



EUROPEAN MEDICINES AGENCY
SCIENCE MEDICINES HEALTH

05 May 2017
EMA/382258/2015
Human Medicines Evaluation Division

List of nationally authorised medicinal products

Active substance(s): drospirenone / ethinylestradiol

Procedure No.: PSUSA/00010217/201609



Product full name (in authorisation country)	MRP /DCP or CP Authorisation number	National Authorisation Number	MAH of product in the member state	Member State where product is authorised
aida® 0,020 mg/3,0 mg Filmtabletten	NL/H/0703/001	64270.00.00	JENAPHARM GMBH & CO KG	DE
Aliane 0,02 mg/3 mg Filmtabletten	NL/H/0702/001	1-26450	BAYER AUSTRIA GMBH	AT
Aliane 3 mg + 0,02 mg comprimidos revestidos por película	NL/H/0702/001	5866389	LUSAL LDA	PT
Aliane 3 mg + 0,02 mg comprimidos revestidos por película	NL/H/0702/001	5866488	LUSAL LDA	PT
Belanette 0,02 mg/3 mg potahované tablety	NL/H/0702/001	17/197/06-C	BAYER PHARMA AG	CZ
BELANETTE 0,02 mg/3 mg, comprimé pelliculé	NL/H/0702/001	376 418-1	BAYER HEALTHCARE	FR
BELANETTE 0,02 mg/3 mg, comprimé pelliculé	NL/H/0702/001	376 419-8	BAYER HEALTHCARE	FR
Belanette 28, filmomhulde tabletten, 0,02 mg/3 mg	NL/H/705/01	RVG 33189	BAYER BV	NL
Belanette, filmomhulde tabletten, 0,02 mg/3 mg	NL/H/0702/001	RVG 33186	BAYER BV	NL
CLEODETTE 0,02 MG/3 MG, POTAHOVANÉ TABLETY	PT/H/0707/001	17/012/14-C	ACTAVIS GROUP PTC EHF.	CZ
CLEODETTE 0,02 MG/3 MGFILMOM OBALENÉ TABLETY	PT/H/0707/001	17/0066/14-S	ACTAVIS GROUP PTC EHF.	SK
Cleodette 0,03mg/3mg plevele dengtos tabletes	PT/H/0707/002	LT/1/14/3490/006-010	ACTAVIS GROUP PTC EHF.	LT
Cleonita 3 mg/0,02 mg comprimate filmate	PT/H/0707/001	6468/2014/01-05	ACTAVIS GROUP PTC EHF.	RO
CONVULINE 0,03 mg / 3 mg, comprimé pelliculé	NL/H/0218/001	356 169-6	BAYER HEALTHCARE	FR
CONVULINE 0,03 mg/3 mg, comprimé pelliculé	NL/H/0218/01	356 170-4	BAYER HEALTHCARE	FR
Diza 0,02 mg / 3 mg kalvopäällysteiset tabletit	NL/H/2917/001	33348	EXELTIS HEALTHCARE S.L	FI
Diza 0,02 mg/3 mg filmdrasjerte tabletter	NL/H/2917/001	15/04368	EXELTIS HEALTHCARE S.L	NO
Diza 0,02 mg/3 mg, filmdragerade tabletter	NL/H/2917/001	52853	EXELTIS HEALTHCARE S.L	SE
Drosbelalleflex 3 mg /0,02 mg comprimidos recubiertos con película	NL/H/2917/001	80604	EXELTIS HEALTHCARE S.L	ES
Drosiane 0,03 mg/3 mg 21+7 - Filmtabletten	NO/H/0146/002	1-28931	SANDOZ GMBH	AT

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Drosiane mite 0,02 mg/3 mg 21+7 - Filmtabletten	NO/H/0146/001	1-28929	SANDOZ GMBH	AT
Eloine 0,02 mg/ 3 mg, filmuhúðaðar töflur	NL/H/1270/01	IS/1/08/021/01	BAYER AB	IS
Eloine 0,02 mg/3 mg Filmtabletten	NL/H/1270/001	1-27587	BAYER AUSTRIA GMBH	AT
Eloine 0,02 mg/3 mg potahované tablety	NL/H/1270/01	17/315/ 08-C	BAYER PHARMA AG	CZ
Eloine 0,02 mg/3 mg, filmdragerade tabletter	NL/H/1270/01	26273	BAYER AB	SE
Eloine 0.02 mg / 3 mg film coated tablets	NL/H/1270/01	MA513/01301	BAYER PLC	MT
ELOINE 0.02 mg / 3 mg film coated tablets	NL/H/1270/01	PL 00010/0575	BAYER PLC	UK
Eloine 3 mg / 0,02 mg comprimidos recubiertos con película	NL/H/1270/01	70233	BAYER HISPANIA SL	ES
Eloine, fillovertrukne tabletter	NL/H/1270/01	42444	BAYER AB	DK
Ethinylestradiol / Drospirenon 0,03 mg / 3 mg Berlipharm, filmomhulde tabletten	NL/H/0218/001	RVG 25414	BERLIPHARM BV	NL
Ethinylestradiol/Drospirenon 24+4 0,02 mg/3 mg Berlipharm filmomhulde tabletten	NL/H/1270/001	RVG 100827	BERLIPHARM BV	NL
ÉTHINYLESTRADIOL/DROSPIRÉNONE BIOGARAN® 0,02 mg/3 mg, comprimé pelliculé	not available	3400949972142	BIOGARAN	FR
ÉTHINYLESTRADIOL/DROSPIRÉNONE BIOGARAN® 0,02 mg/3 mg, comprimé pelliculé	not available	3400949972081	BIOGARAN	FR
ÉTHINYLESTRADIOL/DROSPIRÉNONE BIOGARANcontinu® 0,02 mg/3 mg, comprimé pelliculé	not available	3400949972951	BIOGARAN	FR
ÉTHINYLESTRADIOL/DROSPIRÉNONE BIOGARANcontinu® 0,02 mg/3 mg, comprimé pelliculé	not available	3400949973033	BIOGARAN	FR
Flexyess 3 mg + 0,02 mg comprimidos revestidos por película	NL/H/2041/001	5559109	BERLEX ESPECIALIDADES FARMACEUTICAS LDA	PT

Product full name (in authorisation country)	MRP / DCP or CP Authorisation number	National Authorisation Number	MAH of product in the member state	Member State where product is authorised
Flexyess 3 mg + 0,02 mg comprimidos revestidos por película	NL/H/2041/001	5559067	BERLEX ESPECIALIDADES FARMACEUTICAS LDA	PT
Flexyess 3 mg + 0,02 mg comprimidos revestidos por película	NL/H/2041/001	5559075	BERLEX ESPECIALIDADES FARMACEUTICAS LDA	PT
Jangee Flexibilis 3 mg/0,02 mg filmtabletta	NL/H/2917/001	OGYI-T-22993/01	EXELTIS MAGYARORSZÁG KFT.	HU
JASMINE 0,03 mg/3 mg, comprimé pelliculé	NL/H/215/01	356 055-0	BAYER HEALTHCARE	FR
JASMINE 0,03 mg/3 mg, comprimé pelliculé	NL/H/215/01	356 056-7	BAYER HEALTHCARE	FR
JASMINELLE 0,02 mg/3 mg, comprimé pelliculé	NL/H/701/01	376 400-5	BAYER HEALTHCARE	FR
JASMINELLE 0,02 mg/3 mg, comprimé pelliculé	NL/H/701/01	376 398-0	BAYER HEALTHCARE	FR
JASMINELLECONTINU 0,02 mg/3 mg, comprimé pelliculé	NL/H/704/001	376 444-2	BAYER HEALTHCARE	FR
JASMINELLECONTINU 0,02 mg/3 mg, comprimé pelliculé	NL/H/704/001	376 445-9	BAYER HEALTHCARE	FR
Linatera 0,02 mg/3 mg filmdragerade tabletter	NL/H/1270/001	24322	BAYER OY	FI
Linatera 0,02 mg/3 mg kalvopäällysteiset tabletit	NL/H/1270/01	24322	BAYER OY	FI
Linatera 3 mg + 0,02 mg comprimidos revestidos por película	NL/H/1270/001	5114145	BERLIFARMA ESPECIALIDADES FARMACEUTICAS LDA	PT
Linatera 3 mg + 0,02 mg comprimidos revestidos por película	NL/H/1270/001	5114152	BERLIFARMA ESPECIALIDADES FARMACEUTICAS LDA	PT
Liofora 0,02 mg / 3 mg kalvopäällysteiset tabletit	NL/H/702/01	21724	BAYER OY	FI
LIOFORA 0,02 mg/3 mg filmdragerade tabletter	NL/H/0702/001	21724	BAYER OY	FI
Liofora 3 mg / 0,02 mg comprimidos recubiertos con película	NL/H/0702/001	68.026	BAYER HISPANIA SL	ES
Liofora Diario 3 mg / 0,02 mg comprimidos recubiertos con película	NL/H/0705/001	68.025	BAYER HISPANIA SL	ES

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Liofora, filmomhulde tabletten, 0,02 mg/3 mg	NL/H/0703/001	RVG 33188	BAYER BV	NL
Naraya Flex, 0,02 mg + 3 mg, tabletki powlekane	NL/H/2917/001	23261	EXELTIS POLAND SP. Z O.O.	PL
Perliq 0,02 mg/3 mg compresse rivestite con film	NL/H/2917/001	044159010	EXELTIS HEALTHCARE S.L	IT
Perliq 0.02 mg/3 mg film coated tablets	NL/H/2917/001	PL 44081/0002	EXELTIS HEALTHCARE S.L	UK
Pernella 0,02 mg/3 mg comprimés pelliculés	NL/H/2917/001	BE485955	EXELTIS GERMANY GMBH	BE
Pernella 0,02 mg/3 mg comprimés pelliculés	NL/H/2917/001	2016060211	EXELTIS GERMANY GMBH	LU
Pernella 0,02 mg/3 mg filmomhulde tabletten	NL/H/2917/001	BE485955	EXELTIS GERMANY GMBH	BE
Petibelle 3 mg + 0,03 mg comprimidos revestidos por película	NL/H/0218/001	3350188	LUSAL LDA	PT
Petibelle 3 mg + 0,03 mg comprimidos revestidos por película	NL/H/0218/001	3350287	LUSAL LDA	PT
Petibelle 3 mg + 0,03 mg comprimidos revestidos por película	NL/H/0218/001	3767787	LUSAL LDA	PT
Petibelle 3 mg + 0,03 mg comprimidos revestidos por película	NL/H/0218/001	3767886	LUSAL LDA	PT
Petibelle® 0,03 mg/3 mg Filmtabletten	NL/H/0218/001	49652.00.00	JENAPHARM GMBH & CO KG	DE
Talia 0,02 mg/3 mg apvalkotās tabletes	NL/H/2917/001	15-0347	UAB EXELTIS BALTICS	LV
Talia 0,02 mg/3 mg filmomhulde tabletten	NL/H/2917/001	RVG 113422	QUALITEC EUROPA, S.L.	NL
Talia 3 mg/0,02 mg õhukese polümeerikattega tabletid	NL/H/2917/001	900515	UAB EXELTIS BALTICS	EE
Velmari 0,02 mg/3 mg comprimidos revestidos por película	NL/H/2917/001	5672860	EXELTIS HEALTHCARE S.L	PT
VELMARI Langzyklus 0,02 mg/ 3 mg Filmtabletten	NL/H/2917/001	89823.00.00	EXELTIS GERMANY GMBH	DE
VELMARI Langzyklus 0,02 mg/3 mg Filmtabletten	NL/H/2917/001	137109	EXELTIS GERMANY GMBH	AT
YADINE	not available	17 / 606 / 00-C	BAYER PHARMA AG	CZ
Yadine 0,03 mg/3 mg filmom obalené tablety	not available	17/0098/02-S	BAYER PHARMA AG	SK

Product full name (in authorisation country)	MRP / DCP or CP Authorisation number	National Authorisation Number	MAH of product in the member state	Member State where product is authorised
Yadine filmtabletta	not available	OGYI-T-7473/02	BAYER PHARMA AG	HU
Yadine filmtabletta	not available	OGYI-T-7473/01	BAYER PHARMA AG	HU
Yarina 0,03 mg/3 mg filmsko obložene tablete	not available	H/01/01685/001	BAYER PHARMA AG	SI
Yarina 3000/ 30 mikrogramu apvalkotās tabletes	not available	00-0463	BAYER PHARMA AG	LV
Yarina 3000/30 mikrogramų plėvele dengtos tabletės	not available	LT/1/01/2564/001	BAYER PHARMA AG	LT
Yarina, 3000 µg/30 µg õhukese polümeerikattega tabletid	not available	342401	BAYER PHARMA AG	EE
Yasmin 0,03 / 3 mg filmuhúðaðar töflur	NL/H/215/01	IS/1/00/013/01	BAYER PHARMA AG	IS
Yasmin 0,03 mg / 3 mg filmom obložene tablete	NL/H/0215/001	HR-H-993460030	BAYER DOO	HR
Yasmin 0,03 mg / 3 mg filmom obložene tablete	NL/H/0215/001	HR-H-993460030	BAYER DOO	HR
Yasmin 0,03 mg / 3 mg filmom obložene tablete	NL/H/0215/001	HR-H-993460030	BAYER DOO	HR
Yasmin 0,03 mg / 3 mg filmom obložene tablete	NL/H/0215/001	HR-H-993460030	BAYER DOO	HR
Yasmin 0,03 mg/3 mg - Filmtabletten	NL/H/0215/001	1-23811	BAYER AUSTRIA GMBH	AT
Yasmin 0,03 mg/3 mg compresse rivestite con film	NL/H/0215/001	035023035	BAYER SPA	IT
Yasmin 0,03 mg/3 mg compresse rivestite con film	NL/H/0215/001	035023011	BAYER SPA	IT
Yasmin 0,03 mg/3 mg compresse rivestite con film	NL/H/0215/001	035023047	BAYER SPA	IT
Yasmin 0,03 mg/3 mg compresse rivestite con film	NL/H/0215/001	035023023	BAYER SPA	IT
Yasmin 0,03 mg/3 mg comprimés pelliculés	NL/H/0215/001	BE218066	BAYER SA NV	BE
Yasmin 0,03 mg/3 mg comprimés pelliculés	NL/H/215/01	2009050310	BAYER SA NV	LU
Yasmin 0,03 mg/3 mg filmdrasjerte tabletter	NL/H/215/01	00-2880	BAYER PHARMA AG	NO
Yasmin 0,03 mg/3 mg filmomhulde tabletten	NL/H/0215/001	BE218066	BAYER SA NV	BE

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Yasmin 0,03 mg/3 mg Filmtabletten	NL/H/0215/001	BE218066	BAYER SA NV	BE
Yasmin 0,03 mg/3 mg Filmtabletten	NL/H/215/01	2009050310	BAYER SA NV	LU
Yasmin 0,03 mg/3 mg kalvopäällysteiset tabletit	NL/H/0215/001	15767	BAYER OY	FI
Yasmin 0,03 mg/3 mg, comprimate filmate	not available	2796/2010/01	BAYER PHARMA AG	RO
Yasmin 0.03 mg / 3 mg film-coated tablets	NL/H/0215/001	PA1410/23/1	BAYER LTD	IE
Yasmin 28 0,03 / 3 mg filmuhúðaðar töflur	NL/H/217/01	IS/1/00/014/01	BAYER PHARMA AG	IS
Yasmin 28 0,03 mg/3mg, filmdrasjerte tabletter	NL/H/217/01	00-2881	BAYER PHARMA AG	NO
Yasmin 28, 0,03 mg/3 mg, filmdragerade tabletter	NL/H/0217/001	16328	BAYER PHARMA AG	SE
Yasmin 28, filmomhulde tabletten, 0,03 mg/3 mg	NL/H/0217/001	RVG 25413	BAYER BV	NL
Yasmin 28, filmovertrukne tabletter	NL/H/217/01	31586	BAYER PHARMA AG	DK
Yasmin 3 mg / 0,03 mg comprimidos recubiertos con película	NL/H/215/01	63.576	BAYER HISPANIA SL	ES
Yasmin 3 mg + 0,03 mg comprimidos revestidos por película	NL/H/215/01	3339181	BAYER PORTUGAL LDA	PT
Yasmin 3 mg + 0,03 mg comprimidos revestidos por película	NL/H/215/01	3339280	BAYER PORTUGAL LDA	PT
Yasmin 3 mg + 0,03 mg comprimidos revestidos por película	NL/H/215/01	3767985	BAYER PORTUGAL LDA	PT
Yasmin 3 mg + 0,03 mg comprimidos revestidos por película	NL/H/215/01	3768082	BAYER PORTUGAL LDA	PT
Yasmin Diario 3 mg / 0,03 mg comprimidos recubiertos con película	NL/H/217/01	63.575	BAYER HISPANIA SL	ES
Yasmin, 0,03 mg + 3,00 mg, tabletki powlekane	not available	11011	BAYER PHARMA AG	PL
Yasmin, 0,03 mg/3 mg, filmdragerade tabletter	NL/H/0215/001	16327	BAYER PHARMA AG	SE
Yasmin, film-coated tablets, 0.03 mg/3 mg	not available	MA 513/03201	BAYER PLC	MT

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Yasmin, film-coated tablets, 0.03 mg/3 mg	NL/H/215/01	PL 00010/0571	BAYER PLC	UK
Yasmin, filmomhulde tabletten, 0,03 mg/3 mg	NL/H/0215/001	RVG 23827	BAYER BV	NL
Yasmin, filmovertrukne tabletter	NL/H/0215/001	31587	BAYER PHARMA AG	DK
Yasmin® 0,03 mg/3 mg filmdragerade tabletter	NL/H/0215/001	15767	BAYER OY	FI
Yasmin® 0,03 mg/3 mg Filmtabletten	NL/H/0215/001	49651.00.00	JENAPHARM GMBH & CO KG	DE
Yasmin® επικαλυμμένο με λεπτό υμένιο δισκίο 0,03 mg / 3 mg	NL/H/0215/001	4016/21-01-2016	BAYER HELLAS SA	GR
Yasminelle 0,02 g/3 mg filmom obalené tablety	NL/H/0701/001	17/0259/06-S	BAYER PHARMA AG	SK
Yasminelle 0,02 mg / 3 mg comprimés pelliculés	NL/H/0701/001	BE288206	BAYER SA NV	BE
Yasminelle 0,02 mg / 3 mg comprimés pelliculés	NL/H/701/01	0442/06110022	BAYER SA NV	LU
Yasminelle 0,02 mg / 3 mg comprimés pelliculés	NL/H/701/01	2006110022	BAYER SA NV	LU
Yasminelle 0,02 mg / 3 mg filmomhulde tabletten	NL/H/0701/001	BE288206	BAYER SA NV	BE
Yasminelle 0,02 mg / 3 mg Filmtabletten	NL/H/0701/001	BE288206	BAYER SA NV	BE
Yasminelle 0,02 mg / 3 mg Filmtabletten	NL/H/701/01	0442/06110022	BAYER SA NV	LU
Yasminelle 0,02 mg / 3 mg Filmtabletten	NL/H/701/01	2006110022	BAYER SA NV	LU
Yasminelle 0,02 mg / 3 mg plėvele dengtos tabletės	NL/H/0701/001	LT/1/06/0520/001	BAYER PHARMA AG	LT
Yasminelle 0,02 mg / 3 mg plėvele dengtos tabletės	NL/H/0701/001	LT/1/06/0520/003	BAYER PHARMA AG	LT
Yasminelle 0,02 mg / 3 mg plėvele dengtos tabletės	NL/H/0701/001	LT/1/06/0520/004	BAYER PHARMA AG	LT
Yasminelle 0,02 mg / 3 mg plėvele dengtos tabletės	NL/H/0701/001	LT/1/06/0520/002	BAYER PHARMA AG	LT
Yasminelle 0,02 mg /3 mg, filmuhúðaðar töflur	NL/H/701/01	IS/1/06/008/01	BAYER PHARMA AG	IS
Yasminelle 0,02 mg/3 mg apvalkotās tabletes	NL/H/0701/001	06-0161	BAYER PHARMA AG	LV

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Yasminelle 0,02 mg/3 mg compresse rivestite con film	NL/H/0701/001	037199039	BAYER SPA	IT
Yasminelle 0,02 mg/3 mg compresse rivestite con film	NL/H/0701/001	037199015	BAYER SPA	IT
Yasminelle 0,02 mg/3 mg compresse rivestite con film	NL/H/0701/001	037199027	BAYER SPA	IT
Yasminelle 0,02 mg/3 mg compresse rivestite con film	NL/H/0701/001	037199041	BAYER SPA	IT
Yasminelle 0,02 mg/3 mg filmdragerade tabletter	NL/H/701/01	21723	BAYER OY	FI
Yasminelle 0,02 mg/3 mg filmsko obložene tablete	NL/H/701/01	H/06/01686/003	BAYER PHARMA AG	SI
Yasminelle 0,02 mg/3 mg filmsko obložene tablete	NL/H/701/01	H/06/01686/002	BAYER PHARMA AG	SI
Yasminelle 0,02 mg/3 mg filmsko obložene tablete	NL/H/701/01	H/06/01686/001	BAYER PHARMA AG	SI
Yasminelle 0,02 mg/3 mg filmsko obložene tablete	NL/H/701/01	H/06/01686/004	BAYER PHARMA AG	SI
Yasminelle 0,02 mg/3 mg Filmtabletten	NL/H/701/01	1-26451	BAYER AUSTRIA GMBH	AT
Yasminelle 0,02 mg/3 mg kalvopäällysteiset tabletit	NL/H/701/01	21723	BAYER OY	FI
Yasminelle 0,02 mg/3 mg potahované tablety	NL/H/701/01	17/192/06-C	BAYER PHARMA AG	CZ
Yasminelle 0.02 mg / 3 mg film-coated tablets	NL/H/701/01	PA 1410/024/001	BAYER LTD	IE
Yasminelle 0.02 mg / 3 mg film-coated tablets	NL/H/701/01	MA639/00101	BAYER LTD	MT
Yasminelle 28, 0,02 mg/3 mg, filmdragerade tabletter	NL/H/704/01	23086	BAYER PHARMA AG	SE
Yasminelle 28, 0,02 mg/3 mg, filmdrasjerte tabletter	NL/H/704/01	05-3805	BAYER PHARMA AG	NO
Yasminelle 28, filmomhulde tabletten, 0,02 mg/3 mg	NL/H/704/01	RVG 33187	BAYER BV	NL
Yasminelle 28, filmovertukne tabletter	NL/H/704/01	38688	BAYER PHARMA AG	DK
Yasminelle 3 mg / 0,02 mg comprimidos recubiertos con película	NL/H/701/01	67.982	BAYER HISPANIA SL	ES

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Yasminelle 3 mg + 0,02 mg comprimidos revestidos por película	NL/H/0701/001	5866280	BERLEX ESPECIALIDADES FARMACEUTICAS LDA	PT
Yasminelle 3 mg + 0,02 mg comprimidos revestidos por película	NL/H/0701/001	5866181	BERLEX ESPECIALIDADES FARMACEUTICAS LDA	PT
Yasminelle Diario 3 mg / 0,02 mg comprimidos recubiertos con película	NL/H/704/01	67.983	BAYER HISPANIA SL	ES
Yasminelle filmtabletta	NL/H/702/01	OGYI-T 20121/01	BAYER PHARMA AG	HU
Yasminelle filmtabletta	NL/H/702/01	OGYI-T 20121/02	BAYER PHARMA AG	HU
Yasminelle, 0,02 mg + 3 mg, tabletki powlekane	NL/H/701/01	12364	BAYER PHARMA AG	PL
Yasminelle, 0,02 mg/3 mg õhukese polümeerikattega tabletid	NL/H/701/01	514706	BAYER PHARMA AG	EE
Yasminelle, 0,02 mg/3 mg, filmdragerade tabletter	NL/H/701/01	23085	BAYER PHARMA AG	SE
Yasminelle, 0,02 mg/3 mg, filmdrasjerte tabletter	NL/H/701/01	05-3792	BAYER PHARMA AG	NO
Yasminelle, filmomhulde tabletten, 0,02 mg/3 mg	NL/H/701/01	RVG 31781	BAYER BV	NL
Yasminelle, filmovertrukne tabletter	NL/H/701/01	38687	BAYER PHARMA AG	DK
Yasminelle® 0,02 mg/3 mg Filmtabletten	NL/H/701/01	64268.00.00	JENAPHARM GMBH & CO KG	DE
Yasminelle® επικαλυμμένα με λεπτό υμένιο δισκία 0,02 mg / 3 mg	NL/H/701/01	20125	BAYER HELLAS SA	CY
Yasminelle® επικαλυμμένα με λεπτό υμένιο δισκία 0,02 mg / 3 mg	NL/H/701/01	4022/21-1-2016	BAYER HELLAS SA	GR
YAZ 0,02 mg / 3 mg apvalkotās tabletes	NL/H/1269/01	08-0167	BAYER PHARMA AG	LV
YAZ 0,02 mg / 3 mg filmom obložene tablete	NL/H/1269/01	HR-H-604240169-01	BAYER DOO	HR
YAZ 0,02 mg / 3 mg filmom obložene tablete	NL/H/1269/01	HR-H-604240169-02	BAYER DOO	HR
YAZ 0,02 mg / 3 mg filmom obložene tablete	NL/H/1269/01	HR-H-604240169-03	BAYER DOO	HR
YAZ 0,02 mg / 3 mg filmom obložene tablete	NL/H/1269/01	HR-H-604240169-04	BAYER DOO	HR
YAZ 0,02 mg / 3 mg, compresse rivestite con film	NL/H/1269/01	038542015	BAYER SPA	IT

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YAZ 0,02 mg / 3 mg, compresse rivestite con film	NL/H/1269/01	038542039	BAYER SPA	IT
YAZ 0,02 mg / 3 mg, compresse rivestite con film	NL/H/1269/01	038542027	BAYER SPA	IT
YAZ 0,02 mg / 3 mg, comprimé pelliculé.	NL/H/1269/01	388 404-0	BAYER HEALTHCARE	FR
YAZ 0,02 mg / 3 mg, comprimé pelliculé.	NL/H/1269/01	388 405-7	BAYER HEALTHCARE	FR
YAZ 0,02 mg / 3 mg, comprimé pelliculé.	NL/H/1269/01	574 189-9	BAYER HEALTHCARE	FR
YAZ 0,02 mg / 3 mg filmom obalené tablety	NL/H/1269/01	17/0217/08-S	BAYER SPOL SRO	SK
YAZ 0,02 mg/ 3 mg, filmuhúðaðar töflur	NL/H/1269/01	IS/1/08/020/01	BAYER AB	IS
YAZ 0,02 mg/3 mg comprimate filmate	NL/H/1269/01	5060/2012/01-03	BAYER PHARMA AG	RO
Yaz 0,02 mg/3 mg comprimés pelliculés	NL/H/1269/01	BE321386	BAYER SA NV	BE
Yaz 0,02 mg/3 mg comprimés pelliculés	NL/H/1269/01	2008090015	BAYER SA NV	LU
Yaz 0,02 mg/3 mg filmomhulde tabletten	NL/H/1269/01	BE321386	BAYER SA NV	BE
YAZ 0,02 mg/3 mg filmsko obložene tablete	NL/H/1269/01	H/08/01688/001	BAYER D.O.O	SI
YAZ 0,02 mg/3 mg filmsko obložene tablete	NL/H/1269/01	H/08/01688/002	BAYER D.O.O	SI
YAZ 0,02 mg/3 mg filmsko obložene tablete	NL/H/1269/01	H/08/01688/004	BAYER D.O.O	SI
YAZ 0,02 mg/3 mg filmsko obložene tablete	NL/H/1269/01	H/08/01688/003	BAYER D.O.O	SI
YAZ 0,02 mg/3 mg Filmtabletten	NL/H/1269/01	1-27586	BAYER AUSTRIA GMBH	AT
Yaz 0,02 mg/3 mg Filmtabletten	NL/H/1269/01	BE321386	BAYER SA NV	BE
Yaz 0,02 mg/3 mg Filmtabletten	NL/H/1269/01	2008090015	BAYER SA NV	LU
Yaz 0,02 mg/3 mg kalvopäällysteiset tabletit	NL/H/1269/01	24321	BAYER OY	FI
YAZ 0,02 mg/3 mg plėvele dengtos tabletės	NL/H/1269/01	LT/1/08/1204/003	BAYER PHARMA AG	LT
YAZ 0,02 mg/3 mg plėvele dengtos tabletės	NL/H/1269/01	LT/1/08/1204/001	BAYER PHARMA AG	LT

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YAZ 0,02 mg/3 mg plèvele dengtos tabletės	NL/H/1269/01	LT/1/08/1204/004	BAYER PHARMA AG	LT
YAZ 0,02 mg/3 mg plèvele dengtos tabletės	NL/H/1269/01	LT/1/08/1204/002	BAYER PHARMA AG	LT
Yaz 0,02 mg/3 mg tabletter, filmbrasjerte	NL/H/1269/01	07-5386	BAYER AB	NO
Yaz 0,02 mg/3 mg, filmdragerade tabletter	NL/H/1269/01	26267	BAYER AB	SE
Yaz 0,02 mg / 3 mg filmdragerade tabletter	NL/H/1269/01	24321	BAYER OY	FI
YAZ 0,02 mg /3 mg potahované tablety	NL/H/1269/001	17/ 316/ 08 - C	BAYER PHARMA AG	CZ
Yaz 0.02 mg / 3 mg film coated tablets	NL/H/1269/001	MA639/00601	BAYER LTD	MT
Yaz 0.02 mg / 3 mg film-coated tablets	NL/H/1269/01	PA1410/056/001	BAYER LTD	IE
YAZ 24+4 0,02 mg/3 mg filmomhulde tabletten	NL/H/1269/001	RVG 33842	BAYER BV	NL
YAZ 3 mg / 0,02 mg comprimidos recubiertos con película	NL/H/1269/01	70093	BAYER HISPANIA SL	ES
Yaz 3 mg + 0,02 mg comprimidos revestidos por película	NL/H/1269/01	5114137	BERLEX ESPECIALIDADES FARMACEUTICAS LDA	PT
Yaz 3 mg + 0,02 mg comprimidos revestidos por película	NL/H/1269/01	5114129	BERLEX ESPECIALIDADES FARMACEUTICAS LDA	PT
Yaz, 0,02 mg + 3,0 mg, tabletki powlekane	NL/H/1269/01	14780	BAYER PHARMA AG	PL
YAZ, 0,02 mg/3 mg õhukese polümeerikattega tabletid	NL/H/1269/01	586208	BAYER PHARMA AG	EE
Yaz, filmovertukne tabletter	NL/H/1269/01	42417	BAYER AB	DK
YAZ® 0,02 mg / 3 mg Filmtabletten	NL/H/1269/01	71898.00.00	JENAPHARM GMBH & CO KG	DE
Yira 3 mg / 0,03 mg comprimidos recubiertos con película	NL/H/218/01	63.577	BAYER HISPANIA SL	ES
Yirala 0,03 mg/3 mg - Filmtabletten	NL/H/218/01	1-23812	BAYER AUSTRIA GMBH	AT
Yvidually 0,02 mg/3 mg, filmomhulde tabletten	NL/H/2041/001	RVG 108807	BAYER BV	NL
Yvidually® 0,02 mg/3 mg Filmtabletten	NL/H/2041/001	83235.00.00	JENAPHARM GMBH & CO KG	DE
Джаз 0,02 mg/3 mg филмирани таблетки	NL/H/1269/01	20090142	BAYER PHARMA AG	BG

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Елоин 0,02 mg/3 mg филмирани таблетки	NL/H/1270/01	20090309	BAYER PHARMA AG	BG
Флексиес 0,02 mg/3 mg филмирани таблетки	NL/H/2041/001	20120567	BAYER PHARMA AG	BG
Ясмин 0,03 mg/3 mg филмирани таблетки	not available	20010803	BAYER PHARMA AG	BG