



EUROPEAN MEDICINES AGENCY
SCIENCE MEDICINES HEALTH

8 June 2023
EMA/547438/2023
Human Medicines Evaluation Division

List of nationally authorised medicinal products

Active substance(s): drospirenone

Procedure no.: PSUSA/00010853/202211



Product Name (in authorisation country)	MRP/DCP Authorisation number	National Authorisation Number	MAH of product in the member state	Member State where product is authorised
Slinda 4 mg filmtabletta	SE/H/1810/001	OGYI-T-23612/01	EXELTIS MAGYARORSZÁG KFT.	HU
Slinda 4 mg filmtabletta	SE/H/1810/001	OGYI-T-23612/02	EXELTIS MAGYARORSZÁG KFT.	HU
Stelista, 4 mg filmdragerad tablett	SE/H/1810/001	58527	EXELTIS HEALTHCARE S.L	SE
Zlynda 4 mg filmom obalené tablety	SE/H/1810/001	17/0381/19-S	EXELTIS SLOVAKIA	SK
Slinda, 4 mg, tabletki powlekane	SE/H/1810/001	25666	EXELTIS POLAND SP. Z O.O.	PL
Zlynda 4 mg potahované tablety	SE/H/1810/001	17/385/18-C	EXELTIS CZECH S.R.O.	CZ
Slinda 4 mg filmsko obložene tablete	SE/H/1810/001	H/21/02860/001	EXELTIS HEALTHCARE S.L	SI
Slinda 4 mg filmsko obložene tablete	SE/H/1810/001	H/21/02860/002	EXELTIS HEALTHCARE S.L	SI
Slinda 4 mg filmsko obložene tablete	SE/H/1810/001	H/21/02860/003	EXELTIS HEALTHCARE S.L	SI
Slinda 4 mg filmsko obložene tablete	SE/H/1810/001	H/21/02860/004	EXELTIS HEALTHCARE S.L	SI
SLINDA 4 mg comprimidos recubiertos con película	SE/H/1869/001	84603	EXELTIS HEALTHCARE S.L	ES
Drospirenone Exeltis 4 mg filmdragerade tabletter	SE/H/1869/001	58259	EXELTIS HEALTHCARE S.L	SE
Slenma 4 mg filmdragerade tabletter	SE/H/1893/001	58261	EXELTIS HEALTHCARE S.L	SE

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Slinda 4 mg comprimé pelliculé	SE/H/1893/001	BE548284	EXELTIS GERMANY GMBH	BE
Slinda 4 mg filmomhulde tabletten	SE/H/1893/001	BE548284	EXELTIS GERMANY GMBH	BE
Slinda 4 mg Filmtabletten	SE/H/1893/001	BE548284	EXELTIS GERMANY GMBH	BE
SLINDA 4 mg Comprimido revestido por película	SE/H/1893/001	5785829	EXELTIS HEALTHCARE S.L	PT
Lyzbet 4 mg Filmtabletten	SE/H/1893/001	139227	EXELTIS GERMANY GMBH	AT
SLINDA 4 mg Comprimido revestido por película	SE/H/1893/001	5785837	EXELTIS HEALTHCARE S.L	PT
SLINDA 4 mg Comprimido revestido por película	SE/H/1893/001	5785811	EXELTIS HEALTHCARE S.L	PT
Slinda 4 mg Filmtabletten	SE/H/1893/001	7002248.00.00	EXELTIS GERMANY GMBH	DE
SLINDA 4 mg Comprimé pelliculé	SE/H/1893/001	2021040055	EXELTIS GERMANY GMBH	LU
Zlynda 4 mg plēvele dengtos tabletės	SE/H/1893/001	LT/1/21/4721/001	UAB EXELTIS BALTICS	LT
Zlynda 4 mg plēvele dengtos tabletės	SE/H/1893/001	LT/1/21/4721/002	UAB EXELTIS BALTICS	LT
Zlynda 4 mg plēvele dengtos tabletės	SE/H/1893/001	LT/1/21/4721/003	UAB EXELTIS BALTICS	LT
Zlynda 4 mg plēvele dengtos tabletės	SE/H/1893/001	LT/1/21/4721/004	UAB EXELTIS BALTICS	LT
Zlynda 4 mg õhukese polümeerikattega tabletid	SE/H/1893/001	1027821	UAB EXELTIS BALTICS	EE
Zlynda 4 mg apvalkotās tabletes	SE/H/1893/001	21-0082	UAB EXELTIS BALTICS	LV

Product Name (in authorisation country)	MRP/DCP Authorisation number	National Authorisation Number	MAH of product in the member state	Member State where product is authorised
SLYND 4 mg film-coated tablets	SE/H/1893/001	PL 44081/0005	EXELTIS HEALTHCARE S.L	XI
SLINDA 4 mg filmomhulde tabletten	SE/H/1893/001	RVG 127386	EXELTIS HEALTHCARE S.L	NL
Slyndy 4 mg film-coated tablets	SE/H/1893/001	PA22998/002/001	EXELTIS HEALTHCARE S.L	IE
Nusvelta 4 mg filmdragerade tabletter	SE/H/1868/001	58275	EXELTIS HEALTHCARE S.L	SE
SLINDA "4 mg compresse rivestite con film"	SE/H/1868/001	047048020	EXELTIS HEALTHCARE S.L	IT
SLINDA "4 mg compresse rivestite con film"	SE/H/1868/001	047048032	EXELTIS HEALTHCARE S.L	IT
SLINDA "4 mg compresse rivestite con film"	SE/H/1868/001	047048044	EXELTIS HEALTHCARE S.L	IT
SLINDA "4 mg compresse rivestite con film"	SE/H/1868/001	047048018	EXELTIS HEALTHCARE S.L	IT
Slinda, fillovertrukne tabletter	SE/H/1809/001	61678	EXELTIS HEALTHCARE S.L	DK
Slinda 4 mg filmdragerade tabletter	SE/H/1809/001	MTNR: 58256	EXELTIS HEALTHCARE S.L	SE
Slinda 4 mg kalvopäällysteinen tabletti	SE/H/1809/001	36346	EXELTIS HEALTHCARE S.L	FI
Slinda 4 mg filmdrasjerte tabletter	SE/H/1809/001	18-12450	EXELTIS HEALTHCARE S.L	NO
Slinda 4 mg filmuhúðaðar töflur	SE/H/1809/001	IS/1/19/084/01	EXELTIS HEALTHCARE S.L	IS
Droslind 4 mg filmdragerade tabletter	SE/H/1811/001/DC	58258	EXELTIS HEALTHCARE S.L	SE
SLINDA 4 mg, comprimé pelliculé	SE/H/1811/001	34009 301 927 4 0	EXELTIS HEALTHCARE S.L	FR

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SLINDA 4 mg, comprimé pelliculé	SE/H/1811/001	34009 301 927 5 7	EXELTIS HEALTHCARE S.L	FR
SLINDA 4 mg, comprimé pelliculé	SE/H/1811/001	34009 550 698 2 9	EXELTIS HEALTHCARE S.L	FR
SLINDA 4 mg, comprimé pelliculé	SE/H/1811/001	34009 550 698 3 6	EXELTIS HEALTHCARE S.L	FR