



EUROPEAN MEDICINES AGENCY
SCIENCE MEDICINES HEALTH

27 June 2024
EMA/391410/2024
Human Medicines Evaluation Division

List of nationally authorised medicinal products

Active substance(s): hydrochlorothiazide / nebivolol

Procedure no.: PSUSA/00001658/202311

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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
Hyporetic 5 mg / 12,5 mg comprimés pelliculés	NL/H/1068/001	BE335973	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	BE
Hyporetic 5 mg / 25 mg comprimés pelliculés	NL/H/1068/002	BE335982	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	BE
HYPORETIC 5 mg / 12,5 mg Filmdabletten	NL/H/1068/001	BE335973	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	BE
HYPORETIC 5 mg / 25 mg Filmdabletten	NL/H/1068/002	BE335982	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	BE
Nomexor® plus HCT 5 mg/12,5 mg Filmdabletten	NL/H/1068/001	1-28414	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	AT
Nomexor® plus HCT 5 mg/25 mg Filmdabletten	NL/H/1068/002	1-28415	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	AT
Hyporetic 5 mg / 12,5 mg filmomhulde tableten	NL/H/1068/001	BE335973	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	BE
Hyporetic 5 mg / 25 mg filmomhulde tableten	NL/H/1068/002	BE335982	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	BE
CONEBILOX 5 mg/12,5 mg, comprimé pelliculé	NL/H/1068/001	34009 393 963 4 7	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	FR
CONEBILOX 5 mg/12,5 mg, comprimé pelliculé	NL/H/1068/001	34009 393 966 3 7	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	FR
CONEBILOX 5 mg/25 mg, comprimé pelliculé	NL/H/1068/002	34009 393 971 7 7	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	FR
CONEBILOX 5 mg/25 mg, comprimé pelliculé	NL/H/1068/002	34009 393 975 2 8	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	FR
Hypoloc-plus 5 mg/12,5 mg επικαλυμμένα με λεπτό υμένιο δισκία.	NL/H/1068/001	76801	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	GR
Hypoloc-plus 5 mg/25 mg επικαλυμμένα με λεπτό υμένιο δισκία.	NL/H/1068/002	76802	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	GR
Nebilet Plus 5 mg / 12.5 mg film-coated tablets	NL/H/1068/001	PA 865/15/1	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IE
LOBIDIUR 5 mg/12,5 mg compresse rivestita con film	NL/H/1068/001	039181033	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT

Product Name (in authorisation country)	MRP/DCP Authorisation number	National Authorisation Number	MAH of product in the member state	Member State where product is authorised
LOBIDIUR 5 mg/12,5 mg compresse rivestite con film	NL/H/1068/001	039181058	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
LOBIDIUR 5 mg/12,5 mg compresse rivestite con film	NL/H/1068/001	039181045	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
Nebile Plus 5 mg / 25 mg film-coated tablets	NL/H/1068/002	PA 865/15/2	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IE
LOBIDIUR 5 mg/25 mg compresse rivestite con film	NL/H/1068/002	039181072	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
LOBIDIUR 5 mg/25 mg compresse rivestite con film	NL/H/1068/002	039181096	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
LOBIDIUR 5 mg/25 mg compresse rivestite con film	NL/H/1068/002	039181084	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
LOBIDIUR 5 mg/25 mg compresse rivestite con film	NL/H/1068/002	039181108	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
LOBIDIUR 5 mg/12,5 mg compresse rivestite con film	NL/H/1068/001	039181060	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
LOBIDIUR 5 mg/25 mg compresse rivestite con film	NL/H/1068/002	039181110	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
LOBIDIUR 5 mg/12,5 mg compresse rivestite con film	NL/H/1068/001	039181019	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
LOBIDIUR 5 mg/25 mg compresse rivestite con film	NL/H/1068/002	039181122	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
LOBIDIUR 5 mg/12,5 mg compresse rivestite con film	NL/H/1068/001	039181021	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
Hyporetic 5 mg / 12,5 mg filmomhulde tabletten	NL/H/1068/001	RVG 35174	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	NL
Hyporetic 5 mg / 25 mg filmomhulde tabletten	NL/H/1068/002	RVG 35175	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	NL
Silostar Plus 5 mg/ 12,5 mg comprimidos recubiertos con película	NL/H/1068/001	71.256	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	ES
Silostar Plus 5 mg/ 25 mg comprimidos recubiertos con película	NL/H/1068/002	71.255	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	ES
Невотенс Плюс 5 mg/12,5 mg	not available	20160398	TEVA PHARMA EAD	BG

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филмирани таблетки				
Hypoloc Comp 5 mg/12,5 mg filmdragerade tabletter	NL/H/1067/001	23313	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	FI
Nobiretic 5 mg / 12,5 mg comprimés pelliculés	NL/H/1067/001	BE335991	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	BE
Nobiretic 5 mg / 25 mg comprimés pelliculés	NL/H/1067/002	BE336007	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	BE
NOBIRETIC 5 mg / 25 mg Filmtabletten	NL/H/1067/002	BE336007	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	BE
NOBIRETIC 5 mg / 12,5 mg Filmtabletten	NL/H/1067/001	BE335991	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	BE
Co-Nebilet 5 mg/12,5 mg comprimate filmate	NL/H/1067/001	5382/2013/01	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	RO
Co-Nebilet 5 mg/12,5 mg comprimate filmate	NL/H/1067/001	5382/2013/03	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	RO
Co-Nebilet 5 mg/12,5 mg comprimate filmate	NL/H/1067/001	5382/2013/02	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	RO
Co-Nebilet 5 mg/25 mg comprimate filmate	NL/H/1067/002	5383/2013/02	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	RO
Co-Nebilet 5 mg/25 mg comprimate filmate	NL/H/1067/002	5383/2013/01	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	RO
Co-Nebilet 5 mg/25 mg comprimate filmate	NL/H/1067/002	5383/2013/03	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	RO
Nobiretic 5 mg / 25 mg comprimés pelliculés	NL/H/1067/002	2009080047	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	LU
Nobiretic 5 mg / 12,5 mg comprimés pelliculés	NL/H/1067/001	2009080046	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	LU
Co-Nebilet 5 mg/12,5 mg filmsko obložene tablete	NL/H/1067/001	H/09/00416/006	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	SI
Co-Nebilet 5 mg/12,5 mg filmsko obložene tablete	NL/H/1067/001	H/09/00416/004	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	SI
Co-Nebilet 5 mg/12,5 mg filmsko obložene tablete	NL/H/1067/001	H/09/00416/003	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	SI
Co-Nebilet 5 mg/12,5 mg filmsko obložene tablete	NL/H/1067/001	H/09/00416/005	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	SI

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Co-Nebilet 5 mg/25 mg filmsko obložene tablete	NL/H/1067/002	H/09/00416/010	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	SI
Co-Nebilet 5 mg/25 mg filmsko obložene tablete	NL/H/1067/002	H/09/00416/009	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	SI
Co-Nebilet 5 mg/25 mg filmsko obložene tablete	NL/H/1067/002	H/09/00416/012	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	SI
Co-Nebilet 5 mg/25 mg filmsko obložene tablete	NL/H/1067/002	H/09/00416/011	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	SI
Небилет плюс 5 mg/12,5 mg филмирани таблетки	NL/H/1067/001	20090096	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	BG
Nobiretic 5 mg / 25 mg filmomhulde tabletten	NL/H/1067/002	BE336007	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	BE
Nobiretic 5 mg / 12,5 mg filmomhulde tabletten	NL/H/1067/001	BE335991	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	BE
Nebilet Plus H 5 mg/12,5 mg potahované tablety	NL/H/1067/001	58/306/09-C	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	CZ
Nebilet Plus H 5 mg/25 mg potahované tablety	NL/H/1067/002	58/307/09-C	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	CZ
Nebilet Plus, 5 mg/25 mg õhukese polümeerikattega tabletid	NL/H/1067/002	625109	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	EE
Hypoloc Comp, filmovertukne tabletter 5 mg/25 mg	NL/H/1067/002	40962	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	DK
Hypoloc Comp, filmovertukne tabletter 5 mg/12,5 mg	NL/H/1067/001	40961	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	DK
Nebilet Plus, 5 mg/12,5 mg õhukese polümeerikattega tabletid	NL/H/1067/001	625209	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	EE
TEMERITDUO 5 mg/12,5 mg, comprimé pelliculé	NL/H/1067/001	34009 393 976 9 6	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	FR
TEMERITDUO 5 mg/25 mg, comprimé pelliculé	NL/H/1067/002	34009 393 978 1 8	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	FR
TEMERITDUO 5 mg/12,5 mg, comprimé pelliculé	NL/H/1067/001	34009 393 977 5 7	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	FR
Hypoloc Comp 5 mg/12,5 mg	NL/H/1067/001	23313	MENARINI INTERNATIONAL	FI

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kalvopäälysteiset tabletit			OPERATIONS LUXEMBOURG S.A.	
TEMERITDUO 5 mg/25 mg, comprimé pelliculé	NL/H/1067/002	34009 393 979 8 6	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	FR
Lobivon-plus 5 mg/12,5 mg επικαλυμμένα με λεπτό υμένιο δισκία.	NL/H/1067/001	76800	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	GR
Lobivon-plus 5 mg/25 mg επικαλυμμένα με λεπτό υμένιο δισκία.	NL/H/1067/002	76799	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	GR
Nebilet Plus 5 mg/12,5 mg filmdoublet	NL/H/1067/001	OGYI-T-21055/03	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	HU
Nebilet Plus 5 mg/25 mg filmdoublet	NL/H/1067/002	OGYI-T-21055/05	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	HU
Nebilet Plus 5 mg/12,5 mg filmdoublet	NL/H/1067/001	OGYI-T-21055/02	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	HU
Hypoloc Plus 5 mg / 12.5 mg film-coated tablets	NL/H/1067/001	PA 865/16/1	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IE
Nebilet Plus 5 mg/25 mg filmdoublet	NL/H/1067/002	OGYI-T-21055/04	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	HU
Nebilet Plus 5 mg/12,5 mg filmdoublet	NL/H/1067/001	OGYI-T-21055/01	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	HU
Nebilet Plus 5 mg/25 mg filmdoublet	NL/H/1067/002	OGYI-T-21055/06	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	HU
Hypoloc Plus 5 mg / 25 mg film-coated tablets	NL/H/1067/002	PA 865/16/2	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IE
ALONEB 5 mg/12,5 mg compresse rivestita con film	NL/H/1067/001	039180031	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
ALONEB 5 mg/12,5 mg compresse rivestite con film	NL/H/1067/001	039180029	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
ALONEB 5 mg/25 mg compresse rivestite con film	NL/H/1067/002	039180082	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
ALONEB 5 mg/12,5 mg compresse rivestite con film	NL/H/1067/001	039180043	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
ALONEB 5 mg/12,5 mg compresse rivestite con film	NL/H/1067/001	039180056	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT

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ALONEB 5 mg/25 mg comprese rivestite con film	NL/H/1067/002	039180118	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
ALONEB 5 mg/25 mg comprese rivestite con film	NL/H/1067/002	039180094	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
ALONEB 5 mg/12,5 mg comprese rivestite con film	NL/H/1067/001	039180068	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
ALONEB 5 mg/12,5 mg comprese rivestite con film	NL/H/1067/001	039180017	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
ALONEB 5 mg/25 mg comprese rivestite con film	NL/H/1067/002	039180070	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
ALONEB 5 mg/25 mg comprese rivestite con film	NL/H/1067/002	039180120	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
Nebilet Plus 5 mg/12,5 mg apvalkotās tabletes	NL/H/1067/001	09-0074	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	LV
ALONEB 5 mg/25 mg comprese rivestite con film	NL/H/1067/002	039180106	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
Nebilet Plus 5 mg/25 mg apvalkotās tabletes	NL/H/1067/002	09-0075	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	LV
Nebilet Plus 5 mg/12,5 mg plėvele dengtos tabletės	NL/H/1067/001	LT/1/09/1554/006	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	LT
Nebilet Plus 5 mg/12,5 mg plėvele dengtos tabletės	NL/H/1067/001	LT/1/09/1554/001	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	LT
Nebilet Plus 5 mg/25 mg plėvele dengtos tabletės	NL/H/1067/002	LT/1/09/1554/010	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	LT
Nebilet Plus 5 mg/25 mg plėvele dengtos tabletės	NL/H/1067/002	LT/1/09/1554/009	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	LT
Nebilet Plus 5 mg/25 mg plėvele dengtos tabletės	NL/H/1067/002	LT/1/09/1554/008	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	LT
Nebilet Plus 5 mg/25 mg plėvele dengtos tabletės	NL/H/1067/002	LT/1/09/1554/011	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	LT
Nebilet Plus 5 mg/12,5 mg plėvele dengtos tabletės	NL/H/1067/001	LT/1/09/1554/004	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	LT
Nebilet Plus 5 mg/12,5 mg plėvele dengtos tabletės	NL/H/1067/001	LT/1/09/1554/003	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	LT
Nebilet Plus 5 mg/12,5 mg	NL/H/1067/001	LT/1/09/1554/005	MENARINI INTERNATIONAL	LT

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plèvele dengtos tabletės			OPERATIONS LUXEMBOURG S.A.	
Nebilet Plus 5 mg/12,5 mg plèvele dengtos tabletės	NL/H/1067/001	LT/1/09/1554/002	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	LT
Nebilet Plus 5 mg/25 mg plèvele dengtos tabletės	NL/H/1067/002	LT/1/09/1554/007	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	LT
Nebilet Plus 5 mg/25 mg plèvele dengtos tabletės	NL/H/1067/002	LT/1/09/1554/012	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	LT
Nebiretic 5 mg / 25 mg filmomhulde tabletten	NL/H/1067/002	RVG 35177	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	NL
Nebilet Plus 5 mg / 25 mg film-coated tablets	NL/H/1067/002	MA204/00203	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	MT
Nebiretic 5 mg / 12,5 mg filmomhulde tabletten	NL/H/1067/001	RVG 35176	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	NL
Nebilet Plus 5 mg / 12.5 mg film-coated tablets	NL/H/1067/001	MA204/00202	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	MT
NEBILET HCT, 5 mg + 25 mg, tabletki powlekane	NL/H/1067/002	16085	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	PL
NEBILET HCT, 5 mg + 12,5 mg, tabletki powlekane	NL/H/1067/001	16084	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	PL
Nebilet HCT 5 mg + 12,5 mg comprimidos revestidos por película	NL/H/1067/001	5400338	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	PT
Nebilet HCT 5 mg + 25 mg comprimidos revestidos por película	NL/H/1067/002	5400312	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	PT
Nebilet HCT 5 mg + 25 mg comprimidos revestidos por película	NL/H/1067/002	5400320	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	PT
Nebilet HCT 5 mg + 12,5 mg comprimidos revestidos por película	NL/H/1067/001	5400346	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	PT
Co-Nebilet 5 mg/12,5 mg filmsko obložene tablete	NL/H/1067/001	H/09/00416/002	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	SI
Nebilet HCTZ 5 mg/25 mg filmom obalené tablety	NL/H/1067/002	58/0203/09-S	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	SK

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Co-Nebilet 5 mg/12,5 mg filmsko obložene tablete	NL/H/1067/001	H/09/00416/001	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	SI
Co-Nebilet 5 mg/25 mg filmsko obložene tablete	NL/H/1067/002	H/09/00416/007	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	SI
Lobivon Plus 5 mg/ 12,5 mg comprimidos recubiertos con película	NL/H/1067/001	71.268	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	ES
Co-Nebilet 5 mg/25 mg filmsko obložene tablete	NL/H/1067/002	H/09/00416/008	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	SI
Nebilet HCTZ 5 mg/12,5 mg filmom obalené tablety	NL/H/1067/001	58/0202/09-S	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	SK
Lobivon Plus 5 mg/ 25 mg comprimidos recubiertos con película	NL/H/1067/002	71.267	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	ES
Lobivon-plus 5 mg/12,5 mg επικαλυμμένα με λεπτό υμένιο δισκία.	NL/H/1067/001	21788	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	CY
Lobivon-plus 5 mg/25 mg επικαλυμμένα με λεπτό υμένιο δισκία.	NL/H/1067/002	21789	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	CY
NOBIZIDE 5 mg/25 mg compresse rivestite con film	NL/H/1069/002	039182086	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
NOBIZIDE 5 mg/25 mg compresse rivestite con film	NL/H/1069/002	039182098	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
NOBIZIDE 5 mg/25 mg compresse rivestite con film	NL/H/1069/002	039182100	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
NOBIZIDE 5 mg/25 mg compresse rivestite con film	NL/H/1069/002	039182074	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
NOBIZIDE 5 mg/25 mg compresse rivestite con film	NL/H/1069/002	039182112	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
NOBIZIDE 5 mg/12,5 mg compresse rivestite con film	NL/H/1069/001	039182047	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
NOBIZIDE 5 mg/12,5 mg compresse rivestite con film	NL/H/1069/001	039182050	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
NOBIZIDE 5 mg/12,5 mg	NL/H/1069/001	039182035	MENARINI INTERNATIONAL	IT

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comprese rivestite con film			OPERATIONS LUXEMBOURG S.A.	
NOBIZIDE 5 mg/12,5 mg comprese rivestite con film	NL/H/1069/001	039182023	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
NOBIZIDE 5 mg/12,5 mg comprese rivestite con film	NL/H/1069/001	039182011	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
NOBIZIDE 5 mg/12,5 mg comprese rivestite con film	NL/H/1069/001	039182062	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
NOBIZIDE 5 mg/25 mg comprese rivestite con film	NL/H/1069/002	039182124	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
Lobiretic 5 mg / 12,5 mg filmomhulde tabletten	NL/H/1069/001	RVG 35172	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	NL
Lobiretic 5 mg / 25 mg filmomhulde tabletten	NL/H/1069/002	RVG 35173	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	NL
Nebile Plus 5 mg/12,5 mg filmom obložene tablete	not available	HR-H-396660874-01	BERLIN-CHEMIE MENARINI HRVATSKA D.O.O.	HR
Nebile Plus 5 mg/25 mg filmom obložene tablete	not available	HR-H-340964911-01	BERLIN-CHEMIE MENARINI HRVATSKA D.O.O.	HR
Nebivololo e Idroclorotiazide Sandoz 5 mg / 12.5 mg comprese rivestite con film	IT/H/0680/001	044999011	SANDOZ S.P.A.	IT
Nebivololo e Idroclorotiazide Sandoz 5 mg / 25 mg comprese rivestite con film	IT/H/0680/002	044999023	SANDOZ S.P.A.	IT
Nebivolol/HCT EG 5 mg/12,5 mg comprimés pelliculés	DE/H/4355/001	BE503315	EUROGENERICS N.V./S.A.	BE
Nebivolol/HCT EG 5 mg/12,5 mg Filmtabletten	DE/H/4355/001	BE503315	EUROGENERICS N.V./S.A.	BE
Nebivolol/HCT EG 5 mg/25 mg Filmtabletten	DE/H/4355/002	BE503333	EUROGENERICS N.V./S.A.	BE
Nebivolol/HCT EG 5 mg/25 mg Filmtabletten	DE/H/4355/002	BE503342	EUROGENERICS N.V./S.A.	BE
Nebivolol/HCT EG 5 mg/12,5 mg Filmtabletten	DE/H/4355/001	BE503324	EUROGENERICS N.V./S.A.	BE
Nebivolol/HCT EG 5 mg/12,5 mg filmomhulde tabletten	DE/H/4355/001	BE503315	EUROGENERICS N.V./S.A.	BE

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Nebivolol/HCT EG 5 mg/12,5 mg comprimés pelliculés	DE/H/4355/001	BE503324	EUROGENERICS N.V./S.A.	BE
Nebivolol/HCT EG 5 mg/25 mg comprimés pelliculés	DE/H/4355/002	BE503342	EUROGENERICS N.V./S.A.	BE
Nebivolol/HCT EG 5 mg/25 mg comprimés pelliculés	DE/H/4355/002	BE503333	EUROGENERICS N.V./S.A.	BE
Nebivolol/HCT EG 5 mg/12,5 mg filmomhulde tabletten	DE/H/4355/001	BE503324	EUROGENERICS N.V./S.A.	BE
Nebivolol/HCT EG 5 mg / 25 mg filmomhulde tabletten	DE/H/4355/002	BE503333	EUROGENERICS N.V./S.A.	BE
Nebivolol/HCT EG 5 mg / 25 mg filmomhulde tabletten	DE/H/4355/002	BE503342	EUROGENERICS N.V./S.A.	BE
NEBIVOLOLO E IDROCLOROTIAZIDE EG 5 mg/12,5 mg compresse rivestite con film	DE/H/4355/001	044321014	EG S.P.A.	IT
NEBIVOLOLO E IDROCLOROTIAZIDE EG 5 mg/12,5 mg compresse rivestite con film	DE/H/4355/001	044321139	EG S.P.A.	IT
NEBIVOLOLO E IDROCLOROTIAZIDE EG 5 mg/12,5 mg compresse rivestite con film	DE/H/4355/001	044321180	EG S.P.A.	IT
NEBIVOLOLO E IDROCLOROTIAZIDE EG 5 mg/12,5 mg compresse rivestite con film	DE/H/4355/001	044321026	EG S.P.A.	IT
NEBIVOLOLO E IDROCLOROTIAZIDE EG 5 mg/12,5 mg compresse rivestite con film	DE/H/4355/001	044321154	EG S.P.A.	IT
NEBIVOLOLO E IDROCLOROTIAZIDE EG 5 mg/12,5 mg compresse	DE/H/4355/001	044321141	EG S.P.A.	IT

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rivestite con film				
NEBIVOLOLO E IDROCLOROTIAZIDE EG 5 mg/12,5 mg compresse rivestite con film	DE/H/4355/001	044321127	EG S.P.A.	IT
NEBIVOLOLO E IDROCLOROTIAZIDE EG 5 mg/12,5 mg compresse rivestite con film	DE/H/4355/001	044321040	EG S.P.A.	IT
NEBIVOLOLO E IDROCLOROTIAZIDE EG 5 mg/12,5 mg compresse rivestite con film	DE/H/4355/001	044321115	EG S.P.A.	IT
NEBIVOLOLO E IDROCLOROTIAZIDE EG 5 mg/12,5 mg compresse rivestite con film	DE/H/4355/001	044321204	EG S.P.A.	IT
NEBIVOLOLO E IDROCLOROTIAZIDE EG 5 mg/12,5 mg compresse rivestite con film	DE/H/4355/001	044321038	EG S.P.A.	IT
NEBIVOLOLO E IDROCLOROTIAZIDE EG 5 mg/12,5 mg compresse rivestite con film	DE/H/4355/001	044321178	EG S.P.A.	IT
NEBIVOLOLO E IDROCLOROTIAZIDE EG 5 mg/12,5 mg compresse rivestite con film	DE/H/4355/001	044321166	EG S.P.A.	IT
NEBIVOLOLO E IDROCLOROTIAZIDE EG 5 mg/12,5 mg compresse rivestite con film	DE/H/4355/001	044321192	EG S.P.A.	IT
NEBIVOLOLO E IDROCLOROTIAZIDE EG 5	DE/H/4355/001	044321053	EG S.P.A.	IT

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mg/12,5 mg compresse rivestite con film				
Nebivolol/HCT STADA Arzneimittel AG 5 mg/12,5 mg Filmtabletten	DE/H/4355/001	94546.00.00	STADA ARZNEIMITTEL AG	DE
Nebivolol/HCT STADA Arzneimittel AG 5 mg/25 mg Filmtabletten	DE/H/4355/002	94547.00.00	STADA ARZNEIMITTEL AG	DE
NEBIVOLOLO E IDROCLOROTIAZIDE EG 5 mg/25 mg compresse rivestite con film	DE/H/4355/002	044321279	EG S.P.A.	IT
NEBIVOLOLO E IDROCLOROTIAZIDE EG 5 mg/25 mg compresse rivestite con film	DE/H/4355/002	044321293	EG S.P.A.	IT
NEBIVOLOLO E IDROCLOROTIAZIDE EG 5 mg/25 mg compresse rivestite con film	DE/H/4355/002	044321281	EG S.P.A.	IT
NEBIVOLOLO E IDROCLOROTIAZIDE EG 5 mg/25 mg compresse rivestite con film	DE/H/4355/002	044321255	EG S.P.A.	IT
NEBIVOLOLO E IDROCLOROTIAZIDE EG 5 mg/25 mg compresse rivestite con film	DE/H/4355/002	044321305	EG S.P.A.	IT
NEBIVOLOLO E IDROCLOROTIAZIDE EG 5 mg/25 mg compresse rivestite con film	DE/H/4355/002	044321228	EG S.P.A.	IT
NEBIVOLOLO E IDROCLOROTIAZIDE EG 5 mg/25 mg compresse rivestite	DE/H/4355/002	044321267	EG S.P.A.	IT

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con film				
NEBIVOLOLO E IDROCLOROTIAZIDE EG 5 mg/25 mg compresse rivestite con film	DE/H/4355/002	044321103	EG S.P.A.	IT
NEBIVOLOLO E IDROCLOROTIAZIDE EG 5 mg/25 mg compresse rivestite con film	DE/H/4355/002	044321065	EG S.P.A.	IT
NEBIVOLOLO E IDROCLOROTIAZIDE EG 5 mg/25 mg compresse rivestite con film	DE/H/4355/002	044321230	EG S.P.A.	IT
NEBIVOLOLO E IDROCLOROTIAZIDE EG 5 mg/25 mg compresse rivestite con film	DE/H/4355/002	044321216	EG S.P.A.	IT
NEBIVOLOLO E IDROCLOROTIAZIDE EG 5 mg/25 mg compresse rivestite con film	DE/H/4355/002	044321089	EG S.P.A.	IT
NEBIVOLOLO E IDROCLOROTIAZIDE EG 5 mg/25 mg compresse rivestite con film	DE/H/4355/002	044321242	EG S.P.A.	IT
NEBIVOLOLO E IDROCLOROTIAZIDE EG 5 mg/25 mg compresse rivestite con film	DE/H/4355/002	044321091	EG S.P.A.	IT
NEBIVOLOLO E IDROCLOROTIAZIDE EG 5 mg/25 mg compresse rivestite con film	DE/H/4355/002	044321077	EG S.P.A.	IT
Nebivolol/HCT EG 5 mg/25 mg comprimés pelliculés	DE/H/4355/002	2017070238	EUROGENERICS N.V./S.A.	LU

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Nebivolol/HCT EG 5 mg/12,5 mg comprimés pelliculés	DE/H/4355/001	2017070237	EUROGENERICS N.V./S.A.	LU
NEBIVOLOLO e IDROCLOROTIAZIDE DOC Generici 5 mg/12,5 mg compresse rivestite con film	IT/H/0567/001-002/DC	044322016	DOC GENERICI S.R.L.	IT
NEBIVOLOLO e IDROCLOROTIAZIDE DOC Generici 5 mg/25 mg compresse rivestite con film	IT/H/0567/001-002/DC	044322028	DOC GENERICI S.R.L.	IT