	NL/H/0156/001	H/99/01407/006	ASTRAZENECA AB	SI
Seroquel 25 mg Filmtabletten	NL/H/0156/001	1-23460	ASTRAZENECA OSTERREICH GMBH	AT
Seroquel 25 mg filmuhúðaðar töflur	NL/H/0156/001	IS/1/03/002/01	ASTRAZENECA AB	IS
Seroquel 25 mg tabletti, kalvopäällysteinen	NL/H/0156/001	14757	ASTRAZENECA OY	FI
Seroquel 25 mg επικαλυμμένα με λεπτό υμένιο δισκία	NL/H/0156/001	17716	ASTRAZENECA AB	CY
Seroquel 25 mg επικαλυμμένα με λεπτό υμένιο δισκία	NL/H/0156/001	54572/15.06.2017	ASTRAZENECA S.A	GR
Seroquel 25 mg, comprimés pelliculés	NL/H/0156/001	BE210357	ASTRAZENECA S.A. / N.V.	BE
Seroquel 25 mg, comprimés pelliculés	NL/H/0156/001	2010010629	ASTRAZENECA S.A. / N.V.	LU
SEROQUEL 25 mg, filmomhulde tabletten	NL/H/0156/001	BE210357	ASTRAZENECA S.A. / N.V.	BE
Seroquel 25 mg, filmomhulde tabletten	NL/H/0156/001	RVG 20826	ASTRAZENECA BV	NL
Seroquel 300 mg compresse a rilascio prolungato	NL/H/0156/010	A.I.C 32944148	ASTRAZENECA S.P.A.	IT
Seroquel 300 mg compresse a rilascio prolungato	NL/H/0156/010	A.I.C. 032944670	ASTRAZENECA S.P.A.	IT
Seroquel 300 mg compresse a rilascio prolungato	NL/H/0156/010	A.I.C. 032944643	ASTRAZENECA S.P.A.	IT
Seroquel 300 mg compresse a rilascio	NL/H/0156/010	A.I.C. 032944668	ASTRAZENECA S.P.A.	IT

<b>Product Name (in authorisation country)</b>	<b>MRP/DCP Authorisation number</b>	<b>National Authorisation Number</b>	<b>MAH of product in the member state</b>	<b>Member State where product is authorised</b>
prolungato				
Seroquel 300 mg compresse a rilascio prolungato	NL/H/0156/010	A.I.C. 032944631	ASTRAZENECA S.P.A.	IT
Seroquel 300 mg compresse a rilascio prolungato	NL/H/0156/010	A.I.C. 032944656	ASTRAZENECA S.P.A.	IT
SEROQUEL 300 mg compresse rivestite con film	NL/H/0156/007	A.I.C 032944100	ASTRAZENECA S.P.A.	IT
SEROQUEL 300 mg compresse rivestite con film	NL/H/0156/007	A.I.C 032944098	ASTRAZENECA S.P.A.	IT
SEROQUEL 300 mg compresse rivestite con film	NL/H/0156/007	A.I.C. 032944427	ASTRAZENECA S.P.A.	IT
SEROQUEL 300 mg compresse rivestite con film	NL/H/0156/007	A.I.C. 032944415	ASTRAZENECA S.P.A.	IT
SEROQUEL 300 mg compresse rivestite con film	NL/H/0156/007	A.I.C. 032944403	ASTRAZENECA S.P.A.	IT
SEROQUEL 300 mg compresse rivestite con film	NL/H/0156/007	A.I.C. 032944391	ASTRAZENECA S.P.A.	IT
SEROQUEL 300 mg compresse rivestite con film	NL/H/0156/007	A.I.C. 032944441	ASTRAZENECA S.P.A.	IT
SEROQUEL 300 mg compresse rivestite con film	NL/H/0156/007	A.I.C. 032944454	ASTRAZENECA S.P.A.	IT
SEROQUEL 300 mg compresse rivestite con film	NL/H/0156/007	A.I.C. 032944466	ASTRAZENECA S.P.A.	IT



<b>Product Name (in authorisation country)</b>	<b>MRP/DCP Authorisation number</b>	<b>National Authorisation Number</b>	<b>MAH of product in the member state</b>	<b>Member State where product is authorised</b>
SEROQUEL 300 mg compresse rivestite con film	NL/H/0156/007	A.I.C. 032944439	ASTRAZENECA S.P.A.	IT
Seroquel 300 mg comprimidos recubiertos con película	NL/H/0156/007	64.436	ASTRAZENECA FARMACÉUTICA SPAIN, S.A.	ES
Seroquel 300 mg comprimidos revestidos por película	NL/H/0156/007	3734480	ASTRAZENECA PRODUTOS FARMACEUTICOS LDA	PT
Seroquel 300 mg comprimidos revestidos por película	NL/H/0156/007	3734381	ASTRAZENECA PRODUTOS FARMACEUTICOS LDA	PT
Seroquel 300 mg comprimidos revestidos por película	NL/H/0156/007	3735180	ASTRAZENECA PRODUTOS FARMACEUTICOS LDA	PT
Seroquel 300 mg comprimidos revestidos por película	NL/H/0156/007	3735289	ASTRAZENECA PRODUTOS FARMACEUTICOS LDA	PT
Seroquel 300 mg comprimidos revestidos por película	NL/H/0156/007	3734886	ASTRAZENECA PRODUTOS FARMACEUTICOS LDA	PT
Seroquel 300 mg comprimidos revestidos por película	NL/H/0156/007	3734787	ASTRAZENECA PRODUTOS FARMACEUTICOS LDA	PT
Seroquel 300 mg comprimidos revestidos por película	NL/H/0156/007	3734589	ASTRAZENECA PRODUTOS FARMACEUTICOS LDA	PT
Seroquel 300 mg comprimidos revestidos por película	NL/H/0156/007	3734985	ASTRAZENECA PRODUTOS FARMACEUTICOS LDA	PT
Seroquel 300 mg comprimidos revestidos por película	NL/H/0156/007	3735081	ASTRAZENECA PRODUTOS FARMACEUTICOS LDA	PT
Seroquel 300 mg	NL/H/0156/007	3734688	ASTRAZENECA PRODUTOS	PT

<b>Product Name (in authorisation country)</b>	<b>MRP/DCP Authorisation number</b>	<b>National Authorisation Number</b>	<b>MAH of product in the member state</b>	<b>Member State where product is authorised</b>
comprimidos revestidos por película			FARMACEUTICOS LDA	
Seroquel 300 mg comprimidos revestidos por película	NL/H/0156/007	3734282	ASTRAZENECA PRODUTOS FARMACEUTICOS LDA	PT
Seroquel 300 mg film-coated tablets	NL/H/0156/007	PA 1019/019/004	ASTRAZENECA AB	IE
Seroquel 300 mg film-coated tablets	NL/H/0156/007	PL 50827/0005	LUYE PHARMA LIMITED	UK
Seroquel 300 mg filmdragerade tabletter	NL/H/0156/007	18961	ASTRAZENECA AB	SE
Seroquel 300 mg filmdrasjerte tabletter	NL/H/0156/007	02-1439	ASTRAZENECA AS	NO
Seroquel 300 mg Filmtabletten	NL/H/0156/007	1-24319	ASTRAZENECA OSTERREICH GMBH	AT
Seroquel 300 mg filmuhúðaðar töflur	NL/H/0156/007	IS/1/03/002/05	ASTRAZENECA AB	IS
Seroquel 300 mg tabletti, kalvopäällysteinen	NL/H/0156/007	16559	ASTRAZENECA OY	FI
Seroquel 300 mg επικαλυμμένα με λεπτό υμένιο δισκία	NL/H/0156/007	54578/15.06.2017	ASTRAZENECA S.A	GR
Seroquel 300 mg, comprimés pelliculés	NL/H/0156/007	BE228907	ASTRAZENECA S.A. / N.V.	BE
Seroquel 300 mg, comprimés pelliculés	NL/H/0156/007	2010010631	ASTRAZENECA S.A. / N.V.	LU
SEROQUEL 300 mg, filmomhulde tabletten	NL/H/0156/007	BE228907	ASTRAZENECA S.A. / N.V.	BE
Seroquel 300 mg, filmomhulde tabletten	NL/H/0156/007	RVG 25603	ASTRAZENECA BV	NL
Seroquel 3-daagse Startverpakking (combinatieverpakking)	NL/H/0156/004	BE210341	ASTRAZENECA S.A. / N.V.	BE

<b>Product Name (in authorisation country)</b>	<b>MRP/DCP Authorisation number</b>	<b>National Authorisation Number</b>	<b>MAH of product in the member state</b>	<b>Member State where product is authorised</b>
Seroquel 400 mg compresse a rilascio prolungato	NL/H/0156/011	A.I.C 32944151	ASTRAZENECA S.P.A.	IT
Seroquel 400 mg compresse a rilascio prolungato	NL/H/0156/011	A.I.C. 032944718	ASTRAZENECA S.P.A.	IT
Seroquel 400 mg compresse a rilascio prolungato	NL/H/0156/011	A.I.C. 032944720	ASTRAZENECA S.P.A.	IT
Seroquel 400 mg compresse a rilascio prolungato	NL/H/0156/011	A.I.C. 032944694	ASTRAZENECA S.P.A.	IT
Seroquel 400 mg compresse a rilascio prolungato	NL/H/0156/011	A.I.C. 032944682	ASTRAZENECA S.P.A.	IT
Seroquel 400 mg compresse a rilascio prolungato	NL/H/0156/011	A.I.C. 032944706	ASTRAZENECA S.P.A.	IT
Seroquel 4-daagse Startverpakking	NL/H/0156/005	BE217305	ASTRAZENECA S.A. / N.V.	BE
Seroquel 4-daagse Startverpakking	NL/H/0156/005	RVG 25128	ASTRAZENECA BV	NL
SEROQUEL 4-day Starter Pack, comprimés pelliculés	NL/H/0156/005	2006088995	ASTRAZENECA S.A. / N.V.	LU
Seroquel 4-Tage Starterpackung	NL/H/0156/005	1-23709	ASTRAZENECA OSTERREICH GMBH	AT
Seroquel 50 mg compresse a rilascio prolungato	NL/H/0156/008	A.I.C 32944124	ASTRAZENECA S.P.A.	IT
Seroquel 50 mg compresse a rilascio prolungato	NL/H/0156/008	A.I.C. 032944480	ASTRAZENECA S.P.A.	IT
Seroquel 50 mg	NL/H/0156/008	A.I.C. 032944528	ASTRAZENECA S.P.A.	IT

<b>Product Name (in authorisation country)</b>	<b>MRP/DCP Authorisation number</b>	<b>National Authorisation Number</b>	<b>MAH of product in the member state</b>	<b>Member State where product is authorised</b>
compresse a rilascio prolungato				
Seroquel 50 mg compresse a rilascio prolungato	NL/H/0156/008	A.I.C. 032944492	ASTRAZENECA S.P.A.	IT
Seroquel 50 mg compresse a rilascio prolungato	NL/H/0156/008	A.I.C. 032944516	ASTRAZENECA S.P.A.	IT
Seroquel 50 mg compresse a rilascio prolungato	NL/H/0156/008	A.I.C. 032944504	ASTRAZENECA S.P.A.	IT
Seroquel Confezione Starter 3-Giorni (confezione combinata)	NL/H/0156/004	A.I.C. 032944478	ASTRAZENECA S.P.A.	IT
Seroquel Confezione Starter 4-Giorni	NL/H/0156/005	A.I.C. 032944062	ASTRAZENECA S.P.A.	IT
Seroquel Depot 150 mg depottabletter	NL/H/0156/012	08-5880	ASTRAZENECA AS	NO
Seroquel Depot 150 mg depottabletter	NL/H/0156/012	26908	ASTRAZENECA AB	SE
Seroquel Depot 200 mg depottabletter	NL/H/0156/009	07-5215	ASTRAZENECA AS	NO
Seroquel Depot 200 mg depottabletter	NL/H/0156/009	25987	ASTRAZENECA AB	SE
Seroquel Depot 300 mg depottabletter	NL/H/0156/010	07-5216	ASTRAZENECA AS	NO
Seroquel Depot 300 mg depottabletter	NL/H/0156/010	25988	ASTRAZENECA AB	SE
Seroquel Depot 400 mg depottabletter	NL/H/0156/011	07-5217	ASTRAZENECA AS	NO
Seroquel Depot 400 mg depottabletter	NL/H/0156/011	25989	ASTRAZENECA AB	SE
Seroquel Depot 50 mg depottabletter	NL/H/0156/008	07-5214	ASTRAZENECA AS	NO

<b>Product Name (in authorisation country)</b>	<b>MRP/DCP Authorisation number</b>	<b>National Authorisation Number</b>	<b>MAH of product in the member state</b>	<b>Member State where product is authorised</b>
Seroquel Depot 50 mg depottabletter	NL/H/0156/008	25986	ASTRAZENECA AB	SE
Seroquel Embalagem de início de 3 Dias (25 mg) + (100 mg) comprimidos revestidos por película	NL/H/0156/004	3075686	ASTRAZENECA PRODUTOS FARMACEUTICOS LDA	PT
Seroquel Embalagem de início de 4 Dias (25 mg) + (100 mg) + (200 mg) comprimidos revestidos por película	NL/H/0156/005	3233681	ASTRAZENECA PRODUTOS FARMACEUTICOS LDA	PT
Seroquel Prolong	NL/H/0156/010	42033	ASTRAZENECA A/S	DK
Seroquel Prolong	NL/H/0156/011	42034	ASTRAZENECA A/S	DK
Seroquel Prolong	NL/H/0156/008	42031	ASTRAZENECA A/S	DK
Seroquel Prolong	NL/H/0156/012	43173	ASTRAZENECA A/S	DK
Seroquel Prolong	NL/H/0156/009	42032	ASTRAZENECA A/S	DK
Seroquel Prolong 150 mg comprimidos de liberación prolongada	NL/H/0156/012	70.859	ASTRAZENECA FARMACÉUTICA SPAIN, S.A.	ES
Seroquel Prolong 150 mg depottabletti	NL/H/0156/012	24948	ASTRAZENECA OY	FI
Seroquel Prolong 150 mg forðatöflur	NL/H/0156/012	IS/1/08/081/01	ASTRAZENECA A/S	IS
Seroquel Prolong 200 mg comprimidos de liberación prolongada	NL/H/0156/009	69.646	ASTRAZENECA FARMACÉUTICA SPAIN, S.A.	ES
Seroquel Prolong 200 mg depottabletti	NL/H/0156/009	24065	ASTRAZENECA OY	FI
Seroquel Prolong 200 mg forðatöflur	NL/H/0156/009	IS/1/08/003/02	ASTRAZENECA A/S	IS
Seroquel Prolong 300 mg comprimidos de liberación prolongada	NL/H/0156/010	69.647	ASTRAZENECA FARMACÉUTICA SPAIN, S.A.	ES
Seroquel Prolong 300 mg	NL/H/0156/010	24066	ASTRAZENECA OY	FI

<b>Product Name (in authorisation country)</b>	<b>MRP/DCP Authorisation number</b>	<b>National Authorisation Number</b>	<b>MAH of product in the member state</b>	<b>Member State where product is authorised</b>
depottabletti				
Seroquel Prolong 300 mg forðatöflur	NL/H/0156/010	IS/1/08/003/03	ASTRAZENECA A/S	IS
Seroquel Prolong 400 mg comprimidos de liberación prolongada	NL/H/0156/011	69.648	ASTRAZENECA FARMACÉUTICA SPAIN, S.A.	ES
Seroquel Prolong 400 mg depottabletti	NL/H/0156/011	24067	ASTRAZENECA OY	FI
Seroquel Prolong 400 mg forðatöflur	NL/H/0156/011	IS/1/08/003/04	ASTRAZENECA A/S	IS
Seroquel Prolong 50 mg comprimidos de liberación prolongada	NL/H/0156/008	69.645	ASTRAZENECA FARMACÉUTICA SPAIN, S.A.	ES
Seroquel Prolong 50 mg depottabletti	NL/H/0156/008	24064	ASTRAZENECA OY	FI
Seroquel Prolong 50 mg forðatöflur	NL/H/0156/008	IS/1/08/003/01	ASTRAZENECA A/S	IS
Seroquel Prolong® 150 mg Retardtabletten	NL/H/0156/012	73290.00.00	ASTRAZENECA GMBH	DE
Seroquel Prolong® 200 mg Retardtabletten	NL/H/0156/009	70562.00.00	ASTRAZENECA GMBH	DE
Seroquel Prolong® 300 mg Retardtabletten	NL/H/0156/010	70563.00.00	ASTRAZENECA GMBH	DE
Seroquel Prolong® 400 mg Retardtabletten	NL/H/0156/011	70564.00.00	ASTRAZENECA GMBH	DE
Seroquel Prolong® 50 mg Retardtabletten	NL/H/0156/008	70561.00.00	ASTRAZENECA GMBH	DE
Seroquel SR 150 mg comprimidos de libertação prolongada	NL/H/0156/012	5168455	ASTRAZENECA PRODUTOS FARMACEUTICOS LDA	PT
Seroquel SR 150 mg comprimidos de libertação prolongada	NL/H/0156/012	5168463	ASTRAZENECA PRODUTOS FARMACEUTICOS LDA	PT
Seroquel SR 150 mg	NL/H/0156/012	5456033	ASTRAZENECA PRODUTOS	PT

<b>Product Name (in authorisation country)</b>	<b>MRP/DCP Authorisation number</b>	<b>National Authorisation Number</b>	<b>MAH of product in the member state</b>	<b>Member State where product is authorised</b>
comprimidos de libertação prolongada			FARMACEUTICOS LDA	
Seroquel SR 150 mg tablete s podaljšanim sproščanjem	NL/H/0156/012	H/99/01407/037	ASTRAZENECA AB	SI
Seroquel SR 150 mg tablete s podaljšanim sproščanjem	NL/H/0156/012	H/99/01407/036	ASTRAZENECA AB	SI
Seroquel SR 150 mg tablete s podaljšanim sproščanjem	NL/H/0156/012	H/99/01407/038	ASTRAZENECA AB	SI
Seroquel SR 150 mg tablete s podaljšanim sproščanjem	NL/H/0156/012	H/99/01407/062	ASTRAZENECA AB	SI
Seroquel SR 150 mg tablete s podaljšanim sproščanjem	NL/H/0156/012	H/99/01407/039	ASTRAZENECA AB	SI
Seroquel SR 150 mg tablete s podaljšanim sproščanjem	NL/H/0156/012	H/99/01407/040	ASTRAZENECA AB	SI
Seroquel SR 150 mg tablete s podaljšanim sproščanjem	NL/H/0156/012	H/99/01407/061	ASTRAZENECA AB	SI
Seroquel SR 200 mg comprimidos de libertação prolongada	NL/H/0156/009	5085238	ASTRAZENECA PRODUTOS FARMACEUTICOS LDA	PT
Seroquel SR 200 mg comprimidos de libertação prolongada	NL/H/0156/009	5085220	ASTRAZENECA PRODUTOS FARMACEUTICOS LDA	PT
Seroquel SR 200 mg tablete s podaljšanim sproščanjem	NL/H/0156/009	H/99/01407/041	ASTRAZENECA AB	SI
Seroquel SR 200 mg tablete s podaljšanim sproščanjem	NL/H/0156/009	H/99/01407/043	ASTRAZENECA AB	SI

<b>Product Name (in authorisation country)</b>	<b>MRP/DCP Authorisation number</b>	<b>National Authorisation Number</b>	<b>MAH of product in the member state</b>	<b>Member State where product is authorised</b>
spročanjem				
Seroquel SR 200 mg tablete s podaljšanim sproščanjem	NL/H/0156/009	H/99/01407/045	ASTRAZENECA AB	SI
Seroquel SR 200 mg tablete s podaljšanim sproščanjem	NL/H/0156/009	H/99/01407/064	ASTRAZENECA AB	SI
Seroquel SR 200 mg tablete s podaljšanim sproščanjem	NL/H/0156/009	H/99/01407/063	ASTRAZENECA AB	SI
Seroquel SR 200 mg tablete s podaljšanim sproščanjem	NL/H/0156/009	H/99/01407/044	ASTRAZENECA AB	SI
Seroquel SR 200 mg tablete s podaljšanim sproščanjem	NL/H/0156/009	H/99/01407/042	ASTRAZENECA AB	SI
Seroquel SR 300 mg comprimidos de libertação prolongada	NL/H/0156/010	5085246	ASTRAZENECA PRODUTOS FARMACEUTICOS LDA	PT
Seroquel SR 300 mg comprimidos de libertação prolongada	NL/H/0156/010	5085253	ASTRAZENECA PRODUTOS FARMACEUTICOS LDA	PT
Seroquel SR 300 mg tablete s podaljšanim sproščanjem	NL/H/0156/010	H/99/01407/046	ASTRAZENECA AB	SI
Seroquel SR 300 mg tablete s podaljšanim sproščanjem	NL/H/0156/010	H/99/01407/047	ASTRAZENECA AB	SI
Seroquel SR 300 mg tablete s podaljšanim sproščanjem	NL/H/0156/010	H/99/01407/066	ASTRAZENECA AB	SI
Seroquel SR 300 mg tablete s podaljšanim sproščanjem	NL/H/0156/010	H/99/01407/048	ASTRAZENECA AB	SI



<b>Product Name (in authorisation country)</b>	<b>MRP/DCP Authorisation number</b>	<b>National Authorisation Number</b>	<b>MAH of product in the member state</b>	<b>Member State where product is authorised</b>
Seroquel SR 300 mg tablete s podaljšanim sproščanjem	NL/H/0156/010	H/99/01407/049	ASTRAZENECA AB	SI
Seroquel SR 300 mg tablete s podaljšanim sproščanjem	NL/H/0156/010	H/99/01407/050	ASTRAZENECA AB	SI
Seroquel SR 300 mg tablete s podaljšanim sproščanjem	NL/H/0156/010	H/99/01407/065	ASTRAZENECA AB	SI
Seroquel SR 400 mg comprimidos de libertação prolongada	NL/H/0156/011	5085261	ASTRAZENECA PRODUTOS FARMACEUTICOS LDA	PT
Seroquel SR 400 mg tablete s podaljšanim sproščanjem	NL/H/0156/011	H/99/01407/051	ASTRAZENECA AB	SI
Seroquel SR 400 mg tablete s podaljšanim sproščanjem	NL/H/0156/011	H/99/01407/052	ASTRAZENECA AB	SI
Seroquel SR 400 mg tablete s podaljšanim sproščanjem	NL/H/0156/011	H/99/01407/068	ASTRAZENECA AB	SI
Seroquel SR 400 mg tablete s podaljšanim sproščanjem	NL/H/0156/011	H/99/01407/067	ASTRAZENECA AB	SI
Seroquel SR 400 mg tablete s podaljšanim sproščanjem	NL/H/0156/011	H/99/01407/054	ASTRAZENECA AB	SI
Seroquel SR 400 mg tablete s podaljšanim sproščanjem	NL/H/0156/011	H/99/01407/055	ASTRAZENECA AB	SI
Seroquel SR 400 mg tablete s podaljšanim sproščanjem	NL/H/0156/011	H/99/01407/053	ASTRAZENECA AB	SI
Seroquel SR 50 mg	NL/H/0156/008	5178165	ASTRAZENECA PRODUTOS	PT

<b>Product Name (in authorisation country)</b>	<b>MRP/DCP Authorisation number</b>	<b>National Authorisation Number</b>	<b>MAH of product in the member state</b>	<b>Member State where product is authorised</b>
comprimidos de libertação prolongada			FARMACEUTICOS LDA	
Seroquel SR 50 mg comprimidos de libertação prolongada	NL/H/0156/008	5085212	ASTRAZENECA PRODUTOS FARMACEUTICOS LDA	PT
Seroquel SR 50 mg tablete s podaljšanim sproščanjem	NL/H/0156/008	H/99/01407/031	ASTRAZENECA AB	SI
Seroquel SR 50 mg tablete s podaljšanim sproščanjem	NL/H/0156/008	H/99/01407/035	ASTRAZENECA AB	SI
Seroquel SR 50 mg tablete s podaljšanim sproščanjem	NL/H/0156/008	H/99/01407/032	ASTRAZENECA AB	SI
Seroquel SR 50 mg tablete s podaljšanim sproščanjem	NL/H/0156/008	H/99/01407/060	ASTRAZENECA AB	SI
Seroquel SR 50 mg tablete s podaljšanim sproščanjem	NL/H/0156/008	H/99/01407/034	ASTRAZENECA AB	SI
Seroquel SR 50 mg tablete s podaljšanim sproščanjem	NL/H/0156/008	H/99/01407/059	ASTRAZENECA AB	SI
Seroquel SR 50 mg tablete s podaljšanim sproščanjem	NL/H/0156/008	H/99/01407/033	ASTRAZENECA AB	SI
Seroquel Starter Pack de 3 jours (emballage combiné)	NL/H/0156/004	BE210341	ASTRAZENECA S.A. / N.V.	BE
Seroquel Starter Pack de 3 jours (emballage combiné)	NL/H/0156/004	2006088994	ASTRAZENECA S.A. / N.V.	LU
Seroquel Starter Pack de 4 jours	NL/H/0156/005	BE217305	ASTRAZENECA S.A. / N.V.	BE

<b>Product Name (in authorisation country)</b>	<b>MRP/DCP Authorisation number</b>	<b>National Authorisation Number</b>	<b>MAH of product in the member state</b>	<b>Member State where product is authorised</b>
Seroquel XL 150 mg prolonged-release tablets	NL/H/0156/012	PL 50827/0007	LUYE PHARMA LIMITED	UK
Seroquel XL 200 mg prolonged-release tablets	NL/H/0156/009	PL 50827/0008	LUYE PHARMA LIMITED	UK
Seroquel XL 300 mg prolonged-release tablets	NL/H/0156/010	PL 50827/0009	LUYE PHARMA LIMITED	UK
Seroquel XL 400 mg prolonged-release tablets	NL/H/0156/011	PL 50827/0010	LUYE PHARMA LIMITED	UK
Seroquel XL 50 mg prolonged-release tablets	NL/H/0156/008	PL 50827/0006	LUYE PHARMA LIMITED	UK
Seroquel XR 150 mg comprimate cu eliberare prelungita	NL/H/0156/012	9529/2016/01	ASTRAZENECA AB	RO
Seroquel XR 150 mg comprimate cu eliberare prelungita	NL/H/0156/012	9529/2016/03	ASTRAZENECA AB	RO
Seroquel XR 150 mg comprimate cu eliberare prelungita	NL/H/0156/012	9529/2016/02	ASTRAZENECA AB	RO
Seroquel XR 150 mg comprimate cu eliberare prelungita	NL/H/0156/012	9529/2016/04	ASTRAZENECA AB	RO
Seroquel XR 150 mg comprimate cu eliberare prelungita	NL/H/0156/012	9529/2016/06	ASTRAZENECA AB	RO
Seroquel XR 150 mg comprimate cu eliberare prelungita	NL/H/0156/012	9529/2016/07	ASTRAZENECA AB	RO
Seroquel XR 150 mg comprimate cu eliberare prelungită	NL/H/0156/012	9529/2016/05	ASTRAZENECA AB	RO
Seroquel XR 150 mg prolonged-release tablets	NL/H/0156/012	PA 1019/019/006	ASTRAZENECA AB	IE
Seroquel XR 150 mg	NL/H/0156/012	OGYI-T-5863/08	ASTRAZENECA KFT.	HU

<b>Product Name (in authorisation country)</b>	<b>MRP/DCP Authorisation number</b>	<b>National Authorisation Number</b>	<b>MAH of product in the member state</b>	<b>Member State where product is authorised</b>
retard tableta				
Seroquel XR 150 mg retard tableta	NL/H/0156/012	OGYI-T-5863/09	ASTRAZENECA KFT.	HU
Seroquel XR 150 mg Retardtabletten	NL/H/0156/012	1-28377	ASTRAZENECA OSTERREICH GMBH	AT
Seroquel XR 150 mg tablete s produljenim oslobađanjem	NL/H/0156/012	HR-H-202528018	ASTRAZENECA D.O.O.	HR
SEROQUEL XR 150 mg δισκία παρατεταμένης αποδέσμευσης	NL/H/0156/012	20699	ASTRAZENECA AB	CY
Seroquel XR 150 mg δισκία παρατεταμένης αποδέσμευσης	NL/H/0156/012	54580/15.06.2017	ASTRAZENECA S.A	GR
Seroquel XR 150 mg, comprimés à libération prolongée	NL/H/0156/012	BE331947	ASTRAZENECA S.A. / N.V.	BE
Seroquel XR 150 mg, comprimés à libération prolongée	NL/H/0156/012	2009030009	ASTRAZENECA S.A. / N.V.	LU
Seroquel XR 150 mg, tabletten met verlengde afgifte	NL/H/0156/012	BE331947	ASTRAZENECA S.A. / N.V.	BE
Seroquel XR 150 mg, tabletten met verlengde afgifte	NL/H/0156/012	RVG 102408	ASTRAZENECA BV	NL
Seroquel XR 200 mg comprimate cu eliberare prelungita	NL/H/0156/009	9530/2016/03	ASTRAZENECA AB	RO
Seroquel XR 200 mg comprimate cu eliberare prelungita	NL/H/0156/009	9530/2016/02	ASTRAZENECA AB	RO
Seroquel XR 200 mg comprimate cu eliberare	NL/H/0156/009	9530/2016/01	ASTRAZENECA AB	RO

<b>Product Name (in authorisation country)</b>	<b>MRP/DCP Authorisation number</b>	<b>National Authorisation Number</b>	<b>MAH of product in the member state</b>	<b>Member State where product is authorised</b>
prelungita				
Seroquel XR 200 mg comprimate cu eliberare prelungita	NL/H/0156/009	9530/2016/07	ASTRAZENECA AB	RO
Seroquel XR 200 mg comprimate cu eliberare prelungita	NL/H/0156/009	9530/2016/04	ASTRAZENECA AB	RO
Seroquel XR 200 mg comprimate cu eliberare prelungita	NL/H/0156/009	9530/2016/05	ASTRAZENECA AB	RO
Seroquel XR 200 mg comprimate cu eliberare prelungita	NL/H/0156/009	9530/2016/06	ASTRAZENECA AB	RO
Seroquel XR 200 mg prolonged-release tablets	NL/H/0156/009	PA 1019/019/007	ASTRAZENECA AB	IE
Seroquel XR 200 mg retard tableta	NL/H/0156/009	OGYI-T-5863/11	ASTRAZENECA KFT.	HU
Seroquel XR 200 mg retard tableta	NL/H/0156/009	OGYI-T-5863/14	ASTRAZENECA KFT.	HU
Seroquel XR 200 mg retard tableta	NL/H/0156/009	OGYI-T-5863/03	ASTRAZENECA KFT.	HU
Seroquel XR 200 mg Retardtabletten	NL/H/0156/009	1-27421	ASTRAZENECA OSTERREICH GMBH	AT
Seroquel XR 200 mg tablete s produljenim oslobađanjem	NL/H/0156/009	HR-H-607016913	ASTRAZENECA D.O.O.	HR
Seroquel XR 200 mg tablety s predĺením uvoľňovaním	NL/H/0156/009	68/0231/07-S	ASTRAZENECA AB	SK
Seroquel XR 200 mg δισκία παρατεταμένης αποδέσμευσης	NL/H/0156/009	20357	ASTRAZENECA AB	CY
Seroquel XR 200 mg δισκία παρατεταμένης	NL/H/0156/009	54581/15.06.2017	ASTRAZENECA S.A	GR

<b>Product Name (in authorisation country)</b>	<b>MRP/DCP Authorisation number</b>	<b>National Authorisation Number</b>	<b>MAH of product in the member state</b>	<b>Member State where product is authorised</b>
αποδέσμευσης				
Seroquel XR 200 mg, comprimés à libération prolongée	NL/H/0156/009	BE314246	ASTRAZENECA S.A. / N.V.	BE
Seroquel XR 200 mg, comprimés à libération prolongée	NL/H/0156/009	2008040020	ASTRAZENECA S.A. / N.V.	LU
Seroquel XR 200 mg, tabletten met verlengde afgifte	NL/H/0156/009	BE314246	ASTRAZENECA S.A. / N.V.	BE
Seroquel XR 200 mg, tabletten met verlengde afgifte	NL/H/0156/009	RVG 34626	ASTRAZENECA BV	NL
Seroquel XR 300 mg comprimate cu eliberare prelungita	NL/H/0156/010	9531/2016/04	ASTRAZENECA AB	RO
Seroquel XR 300 mg comprimate cu eliberare prelungita	NL/H/0156/010	9531/2016/05	ASTRAZENECA AB	RO
Seroquel XR 300 mg comprimate cu eliberare prelungita	NL/H/0156/010	9531/2016/01	ASTRAZENECA AB	RO
Seroquel XR 300 mg comprimate cu eliberare prelungita	NL/H/0156/010	9531/2016/06	ASTRAZENECA AB	RO
Seroquel XR 300 mg comprimate cu eliberare prelungita	NL/H/0156/010	9531/2016/02	ASTRAZENECA AB	RO
Seroquel XR 300 mg comprimate cu eliberare prelungita	NL/H/0156/010	9531/2016/07	ASTRAZENECA AB	RO
Seroquel XR 300 mg comprimate cu eliberare prelungita	NL/H/0156/010	9531/2016/03	ASTRAZENECA AB	RO

<b>Product Name (in authorisation country)</b>	<b>MRP/DCP Authorisation number</b>	<b>National Authorisation Number</b>	<b>MAH of product in the member state</b>	<b>Member State where product is authorised</b>
Seroquel XR 300 mg prolonged-release tablets	NL/H/0156/010	PA 1019/019/008	ASTRAZENECA AB	IE
Seroquel XR 300 mg retard tabletta	NL/H/0156/010	OGYI-T-5863/04	ASTRAZENECA KFT.	HU
Seroquel XR 300 mg retard tabletta	NL/H/0156/010	OGYI-T-5863/15	ASTRAZENECA KFT.	HU
Seroquel XR 300 mg retard tabletta	NL/H/0156/010	OGYI-T-5863/12	ASTRAZENECA KFT.	HU
Seroquel XR 300 mg Retardtabletten	NL/H/0156/010	1-27422	ASTRAZENECA OSTERREICH GMBH	AT
Seroquel XR 300 mg tablete s produljenim oslobađanjem	NL/H/0156/010	HR-H-408184728	ASTRAZENECA D.O.O.	HR
Seroquel XR 300 mg tablety s predĺženým uvoľňovaním	NL/H/0156/010	68/0232/07-S	ASTRAZENECA AB	SK
SEROQUEL XR 300 mg δισκία παρατεταμένης αποδέσμευσης	NL/H/0156/010	20358	ASTRAZENECA AB	CY
Seroquel XR 300 mg δισκία παρατεταμένης αποδέσμευσης	NL/H/0156/010	54582/15.06.2017	ASTRAZENECA S.A	GR
Seroquel XR 300 mg, comprimés à libération prolongée	NL/H/0156/010	BE314264	ASTRAZENECA S.A. / N.V.	BE
Seroquel XR 300 mg, comprimés à libération prolongée	NL/H/0156/010	2008040021	ASTRAZENECA S.A. / N.V.	LU
Seroquel XR 300 mg, tabletten met verlengde afgifte	NL/H/0156/010	BE314264	ASTRAZENECA S.A. / N.V.	BE
Seroquel XR 300 mg, tabletten met verlengde afgifte	NL/H/0156/010	RVG 34627	ASTRAZENECA BV	NL

<b>Product Name (in authorisation country)</b>	<b>MRP/DCP Authorisation number</b>	<b>National Authorisation Number</b>	<b>MAH of product in the member state</b>	<b>Member State where product is authorised</b>
Seroquel XR 400 mg comprimate cu eliberare prelungita	NL/H/0156/011	9532/2016/04	ASTRAZENECA AB	RO
Seroquel XR 400 mg comprimate cu eliberare prelungita	NL/H/0156/011	9532/2016/01	ASTRAZENECA AB	RO
Seroquel XR 400 mg comprimate cu eliberare prelungita	NL/H/0156/011	9532/2016/03	ASTRAZENECA AB	RO
Seroquel XR 400 mg comprimate cu eliberare prelungita	NL/H/0156/011	9532/2016/06	ASTRAZENECA AB	RO
Seroquel XR 400 mg comprimate cu eliberare prelungita	NL/H/0156/011	9532/2016/05	ASTRAZENECA AB	RO
Seroquel XR 400 mg comprimate cu eliberare prelungita	NL/H/0156/011	9532/2016/02	ASTRAZENECA AB	RO
Seroquel XR 400 mg comprimate cu eliberare prelungita	NL/H/0156/011	9532/2016/07	ASTRAZENECA AB	RO
Seroquel XR 400 mg prolonged-release tablets	NL/H/0156/011	PA 1019/019/009	ASTRAZENECA AB	IE
Seroquel XR 400 mg retard tableta	NL/H/0156/011	OGYI-T-5863/05	ASTRAZENECA KFT.	HU
Seroquel XR 400 mg retard tableta	NL/H/0156/011	OGYI-T-5863/16	ASTRAZENECA KFT.	HU
Seroquel XR 400 mg retard tableta	NL/H/0156/011	OGYI-T-5863/13	ASTRAZENECA KFT.	HU
Seroquel XR 400 mg Retardtabletten	NL/H/0156/011	1-27419	ASTRAZENECA OSTERREICH GMBH	AT
Seroquel XR 400 mg tablete s produljenim oslobađanjem	NL/H/0156/011	HR-H-789589749	ASTRAZENECA D.O.O.	HR



<b>Product Name (in authorisation country)</b>	<b>MRP/DCP Authorisation number</b>	<b>National Authorisation Number</b>	<b>MAH of product in the member state</b>	<b>Member State where product is authorised</b>
Seroquel XR 400 mg δισκία παρατεταμένης αποδέσμευσης	NL/H/0156/011	20359	ASTRAZENECA AB	CY
Seroquel XR 400 mg δισκία παρατεταμένης αποδέσμευσης	NL/H/0156/011	54583/15.06.2017	ASTRAZENECA S.A	GR
Seroquel XR 400 mg, comprimés à libération prolongée	NL/H/0156/011	BE314282	ASTRAZENECA S.A. / N.V.	BE
Seroquel XR 400 mg, comprimés à libération prolongée	NL/H/0156/011	2008040022	ASTRAZENECA S.A. / N.V.	LU
Seroquel XR 400 mg, tabletten met verlengde afgifte	NL/H/0156/011	BE314282	ASTRAZENECA S.A. / N.V.	BE
Seroquel XR 400 mg, tabletten met verlengde afgifte	NL/H/0156/011	RVG 34628	ASTRAZENECA BV	NL
Seroquel XR 50 mg comprimat cu eliberare prelungită	NL/H/0156/008	9528/2016/07	ASTRAZENECA AB	RO
Seroquel XR 50 mg comprimat cu eliberare prelungită	NL/H/0156/008	9528/2016/05	ASTRAZENECA AB	RO
Seroquel XR 50 mg comprimat cu eliberare prelungită	NL/H/0156/008	9528/2016/04	ASTRAZENECA AB	RO
Seroquel XR 50 mg comprimat cu eliberare prelungită	NL/H/0156/008	9528/2016/01	ASTRAZENECA AB	RO
Seroquel XR 50 mg comprimat cu eliberare prelungită	NL/H/0156/008	9528/2016/06	ASTRAZENECA AB	RO
Seroquel XR 50 mg	NL/H/0156/008	9528/2016/03	ASTRAZENECA AB	RO

<b>Product Name (in authorisation country)</b>	<b>MRP/DCP Authorisation number</b>	<b>National Authorisation Number</b>	<b>MAH of product in the member state</b>	<b>Member State where product is authorised</b>
comprimate cu eliberare prelungită				
Seroquel XR 50 mg comprimate cu eliberare prelungită	NL/H/0156/008	9528/2016/02	ASTRAZENECA AB	RO
Seroquel XR 50 mg prolonged-release tablets	NL/H/0156/008	PA 1019/019/005	ASTRAZENECA AB	IE
Seroquel XR 50 mg retard tabletta	NL/H/0156/008	OGYI-T-5863/02	ASTRAZENECA KFT.	HU
Seroquel XR 50 mg retard tabletta	NL/H/0156/008	OGYI-T-5863/10	ASTRAZENECA KFT.	HU
Seroquel XR 50 mg Retardtabletten	NL/H/0156/008	1-27420	ASTRAZENECA OSTERREICH GMBH	AT
Seroquel XR 50 mg tablete s produjjenim oslobađanjem	NL/H/0156/008	HR-H-955743091	ASTRAZENECA D.O.O.	HR
Seroquel XR 50 mg tablety s predíženým uvolňovaním	NL/H/0156/008	68/0230/07-S	ASTRAZENECA AB	SK
Seroquel XR 50 mg δισκία παρατεταμένης αποδέσμευσης	NL/H/0156/008	20356	ASTRAZENECA AB	CY
Seroquel XR 50 mg δισκία παρατεταμένης αποδέσμευσης	NL/H/0156/008	54579/15.06.2017	ASTRAZENECA S.A	GR
Seroquel XR 50 mg, comprimés à libération prolongée	NL/H/0156/008	BE314221	ASTRAZENECA S.A. / N.V.	BE
Seroquel XR 50 mg, comprimés à libération prolongée	NL/H/0156/008	2008040019	ASTRAZENECA S.A. / N.V.	LU
Seroquel XR 50 mg, tabletten met verlengde afgifte	NL/H/0156/008	BE314221	ASTRAZENECA S.A. / N.V.	BE

<b>Product Name (in authorisation country)</b>	<b>MRP/DCP Authorisation number</b>	<b>National Authorisation Number</b>	<b>MAH of product in the member state</b>	<b>Member State where product is authorised</b>
Seroquel XR 50 mg, tabletten met verlengde afgifte	NL/H/0156/008	RVG 34625	ASTRAZENECA BV	NL
Seroquel XR, 300 mg toimeainet prolongeeritult vabastavad tabletid	NL/H/0156/010	565007	ASTRAZENECA AB	EE
Seroquel XR, 400 mg toimeainet prolongeeritult vabastavad tabletid	NL/H/0156/011	565207	ASTRAZENECA AB	EE
SEROQUEL μικτή συσκευασία έναρξης θεραπείας 3 ημερών (3 Day Starterpack)	NL/H/0156/004	54575/15.06.2017	ASTRAZENECA S.A	GR
SEROQUEL μικτή συσκευασία έναρξης θεραπείας 4 ημερών (4 Day Starterpack)	NL/H/0156/005	54576/15.06.2017	ASTRAZENECA S.A	GR
Seroquel, 100 mg õhukese polümeerikattega tabletid	NL/H/0156/002	257199	ASTRAZENECA AB	EE
Seroquel, 200 mg õhukese polümeerikattega tabletid	NL/H/0156/003	257099	ASTRAZENECA AB	EE
Seroquel, filmovertrukne tabletter	NL/H/0156/005	32337	ASTRAZENECA A/S	DK
Seroquel® 100 mg Filmtabletten	NL/H/0156/002	47291.01.00	ASTRAZENECA GMBH	DE
Seroquel® 200 mg Filmtabletten	NL/H/0156/003	47291.02.00	ASTRAZENECA GMBH	DE
Seroquel® 25 mg	NL/H/0156/001	47291.00.00	ASTRAZENECA GMBH	DE

<b>Product Name (in authorisation country)</b>	<b>MRP/DCP Authorisation number</b>	<b>National Authorisation Number</b>	<b>MAH of product in the member state</b>	<b>Member State where product is authorised</b>
Filmtabletten				
Seroquel® 300 mg Filmtabletten	NL/H/0156/007	47291.04.00	ASTRAZENECA GMBH	DE
Setinin, 300 mg, tabletki powlekane	PL/H/0588/005	16021	+PHARMA ARZNEIMITTEL GMBH	PL
Tomel 150 mg Film-coated tablets	not available	021146	DELORBIS PHARMACEUTICALS LTD	CY
Tomel 300 mg Film-coated tablets	not available	021148	DELORBIS PHARMACEUTICALS LTD	CY
XEROQUEL LP 300 mg, comprimé à libération prolongée	not available	NL 39380	ASTRAZENECA S.A.S.	FR
XEROQUEL LP 400 mg, comprimé à libération prolongée	not available	NL 39381	ASTRAZENECA S.A.S.	FR
XEROQUEL LP 50 mg, comprimé à libération prolongée	not available	NL 39 377	ASTRAZENECA S.A.S.	FR
Квентиакс 150 mg филмирани таблетки	DK/H/1059/003	20100176	KRKA, D.D., NOVO MESTO	BG