



EUROPEAN MEDICINES AGENCY
SCIENCE MEDICINES HEALTH

10 June 2021
EMA/356899/2021
Human Medicines Evaluation Division

List of nationally authorised medicinal products

Active substance: prulifloxacin

Procedure no.: PSUSA/00002569/202010



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
Oliflox 600 mg comprimidos revestidos por película	IT/H/0118/001	5429386	BGP PRODUCTS UNIPESOAL, LDA.	PT
Oliflox 600 mg comprimidos revestidos por película	IT/H/0118/001	5429485	BGP PRODUCTS UNIPESOAL, LDA.	PT
Oliflox 600 mg comprimidos revestidos por película	IT/H/0118/001	5468087	BGP PRODUCTS UNIPESOAL, LDA.	PT
Oliflox 600 mg comprimidos revestidos por película	IT/H/0118/001	5429584	BGP PRODUCTS UNIPESOAL, LDA.	PT
Glimbax 600 mg επικαλυμμένα με λεπτό υμένιο δισκία	IT/H/118/01/MR	107756/14	MEDITRINA PHARMACEUTICALS LTD	GR
KERAFLOX, 600 mg, comprimidos revestidos por película	IT/H/0119/001	5429089	AZIENDE CHIMICHE RIUNITE ANGELINI FRANCESCO - A.C.R.A.F. S.P.A.	PT
KERAFLOX, 600 mg, comprimidos revestidos por película	IT/H/0119/001	5467980	AZIENDE CHIMICHE RIUNITE ANGELINI FRANCESCO - A.C.R.A.F. S.P.A.	PT
KERAFLOX, 600 mg, comprimidos revestidos por película	IT/H/0119/001	5429188	AZIENDE CHIMICHE RIUNITE ANGELINI FRANCESCO - A.C.R.A.F. S.P.A.	PT
KERAFLOX, 600 mg, comprimidos revestidos por película	IT/H/0119/001	5429287	AZIENDE CHIMICHE RIUNITE ANGELINI FRANCESCO - A.C.R.A.F. S.P.A.	PT
Keraflox 600 mg compresse rivestite con film	IT/H/0119/001	035680014	EG S.P.A.	IT
Keraflox 600 mg compresse rivestite con	IT/H/0119/001	035680026	EG S.P.A.	IT

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film.				
Keraflox 600 mg compresse rivestite con film	IT/H/0119/001	035680038	EG S.P.A.	IT
Keraflox 600 mg compresse rivestite con film	IT/H/0119/001	035680040	EG S.P.A.	IT
Unidrox 600 mg compresse rivestite con film.	IT/H/0117/001	035678034	AZIENDE CHIMICHE RIUNITE ANGELINI FRANCESCO - A.C.R.A.F. S.P.A.	IT
Unidrox 600 mg compresse rivestite con film.	IT/H/0117/001	035678046	AZIENDE CHIMICHE RIUNITE ANGELINI FRANCESCO - A.C.R.A.F. S.P.A.	IT
Unidrox 600 mg compresse rivestite con film.	IT/H/0117/001	035678010	AZIENDE CHIMICHE RIUNITE ANGELINI FRANCESCO - A.C.R.A.F. S.P.A.	IT
Unidrox 600 mg compresse rivestite con film.	IT/H/0117/001	035678022	AZIENDE CHIMICHE RIUNITE ANGELINI FRANCESCO - A.C.R.A.F. S.P.A.	IT
Unidrox 600 mg potahované tablety	IT/H/0117/001	42/208/05-C	ANGELINI PHARMA ČESKÁ REPUBLIKA S.R.O.	CZ
Unidrox 600 mg Filmdabletten	IT/H/0117/001	1-26053	ANGELINI PHARMA ÖSTERREICH GMBH	AT
Unidrox 600 mg Filmdabletten	IT/H/0117/001	1-26053	ANGELINI PHARMA ÖSTERREICH GMBH	AT
Unidrox 600 mg Filmdabletten	IT/H/0117/001	1-26053	ANGELINI PHARMA ÖSTERREICH GMBH	AT
Unidrox 600 mg Filmdabletten	IT/H/0117/001	1-26053	ANGELINI PHARMA ÖSTERREICH GMBH	AT
Prixina 600 mg δισκία επικαλυμμένα με υμένιο	IT/H/0117/001	29906/16/29-9-2017	ANGELINI PHARMA HELLAS S.A.	GR

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Unidrox, 600 mg, comprimidos revestidos por película	IT/H/0117/001	5428784	ANGELINI FARMACĚUTICA, LDA	PT
Unidrox, 600 mg, comprimidos revestidos por película	IT/H/0117/001	5467881	ANGELINI FARMACĚUTICA, LDA	PT
Unidrox, 600 mg, comprimidos revestidos por película	IT/H/0117/001	5428883	ANGELINI FARMACĚUTICA, LDA	PT
Unidrox, 600 mg, comprimidos revestidos por película	IT/H/0117/001	5428982	ANGELINI FARMACĚUTICA, LDA	PT
Unidrox 600 mg potahované tablety	IT/H/0117/001	42/208/05-C	ANGELINI PHARMA ĀESKĀ REPUBLIKA S.R.O.	CZ
Unidrox 600 mg potahované tablety	IT/H/0117/001	42/208/05-C	ANGELINI PHARMA ĀESKĀ REPUBLIKA S.R.O.	CZ
Unidrox 600 mg potahované tablety	IT/H/0117/001	42/208/05-C	ANGELINI PHARMA ĀESKĀ REPUBLIKA S.R.O.	CZ
Prixina 600 mg δισκία επικαλυμμένα με υμένιο	IT/H/0117/001	29906/16/29-9-2017	ANGELINI PHARMA HELLAS S.A.	GR
Prixina 600 mg δισκία επικαλυμμένα με υμένιο	IT/H/0117/001	29906/16/29-9-2017	ANGELINI PHARMA HELLAS S.A.	GR
Prixina 600 mg δισκία επικαλυμμένα με υμένιο	IT/H/0117/001	29906/16/29-9-2017	ANGELINI PHARMA HELLAS S.A.	GR
Chinoplus 600 mg compresse rivestite con film.	IT/H/0118/001	035679012	SPA – SOCIETĀ PRODOTTI ANTIBIOTICI S.P.A.	IT
Chinoplus 600 mg compresse rivestite con film.	IT/H/0118/001	035679024	SPA – SOCIETĀ PRODOTTI ANTIBIOTICI S.P.A.	IT
Chinoplus 600 mg compresse rivestite con film.	IT/H/0118/001	035679036	SPA – SOCIETĀ PRODOTTI ANTIBIOTICI S.P.A.	IT
Chinoplus 600 mg compresse rivestite con	IT/H/0118/001	035679048	SPA – SOCIETĀ PRODOTTI ANTIBIOTICI S.P.A.	IT

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film.				