



EUROPEAN MEDICINES AGENCY
SCIENCE MEDICINES HEALTH

16 May 2019
EMA/341412/2019
Human Medicines Evaluation Division

List of nationally authorised medicinal products

Active substance: etonogestrel

Procedure no.: PSUSA/00001331/201809



Product full name (in authorisation country)	MRP/DCP Authorisation number	National Authorisation Number	MAH of product in the member state	Member State where product is authorised
Implanon NXT®, 68 mg Implantat zur subkutanen Anwendung	NL/H/0150/001	BE203271	MSD BELGIUM BVBA/SPRL	BE
Nexplanon 68 mg implantat för subdermal användning	NL/H/0150/001	14924	MERCK SHARP & DOHME BV	SE
Implanon NXT, 68 mg implantaat voor subdermaal gebruik	NL/H/0150/001	RVG 21168	NV ORGANON	NL
Implanon NXT, 68 mg Implantat zur subkutanen Anwendung	NL/H/0150/001	44868.00.00	MSD SHARP & DOHME GMBH	DE
Implanon NXT, 68 mg, implant for subdermal use	NL/H/0150/001	PA 1286/50/1	MERCK SHARP & DOHME IRELAND (HUMAN HEALTH) LTD	IE
Nexplanon, 68 mg, implant for subdermal use	NL/H/0150/001	MA058/03401	MERCK SHARP & DOHME LTD.	MT

Product full name (in authorisation country)	MRP/DCP Authorisation number	National Authorisation Number	MAH of product in the member state	Member State where product is authorised
Implanon NXT 68 mg implante para via subcutânea	NL/H/0150/001	2871382	MERCK SHARP & DOHME, LDA.	PT
Implanon NXT, 68 mg implante	NL/H/0150/001	62.628	MERCK SHARP & DOHME DE ESPAÑA, S.A	ES
Nexplanon, 68 mg, implant for subdermal use	NL/H/0150/001	PL 00025/0563	MERCK SHARP & DOHME LTD.	UK
Implanon NXT 68 mg implantát na subkutánne použitie	NL/H/0150/001	17/0011/08-S	MERCK SHARP & DOHME BV	SK
Implanon NXT®, 68 mg implant pour usage sous-cutané	NL/H/0150/001	2003107744	MSD BELGIUM BVBA/SPRL	LU
Nexplanon 68 mg vefjalyf til notkunar undir húð	NL/H/0150/001	IS/1/00/019/01	MERCK SHARP & DOHME BV	IS

Product full name (in authorisation country)	MRP/DCP Authorisation number	National Authorisation Number	MAH of product in the member state	Member State where product is authorised
Nexplanon, implantat til subkutan anvendelse	NL/H/0150/001	30379	MERCK SHARP & DOHME BV	DK
NEXPLANON 68 mg, implant pour usage sous-cutané	NL/H/0150/001	34009 587 005 9 0	MSD FRANCE	FR
NEXPLANON 68 mg, implant pour usage sous-cutané	NL/H/0150/001	34009 351 544 3 9	MSD FRANCE	FR
Nexplanon 68 mg implantat til subkutan bruk	NL/H/0150/001	00-3543	MERCK SHARP & DOHME BV	NO
Nexplanon 68 mg implantaatti, ihon alle	NL/H/0150/001	13811	MERCK SHARP & DOHME BV	FI
Nexplanon, 68 mg implantat, för subdermal användning	NL/H/0150/001	13811	MERCK SHARP & DOHME BV	FI

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Implanon NXT 68 mg Implantat zur subkutanen Anwendung	NL/H/0150/001	1-22964	MERCK SHARP & DOHME GES.M.B.H.	AT
Nexplanon 68 mg implant	NL/H/0150/001	6256/2014/01	MERCK SHARP & DOHME ROMANIA SRL	RO
Nexplanon 68 mg implant	NL/H/0150/001	6256/2014/02	MERCK SHARP & DOHME ROMANIA SRL	RO
Nexplanon, 68 mg implantaat	NL/H/0150/001	806813	MERCK SHARP & DOHME BV	EE
Implanon NXT, 68 mg, implant podskórny	NL/H/0150/001	21319	MSD POLSKA SP. Z O.O.	PL
Nexplanon, 68 mg impianto per uso sottocutaneo	NL/H/0150/001	034352029	MSD ITALIA S.R.L.	IT

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Nexplanon, 68 mg impianto per uso sottocutaneo	NL/H/0150/001	034352017	MSD ITALIA S.R.L.	IT
Implanon NXT®, 68 mg implant pour usage sous-cutané	NL/H/0150/001	BE203271	MSD BELGIUM BVBA/SPRL	BE
Implanon NXT®, 68 mg implantaat voor subdermaal gebruik	NL/H/0150/001	BE203271	MSD BELGIUM BVBA/SPRL	BE