



EUROPEAN MEDICINES AGENCY
SCIENCE MEDICINES HEALTH

21 February 2018
EMA/208334/2018
Human Medicines Evaluation Division

List of nationally authorised medicinal products

Active substance: ibuprofen / pseudoephedrine

Procedure no.: PSUSA/00001711/201707



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
BoxaGrippal 200 mg/30 mg - Filmtabletten	UK/H/4352/001	1-31668	SANOFI-AVENTIS GMBH OSTERREICH	AT
Dolorflu akut 200 mg/30 mg - Filmtabletten	DE/H/4184/001	136793	SANDOZ GMBH	AT
Ibuduo, 200 mg/30 mg film-coated tablets	AT/H/0614/001	137621	ORIFARM GENERICS A/S	AT
WICK DayMed Duo 200mg/30mg Filmtabletten	IT/H/0331/001	135488	RATIOPHARM ARZNEIMITTEL VERTRIEBS-GMBH	AT
Ibuprofen/Pseudoephedrine HCl Pfizer 200 mg/30 mg comprimés enrobés	IE/H/0420/001	BE463680	PFIZER B.V.	BE
Ibuprofen/Pseudoephedrine HCl Pfizer 200 mg/30 mg omhulde tabletten	IE/H/0420/001	BE463680	PFIZER B.V.	BE
Ibuprofen/Pseudoephedrine HCl Pfizer 200mg/30 mg überzogene Tabletten	IE/H/0420/001	BE463680	PFIZER B.V.	BE
Ibuprofen/Pseudoephedrine hydrochloride Boehringer Ingelheim 200 mg/30 mg Filmtabletten	UK/H/4352/001	BE445164	BOEHRINGER INGELHEIM INTERNATIONAL GMBH	BE
Ibuprofen/Pseudoephedrine hydrochloride Boehringer Ingelheim 200 mg/30 mg filmomhulde tabletten	UK/H/4352/001	BE445164	BOEHRINGER INGELHEIM INTERNATIONAL GMBH	BE
IBUPROFEN/PSEUDOEPHEDRINE HYDROCHLORIDE SANOFI 200 mg/30 mg comprimés pelliculés	UK/H/4352/001	BE445164	SANOFI BELGIUM	BE
Ibuprofen/pseudoephedrine hydrochloride Sanofi 200 mg/30 mg filmomhulde tabletten	UK/H/4352/001	BE445164	SANOFI BELGIUM	BE

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Ibuprofen/pseudoephedrine hydrochloride Sanofi 200 mg/30 mg Filmtabletten	UK/H/4352/001	BE445164	SANOFI BELGIUM	BE
Sinuphen 200 mg/30 mg comprimés pelliculés	DE/H/4183/001/DC	BE484782	JOHNSON & JOHNSON CONSUMER N.V./S.A.	BE
Sinuphen 200 mg/30 mg comprimés pelliculés	DE/H/4183/001/DC	BE484782	JOHNSON & JOHNSON CONSUMER N.V./S.A.	BE
Sinuphen 200 mg/30 mg comprimés pelliculés	DE/H/4183/001/DC	BE484782	JOHNSON & JOHNSON CONSUMER N.V./S.A.	BE
Sinuphen 200 mg/30 mg comprimés pelliculés	DE/H/4183/001/DC	BE484782	JOHNSON & JOHNSON CONSUMER N.V./S.A.	BE
Sinuphen 200 mg/30 mg filmomhulde tabletten	DE/H/4183/001/DC	BE484782	JOHNSON & JOHNSON CONSUMER N.V./S.A.	BE
Sinuphen 200 mg/30 mg filmomhulde tabletten	DE/H/4183/001/DC	BE484782	JOHNSON & JOHNSON CONSUMER N.V./S.A.	BE
Sinuphen 200 mg/30 mg filmomhulde tabletten	DE/H/4183/001/DC	BE484782	JOHNSON & JOHNSON CONSUMER N.V./S.A.	BE
Sinuphen 200 mg/30 mg filmomhulde tabletten	DE/H/4183/001/DC	BE484782	JOHNSON & JOHNSON CONSUMER N.V./S.A.	BE
Sinuphen 200 mg/30 mg Filmtabletten	DE/H/4183/001/DC	BE484782	JOHNSON & JOHNSON CONSUMER N.V./S.A.	BE
Sinuphen 200 mg/30 mg Filmtabletten	DE/H/4183/001/DC	BE484782	JOHNSON & JOHNSON CONSUMER N.V./S.A.	BE
Sinuphen 200 mg/30 mg Filmtabletten	DE/H/4183/001/DC	BE484782	JOHNSON & JOHNSON CONSUMER N.V./S.A.	BE
ЕКСТРАЛГИН КОЛД 200 mg/30 mg филмирани таблетки	not available	20100639	ZENTIVA, K.S.	BG
ЕКСТРАЛГИН КОЛД 200 mg/30 mg филмирани таблетки	not available	20100639	ZENTIVA, K.S.	BG
ЕКСТРАЛГИН КОЛД 200 mg/30 mg филмирани таблетки	not available	20100639	ZENTIVA, K.S.	BG

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ЕКСТРАЛГИН КОЛД 200 mg/30 mg филмирани таблетки	not available	20100639	ZENTIVA, K.S.	BG
Ибупром Синус 200 mg/30 mg обвити таблетки	not available	20080222	US PHARMACIA SP. Z O.O.	BG
Мукогрип 200 mg/30 mg филмирани таблетки	UK/H/4352/001	20120408	BOEHRINGER INGELHEIM INTERNATIONAL GMBH	BG
Нурофен Стопколд 200 mg/30 mg филмирани таблетки	not available	9800356	RECKITT BENCKISER HEALTHCARE INTERNATIONAL LIMITED	BG
Олитабс 200 mg/30 mg филмирани таблетки	DE/H/4183/001/DC	20160014	MCNEIL PRODUCTS LIMITED	BG
Олитабс 200 mg/30 mg филмирани таблетки	DE/H/4183/001/DC	20160014	MCNEIL PRODUCTS LIMITED	BG
Олитабс 200 mg/30 mg филмирани таблетки	DE/H/4183/001/DC	20160014	MCNEIL PRODUCTS LIMITED	BG
Олитабс 200 mg/30 mg филмирани таблетки	DE/H/4183/001/DC	20160014	MCNEIL PRODUCTS LIMITED	BG
Hexatab Headcold Relief 200 mg/30 mg, επικαλυμμένα με λεπτό υμένιο δισκία	DE/H/4183/001	022528	JOHNSON & JOHNSON HELLAS CONSUMER AE	CY
Hexatab Headcold Relief 200 mg/30 mg, επικαλυμμένα με λεπτό υμένιο δισκία	DE/H/4183/001	022528	JOHNSON & JOHNSON HELLAS CONSUMER AE	CY
Hexatab Headcold Relief 200 mg/30 mg, επικαλυμμένα με λεπτό υμένιο δισκία	DE/H/4183/001	022528	JOHNSON & JOHNSON HELLAS CONSUMER AE	CY
Hexatab Headcold Relief 200 mg/30 mg, επικαλυμμένα με λεπτό υμένιο δισκία	DE/H/4183/001	022528	JOHNSON & JOHNSON HELLAS CONSUMER AE	CY

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Mucocold 200mg/30mg επικαλυμμένα με λεπτό υμένιο δισκία	UK/H/4352/001	21691	BOEHRINGER INGELHEIM INTERNATIONAL GMBH	CY
NUROFEN Cold & Flu	not available	19671	RECKITT BENCKISER HELLAS CHEMICALS ABEE	CY
Grippecton 200 mg/30 mg potahované tablety	DE/H/4188/001	07/155/16-C	KREWEL MEUSELBACH GMBH	CZ
Modafen potahované tablety	not available	07/136/98-C	ZENTIVA, K.S.	CZ
Modafen potahované tablety	not available	07/136/98-C	ZENTIVA, K.S.	CZ
Modafen potahované tablety	not available	07/136/98-C	ZENTIVA, K.S.	CZ
Modafen potahované tablety	not available	07/136/98-C	ZENTIVA, K.S.	CZ
Nurofen StopGrip potahované tablety	not available	07/612/96-C	RECKITT BENCKISER HEALTHCARE INTERNATIONAL LIMITED	CZ
Robicold 200 mg/30 mg obalené tablety	IE/H/0420/001	07/040/15-C	PFIZER CORPORATION AUSTRIA GESELLSCHAFT M.B.H.	CZ
BoxaGrippal Erkältungssaft 200 mg/10 ml + 30 mg/10 ml Suspension zum Einnehmen	UK/H/5545/001	90668.00.00	SANOFI-AVENTIS DEUTSCHLAND GMBH	DE
BoxaGrippal Erkältungstabletten 200 mg / 30 mg Filmtabletten	UK/H/4352/001	85528.00.00	SANOFI-AVENTIS DEUTSCHLAND GMBH	DE
Dolormin Grippal 200 mg/30 mg Filmtabletten	DE/H/4182/001/DC	93267.00.00	JOHNSON & JOHNSON GMBH	DE
Dolormin Grippal 200 mg/30 mg Filmtabletten	DE/H/4182/001/DC	93267.00.00	JOHNSON & JOHNSON GMBH	DE
Dolormin Grippal 200 mg/30 mg Filmtabletten	DE/H/4182/001/DC	93267.00.00	JOHNSON & JOHNSON GMBH	DE
GRIPOMED 200 mg/30 mg Filmtabletten	DE/H/4185/001	93270.00.00	PHARMASWISS ČESKÁ REPUBLIKA S.R.O.	DE

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IbuHEXAL Grippal 200 mg/30 mg Filmtabletten	DE/H/4184/001	93269.00.00	HEXAL AG	DE
Ibuprofen/Pseudoephedrinhydrochlorid Angelini 200 mg/30 mg Filmtabletten	DE/H/4186/001	93271.00.00	AZIENDE CHIMICHE RIUNITE ANGELINI FRANCESCO - A.C.R.A.F. S.P.A.	DE
Ibuprofen/Pseudoephedrinhydrochlorid Krewel 200 mg/30 mg Filmtabletten	DE/H/4188/001	93273.00.00	KREWEL MEUSELBACH GMBH	DE
IbuSinex 200 mg/30 mg Filmtabletten	DE/H/4187/001	93272.00.00	DIAPHARM GMBH & CO. KG	DE
Olytabs 200 mg/30 mg Filmtabletten	DE/H/4183/001/DC	93268.00.00	JOHNSON & JOHNSON GMBH	DE
Olytabs 200 mg/30 mg Filmtabletten	DE/H/4183/001/DC	93268.00.00	JOHNSON & JOHNSON GMBH	DE
Olytabs 200 mg/30 mg Filmtabletten	DE/H/4183/001/DC	93268.00.00	JOHNSON & JOHNSON GMBH	DE
Olytabs 200 mg/30 mg Filmtabletten	DE/H/4183/001/DC	93268.00.00	JOHNSON & JOHNSON GMBH	DE
ratioGrippal 200 mg/30 mg Filmtabletten	not available	92698.00.00	RATIOPHARM GMBH	DE
Spalt Effekt 30mg/200 mg überzogene Tabletten	FR/H/0238/001	62958.00.00	PFIZER CONSUMER HEALTHCARE GMBH	DE
SpaltGrippal 200 mg/30 mg Weichkapseln	UK/H/5671/001	94021.00.00	PFIZER CONSUMER HEALTHCARE GMBH	DE
WICK DuoGrippal 200 mg/30 mg Filmtabletten	IT/H/0331/001	88707.00.00	WICK PHARMA ZWEIGNIEDERLASSUNG DER PROCTER & GAMBLE GMBH	DE
Febrilek 200 mg / 30 mg õhukese polümeerikattega tabletid	DE/H/4185/001	899915	PHARMASWISS ČESKÁ REPUBLIKA S.R.O.	EE
Bisolfren 200 mg/30 mg comprimidos recubiertos con película	UK/H/4352/001	76389	BOEHRINGER INGELHEIM INTERNATIONAL GMBH	ES

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
Gripaduo 200 mg/30 mg comprimidos recubiertos con película	IT/H/0331/001	78831	LABORATORIOS VICKS, S.L.	ES
NUROFEN RHUME, comprimé pelliculé	not available	343 625-8	RECKITT BENCKISER HEALTHCARE FRANCE	FR
RHINADVIL RHUME IBUPROFENE/PSEUDOPHE DRINE, comprimé enrobé	FR/H/0238/001	334 084-8	PFIZER SANTE FAMILIALE SAS	FR
RHINUREFLEX, comprimé pelliculé	not available	338 856-5	RECKITT BENCKISER HEALTHCARE FRANCE	FR
Gripomed 200 mg/30 mg επικαλυμμένα με λεπτό υμένιο δισκία	DE/H/4185/001	78858/14/14.09.16	PHARMASWISS ČESKÁ REPUBLIKA S.R.O.	GR
Hexatab Headcold Relief 200 mg/30 mg, επικαλυμμένα με λεπτό υμένιο δισκία	DE/H/4183/001/DC	70000/23-09-2016	JOHNSON & JOHNSON HELLAS CONSUMER AE	GR
Hexatab Headcold Relief 200 mg/30 mg, επικαλυμμένα με λεπτό υμένιο δισκία	DE/H/4183/001/DC	70000/23-09-2016	JOHNSON & JOHNSON HELLAS CONSUMER AE	GR
Hexatab Headcold Relief 200 mg/30 mg, επικαλυμμένα με λεπτό υμένιο δισκία	DE/H/4183/001/DC	70000/23-09-2016	JOHNSON & JOHNSON HELLAS CONSUMER AE	GR
Hexatab Headcold Relief 200 mg/30 mg, επικαλυμμένα με λεπτό υμένιο δισκία	DE/H/4183/001/DC	70000/23-09-2016	JOHNSON & JOHNSON HELLAS CONSUMER AE	GR
Mucocold 200mg/30mg επικαλυμμένα με λεπτό υμένιο δισκία	UK/H/4352/001	91698	BOEHRINGER INGELHEIM INTERNATIONAL GMBH	GR
NUROFEN Cold & Flu	not available	44616/10	RECKITT BENCKISER HELLAS CHEMICALS ABEE	GR
Fidiprofen flu 200 mg+ 30 mg tablete	not available	UP/I-530-09/11-01/363	FIDIFARM D.O.O.	HR

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
Ibuprofen 200 mg/30 mg film-coated tablets	DE/H/4184/001	HR-H-099217339	SANDOZ D.O.O.	HR
Nurofen Cold and Flu 200 mg + 30 mg film-coated tablets	not available	UP/I-530-09/11-02/20	RECKITT BENCKISER (CROATIA) D.O.O.	HR
Olymbion 200 mg/30 mg film-coated tablets	DE/H/4183/001/DC	HR-H-962261553	JOHNSON & JOHNSON S.E.D.O.O.	HR
Olymbion 200 mg/30 mg film-coated tablets	DE/H/4183/001/DC	HR-H-962261553	JOHNSON & JOHNSON S.E.D.O.O.	HR
Olymbion 200 mg/30 mg film-coated tablets	DE/H/4183/001/DC	HR-H-962261553	JOHNSON & JOHNSON S.E.D.O.O.	HR
Olymbion 200 mg/30 mg film-coated tablets	DE/H/4183/001/DC	HR-H-962261553	JOHNSON & JOHNSON S.E.D.O.O.	HR
RHINORELIEF 200 mg + 30 mg tablets	not available	UP/I-530-09/11-01/379	JADRAN-GALENSKI LABORATORIJ D.D.	HR
Advil Cold 200 mg/30 mg bevonat tabletta	IE/H/0420/001	OGYI-T-22705/02	PFIZER CORPORATION AUSTRIA GESELLSCHAFT M.B.H.	HU
Advil Cold 200 mg/30 mg bevonat tabletta	IE/H/0420/001	OGYI-T-22705/01	PFIZER CORPORATION AUSTRIA GESELLSCHAFT M.B.H.	HU
Advil Cold Rapid 200 mg/30 mg lágy kapszula	UK/H/5670/001	OGYI-T-22705/05	PFIZER CORPORATION AUSTRIA GESELLSCHAFT M.B.H.	HU
Advil Cold Rapid 200 mg/30 mg lágy kapszula	UK/H/5670/001	OGYI-T-22705/03	PFIZER CORPORATION AUSTRIA GESELLSCHAFT M.B.H.	HU
Advil Cold Rapid 200 mg/30 mg lágy kapszula	UK/H/5670/001	OGYI-T-22705/06	PFIZER CORPORATION AUSTRIA GESELLSCHAFT M.B.H.	HU
Advil Cold Rapid 200 mg/30 mg lágy kapszula	UK/H/5670/001	OGYI-T-22705/04	PFIZER CORPORATION AUSTRIA GESELLSCHAFT M.B.H.	HU
ALGOFLEX COLD 200 mg/30 mg filmtabletta	not available	OGYI-T-22831/01	SANOFI-AVENTIS ZRT	HU

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ALGOFLEX COLD 200 mg/30 mg filmtabletta	not available	OGYI-T-22831/03	SANOFI-AVENTIS ZRT	HU
ALGOFLEX COLD 200 mg/30 mg filmtabletta	not available	OGYI-T-22831/02	SANOFI-AVENTIS ZRT	HU
Ibuprofen/Pseudoefedrin Wick 200 mg/30 mg filmtabletta	IT/H0331/001	OGYI-T-22836/01	WICK PHARMA ZWEIGNIEDERLASSUNG DER PROCTER & GAMBLE GMBH	HU
Ibuprofen/Pseudoefedrin Wick 200 mg/30 mg filmtabletta	IT/H/0331/001	OGYI-T-22836/02	WICK PHARMA ZWEIGNIEDERLASSUNG DER PROCTER & GAMBLE GMBH	HU
Ibuprofen/Pseudoefedrin Wick 200 mg/30 mg filmtabletta	IT/H/0331/001	OGYI-T-22836/03	WICK PHARMA ZWEIGNIEDERLASSUNG DER PROCTER & GAMBLE GMBH	HU
Nurofen Cold and Flu 200 mg/30 mg filmtabletta	not available	OGYI-T-6797/02	RECKITT BENCKISER HEALTHCARE INTERNATIONAL LIMITED	HU
Nurofen Cold and Flu 200 mg/30 mg filmtabletta	not available	OGYI-T-6797/01	RECKITT BENCKISER HEALTHCARE INTERNATIONAL LIMITED	HU
Advil Cold & Flu Coated Tablets Ibuprofen 200 mg Pseudoephedrine Hydrochloride 30 mg	IE/H/0420/001	PA 0822/164/001	PFIZER HEALTHCARE IRELAND	IE
Advil Sinus Relief Soft Capsules Ibuprofen 200 mg Pseudoephedrine hydrochloride 30 mg	UK/H/5670/001	PA 822/164/002	PFIZER HEALTHCARE IRELAND	IE
Ibuprofen/Pseudoephedrine Hydrochloride 200mg/30mg film coated tablets	DE/H/4182/001/DC	PA823/067/001	MCNEIL HEALTHCARE (IRELAND) LIMITED	IE
Ibuprofen/Pseudoephedrine Hydrochloride 200mg/30mg film coated tablets	DE/H/4182/001/DC	PA823/067/001	MCNEIL HEALTHCARE (IRELAND) LIMITED	IE

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Ibuprofen/Pseudoephedrine Hydrochloride 200mg/30mg film coated tablets	DE/H/4182/001/DC	PA823/067/001	MCNEIL HEALTHCARE (IRELAND) LIMITED	IE
Ibuprofen/Pseudoephedrine Hydrochloride 200mg/30mg film coated tablets	DE/H/4182/001/DC	PA823/067/001	MCNEIL HEALTHCARE (IRELAND) LIMITED	IE
Non-Drowsy Sudapro Head Cold 200mg / 30mg Film-coated Tablets	DE/H/4183/001/DC	PA 823/68/1	MCNEIL HEALTHCARE (IRELAND) LIMITED	IE
Non-Drowsy Sudapro Head Cold 200mg / 30mg Film-coated Tablets	DE/H/4183/001/DC	PA 823/68/1	MCNEIL HEALTHCARE (IRELAND) LIMITED	IE
Non-Drowsy Sudapro Head Cold 200mg / 30mg Film-coated Tablets	DE/H/4183/001/DC	PA 823/68/1	MCNEIL HEALTHCARE (IRELAND) LIMITED	IE
Non-Drowsy Sudapro Head Cold 200mg / 30mg Film-coated Tablets	DE/H/4183/001/DC	PA 823/68/1	MCNEIL HEALTHCARE (IRELAND) LIMITED	IE
Nurofen Cold & Flu Film-coated Tablets, Ibuprofen 200mg, Pseduoephedrine Hydrochloride 30mg	not available	PA 979/33/1	RECKITT BENCKISER IRELAND LTD.	IE
Nurofen Sinus and Pain Film-Coated Tablets Ibuprofen 200mg Pseudoephedrine Hydrochloride 30mg	not available	PA 979/65/1	RECKITT BENCKISER IRELAND LTD.	IE
Nurofen Sinus and Pain Film-Coated Tablets Ibuprofen 200mg Pseudoephedrine Hydrochloride 30mg	not available	PA 979/65/1	RECKITT BENCKISER IRELAND LTD.	IE
ACTISINU 200 mg/30 mg, compresse rivestite con film	DE/H/4183/001/DC	043681016	JOHNSON & JOHNSON S.P.A.	IT
ACTISINU 200 mg/30 mg, compresse rivestite con film	DE/H/4183/001/DC	043681030	JOHNSON & JOHNSON S.P.A.	IT

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ACTISINU 200 mg/30 mg, compresse rivestite con film	DE/H/4183/001/DC	043681028	JOHNSON & JOHNSON S.P.A.	IT
ACTISINU 200 mg/30 mg, compresse rivestite con film	DE/H/4183/001/DC	043681042	JOHNSON & JOHNSON S.P.A.	IT
Momenxsin 200 mg/30 mg compresse rivestite con film	DE/H/4186/001	043682018	AZIENDE CHIMICHE RIUNITE ANGELINI FRANCESCO - A.C.R.A.F. S.P.A.	IT
Momenxsin 200 mg/30 mg compresse rivestite con film	DE/H/4186/001	043682020	AZIENDE CHIMICHE RIUNITE ANGELINI FRANCESCO - A.C.R.A.F. S.P.A.	IT
Momenxsin 200 mg/30 mg compresse rivestite con film	DE/H/4186/001	043682032	AZIENDE CHIMICHE RIUNITE ANGELINI FRANCESCO - A.C.R.A.F. S.P.A.	IT
Momenxsin 200 mg/30 mg compresse rivestite con film	DE/H/4186/001	043682044	AZIENDE CHIMICHE RIUNITE ANGELINI FRANCESCO - A.C.R.A.F. S.P.A.	IT
NUROFEN INFLUENZA E RAFFREDDORE 200 mg + 30 mg Compresse Rivestite	not available	034246013	RECKITT BENCKISER HEALTHCARE INTERNATIONAL LIMITED	IT
NUROFEN INFLUENZA E RAFFREDDORE 200 mg + 30 mg Compresse Rivestite	not available	034246025	RECKITT BENCKISER HEALTHCARE INTERNATIONAL LIMITED	IT
Vicks Flu Action 200 mg/30 mg compresse rivestite con film	IT/H/0331/001	042499032	PROCTER & GAMBLE S.R.L	IT
Vicks Flu Action 200 mg/30 mg compresse rivestite con film	IT/H/0331/001	042499044	PROCTER & GAMBLE S.R.L	IT
Vicks Flu Action 200 mg/30 mg compresse rivestite con film	IT/H/0331/001	042499057	PROCTER & GAMBLE S.R.L	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
ZerinoDek 200 mg/30 mg compresse rivestite con film	UK/H/4352/001	041218013	SANOFI SPA	IT
ZerinoDek 200 mg/30 mg compresse rivestite con film	UK/H/4352/001	041218025	SANOFI SPA	IT
Daikol 200 mg/30 mg plėvele dengtos tabletės	not available	LT/1/16/3982/001	SIA INGEN PHARMA	LT
ILOXEN 200 mg/30 mg plėvele dengtos tabletės	DE/H/4185/001	LT/1/16/3881/001	PHARMASWISS ČESKÁ REPUBLIKA S.R.O.	LT
ILOXEN 200 mg/30 mg plėvele dengtos tabletės	DE/H/4185/001	LT/1/16/3881/002	PHARMASWISS ČESKÁ REPUBLIKA S.R.O.	LT
ILOXEN 200 mg/30 mg plėvele dengtos tabletės	DE/H/4185/001	LT/1/16/3881/003	PHARMASWISS ČESKÁ REPUBLIKA S.R.O.	LT
ILOXEN 200 mg/30 mg plėvele dengtos tabletės	DE/H/4185/001	LT/1/16/3881/004	PHARMASWISS ČESKÁ REPUBLIKA S.R.O.	LT
Ibuprofen/Pseudoephedrine HCl Pfizer 200 mg / 30 mg Comprimés enrobé	IE/H/0420/001	2015040073	PFIZER B.V.	LU
IBUPROFEN/PSEUDOEPHEDRINE HYDROCHLORIDE SANOFI 200 mg/30 mg comprimés pelliculés	UK/H/4352/001	2014040030	SANOFI BELGIUM	LU
Ibuprofen/Pseudoephedrine hydrochloride Sanofi 200 mg/30 mg Filmtabletten	UK/H/4352/001	2014040030	SANOFI BELGIUM	LU
IbuSinex 200 mg/30 mg comprimé pelliculé	DE/H/4187/001	2016110303	DIAPHARM GMBH & CO. KG	LU
Sinuphen 200 mg/30 mg comprimés pelliculés	DE/H/4183/001/DC	2016110302	JOHNSON & JOHNSON CONSUMER N.V./S.A.	LU
Sinuphen 200 mg/30 mg comprimés pelliculés	DE/H/4183/001/DC	2016110302	JOHNSON & JOHNSON CONSUMER N.V./S.A.	LU
Sinuphen 200 mg/30 mg comprimés pelliculés	DE/H/4183/001/DC	2016110302	JOHNSON & JOHNSON CONSUMER N.V./S.A.	LU
Sinuphen 200 mg/30 mg comprimés pelliculés	DE/H/4183/001/DC	2016110302	JOHNSON & JOHNSON CONSUMER N.V./S.A.	LU
ILOXEN 200 mg/30 mg apvalkotās tabletes	DE/H/4185/001	16-0009	PHARMASWISS ČESKÁ REPUBLIKA S.R.O.	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
Nurofen Antigrip 200 mg/30 mg apvalkotās tabletes	not available	04-0197	RECKITT BENCKISER (POLAND) S.A.	LV
Advil Cold & Flu Coated Tablets	not available	MA 969/00201	PFIZER CONSUMER HEALTHCARE LTD.	MT
Advil Cold & Flu Coated Tablets	not available	MA 969/00201	PFIZER CONSUMER HEALTHCARE LTD.	MT
Advil Sinus Relief Soft Capsules Ibuprofen 200 mg Pseudoephedrine hydrochloride 30 mg	UK/H/5670/001	MA969/00202	PFIZER CORPORATION AUSTRIA GESELLSCHAFT M.B.H.	MT
Nurofen Cold & Flu	not available	MA 190/00404	RECKITT BENCKISER HEALTHCARE INTERNATIONAL LIMITED	MT
Acatar Zatoki, 200 mg + 30 mg, tabletki powlekane	not available	11339	US PHARMACIA SP. Z O.O.	PL
Advil Zatoki, 200 mg + 30 mg, tabletki powlekane	IE/H/0420/001	22369	PFIZER CORPORATION AUSTRIA GESELLSCHAFT M.B.H.	PL
Efedoxin, 200 mg + 30 mg, tabletki powlekane	DE/H/4185/001	23302	PHARMASWISS ČESKÁ REPUBLIKA S.R.O.	PL
IBUM GRIP, 200 mg + 30 mg, tabletki powlekane	not available	10924	HASCO-LEK	PL
Ibum Zatoki Max, 400 mg + 60 mg, tabletki powlekane	not available	23760	HASCO-LEK	PL
Ibum Zatoki, 200 mg + 30 mg, tabletki powlekane	not available	21376	HASCO-LEK	PL
Ibuprofen/Pseudoephedrine hydrochloride Krewel, 200 mg + 30 mg, tabletki powlekane	DE/H/4188/001	23399	KREWEL MEUSELBACH GMBH	PL
IBUPROM ZATOKI, 200 mg + 30 mg, tabletki powlekane	not available	9650	US PHARMACIA SP. Z O.O.	PL
Infex Zatoki	IT-H-0331-001-DC	22111	TEVA PHARMACEUTICALS POLSKA SP. Z O.O.	PL

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
Laboratoria PolfaŁódź ZATOKI, 200 mg + 30 mg, tabletki	not available	21816	BIO PROFIL POLSKA SP Z OO	PL
Metafen ZATOKI, 200 mg + 30 mg, tabletki	not available	11005	ZAKŁADY FARMACEUTYCZNE "POLPHARMA" SPOLKA AKCYJNA	PL
Metafen ZATOKI, 200 mg + 30 mg, tabletki	not available	11005	ZAKŁADY FARMACEUTYCZNE "POLPHARMA" SPOLKA AKCYJNA	PL
MODAFEN EXTRA GRIP, 200 mg + 30 mg, tabletki powlekane	not available	9603	ZENTIVA, K.S.	PL
MODAFEN EXTRA GRIP, 200 mg + 30 mg, tabletki powlekane	not available	9603	ZENTIVA, K.S.	PL
MODAFEN EXTRA GRIP, 200 mg + 30 mg, tabletki powlekane	not available	9603	ZENTIVA, K.S.	PL
Nurofen Zatoki, 200 mg + 30 mg, tabletki powlekane	not available	7787	RECKITT BENCKISER (POLAND) S.A.	PL
Sudafed Extra, 200 mg + 30 mg, tabletki powlekane	DE/H/4183/001/DC	23235	MCNEIL PRODUCTS LIMITED	PL
Sudafed Extra, 200 mg + 30 mg, tabletki powlekane	DE/H/4183/001/DC	23235	MCNEIL PRODUCTS LIMITED	PL
Sudafed Extra, 200 mg + 30 mg, tabletki powlekane	DE/H/4183/001/DC	23235	MCNEIL PRODUCTS LIMITED	PL
Sudafed Extra, 200 mg + 30 mg, tabletki powlekane	DE/H/4183/001/DC	23235	MCNEIL PRODUCTS LIMITED	PL
Bisolpront 200 mg + 30 mg comprimidos revestidos por película	UK/H/4352/001	5666110	SANOFI - PRODUTOS FARMACEUTICOS LDA	PT
Bisolpront 200 mg + 30 mg comprimidos revestidos por película	UK/H/4352/001	5666102	SANOFI - PRODUTOS FARMACEUTICOS LDA	PT

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
Larofen Plus 200mg/30mg comprimate filmate	not available	9424/2016/01	LAROPHARM SRL	RO
Modafen 200 mg/30 mg comprimate filmate	not available	480/2007/02	ZENTIVA, K.S.	RO
Modafen 200 mg/30 mg comprimate filmate	not available	480/2007/03	ZENTIVA, K.S.	RO
Modafen 200 mg/30 mg comprimate filmate	not available	480/2007/04	ZENTIVA, K.S.	RO
Modafen 200 mg/30 mg comprimate filmate	not available	480/2007/01	ZENTIVA, K.S.	RO
Nurofen Răceală și Gripă 200 mg/30 mg comprimate filmate	not available	4144/2011/02	RECKITT BENCKISER HEALTHCARE INTERNATIONAL LIMITED	RO
Nurofen Răceală și Gripă 200 mg/30 mg comprimate filmate	not available	4144/2011/01	RECKITT BENCKISER HEALTHCARE INTERNATIONAL LIMITED	RO
Olytabs 200 mg/30 mg comprimate filmate	DE/H/4183/001/DC	8529/2016/03	MCNEIL PRODUCTS LIMITED	RO
Olytabs 200 mg/30 mg comprimate filmate	DE/H/4183/001/DC	8529/2016/01	MCNEIL PRODUCTS LIMITED	RO
Olytabs 200 mg/30 mg comprimate filmate	DE/H/4183/001/DC	8529/2016/04	MCNEIL PRODUCTS LIMITED	RO
Olytabs 200 mg/30 mg comprimate filmate	DE/H/4183/001/DC	8529/2016/02	MCNEIL PRODUCTS LIMITED	RO
Robicold Sinus & Răceală 200 mg/30 mg capsule moi	UK/H/5670/001	8760/2016/01	PFIZER CORPORATION AUSTRIA GESELLSCHAFT M.B.H.	RO
Robicold Sinus & Răceală 200 mg/30 mg capsule moi	UK/H/5670/001	8760/2016/02	PFIZER CORPORATION AUSTRIA GESELLSCHAFT M.B.H.	RO
Robicold Sinus & Răceală 200 mg/30 mg capsule moi	UK/H/5670/001	8760/2016/08	PFIZER CORPORATION AUSTRIA GESELLSCHAFT M.B.H.	RO
Robicold Sinus & Răceală 200 mg/30 mg capsule moi	UK/H/5670/001	8760/2016/14	PFIZER CORPORATION AUSTRIA GESELLSCHAFT M.B.H.	RO

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
Robicold Sinus & Răceală 200 mg/30 mg capsule moi	UK/H/5670/001	8760/2016/12	PFIZER CORPORATION AUSTRIA GESELLSCHAFT M.B.H.	RO
Robicold Sinus & Răceală 200 mg/30 mg capsule moi	UK/H/5670/001	8760/2016/03	PFIZER CORPORATION AUSTRIA GESELLSCHAFT M.B.H.	RO
Robicold Sinus & Răceală 200 mg/30 mg capsule moi	UK/H/5670/001	8760/2016/10	PFIZER CORPORATION AUSTRIA GESELLSCHAFT M.B.H.	RO
Robicold Sinus & Răceală 200 mg/30 mg capsule moi	UK/H/5670/001	8760/2016/06	PFIZER CORPORATION AUSTRIA GESELLSCHAFT M.B.H.	RO
Robicold Sinus & Răceală 200 mg/30 mg capsule moi	UK/H/5670/001	8760/2016/04	PFIZER CORPORATION AUSTRIA GESELLSCHAFT M.B.H.	RO
Robicold Sinus & Răceală 200 mg/30 mg capsule moi	UK/H/5670/001	8760/2016/16	PFIZER CORPORATION AUSTRIA GESELLSCHAFT M.B.H.	RO
Robicold Sinus & Răceală 200 mg/30 mg capsule moi	UK/H/5670/001	8760/2016/05	PFIZER CORPORATION AUSTRIA GESELLSCHAFT M.B.H.	RO
Robicold Sinus & Răceală 200 mg/30 mg capsule moi	UK/H/5670/001	8760/2016/13	PFIZER CORPORATION AUSTRIA GESELLSCHAFT M.B.H.	RO
Robicold Sinus & Răceală 200 mg/30 mg capsule moi	UK/H/5670/001	8760/2016/07	PFIZER CORPORATION AUSTRIA GESELLSCHAFT M.B.H.	RO
Robicold Sinus & Răceală 200 mg/30 mg capsule moi	UK/H/5670/001	8760/2016/11	PFIZER CORPORATION AUSTRIA GESELLSCHAFT M.B.H.	RO
Robicold Sinus & Răceală 200 mg/30 mg capsule moi	UK/H/5670/001	8760/2016/09	PFIZER CORPORATION AUSTRIA GESELLSCHAFT M.B.H.	RO
Robicold Sinus & Răceală 200 mg/30 mg capsule moi	UK/H/5670/001	8760/2016/15	PFIZER CORPORATION AUSTRIA GESELLSCHAFT M.B.H.	RO

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
TEDOLFEN 200mg/30mg comprimate filmate	IT/H/0331/001	6744/2014/01-04	TEVA PHARMACEUTICALS S.R.L	RO
Ibuseu, 200 mg/30 mg filmdragerade tabletter	AT/H/0614/001	52535	ORIFARM GENERICS A/S	SE
Olytabs 200 mg/30 mg filmsko obložene tablete	DE/H/4183/001/DC	H/16/02163/004	MCNEIL PRODUCTS LIMITED	SI
Olytabs 200 mg/30 mg filmsko obložene tablete	DE/H/4183/001/DC	H/16/02163/002	MCNEIL PRODUCTS LIMITED	SI
Olytabs 200 mg/30 mg filmsko obložene tablete	DE/H/4183/001/DC	H/16/02163/001	MCNEIL PRODUCTS LIMITED	SI
Olytabs 200 mg/30 mg filmsko obložene tablete	DE/H/4183/001/DC	H/16/02163/003	MCNEIL PRODUCTS LIMITED	SI
Grippecton 200 mg/30 mg filmom obalené tablety	DE/H/4188/001	07/0143/16-S	KREWEL MEUSELBACH GMBH	SK
MODAFEN 200 mg/30 mg filmom obalené tablety	not available	07/0263/01-S	ZENTIVA, K.S.	SK
MODAFEN 200 mg/30 mg filmom obalené tablety	not available	07/0263/01-S	ZENTIVA, K.S.	SK
MODAFEN 200 mg/30 mg filmom obalené tablety	not available	07/0263/01-S	ZENTIVA, K.S.	SK
NUROFEN STOPGRIP	not available	07/0662/96-S	RECKITT BENCKISER HEALTHCARE INTERNATIONAL LIMITED	SK
Robicold 200 mg/30 mg obalené tablety	IE/H/0420/001	07/0264/14-S	PFIZER CORPORATION AUSTRIA GESELLSCHAFT M.B.H.	SK
Robicold Rapid 200 mg/30 mg mäkké kapsuly	UK/H/5670/001	07/0235/16-S	PFIZER CORPORATION AUSTRIA GESELLSCHAFT M.B.H.	SK
Boots Cold & Flu Relief with Ibuprofen	not available	00014/0600	THE BOOTS COMPANY PLC	UK
Ibuprofen/ Pseudoephedrine hydrochloride Pfizer 200mg/30mg Capsules	UK/H/5671/001	PL 00165/0389	PFIZER CONSUMER HEALTHCARE LTD.	UK

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
Nurofen Sinus Pressure & Headache Relief 200mg/30mg Tablets	not available	PL 00063/0718	RECKITT BENCKISER HEALTHCARE (UK) LIMITED	UK
Nurofen Cold & Flu	not available	PL 00063/0375	RECKITT BENCKISER HEALTHCARE (UK) LIMITED	UK
Nurofen Sinus Relief	not available	PL 00063/0375	RECKITT BENCKISER HEALTHCARE (UK) LIMITED	UK
RobiCold Cold & Flu Relief 200 mg / 30 mg Soft Capsules	UK/H/5670/001	PL 00165/0388	PFIZER CONSUMER HEALTHCARE LTD.	UK
RobiCold Sinus Relief 200 mg, 30 mg Tablets	IE/H/0420/001	PL 00165/0391	PFIZER CONSUMER HEALTHCARE LTD.	UK
Sudafed Sinus Pressure & Pain 200mg/30mg film-coated tablets	DE/H/4183/001	PL 15513/0396	MCNEIL PRODUCTS LIMITED	UK
Sudafed Sinus Pressure & Pain 200mg/30mg film-coated tablets	DE/H/4183/001	PL 15513/0396	MCNEIL PRODUCTS LIMITED	UK
Sudafed Sinus Pressure & Pain 200mg/30mg film-coated tablets	DE/H/4183/001	PL 15513/0396	MCNEIL PRODUCTS LIMITED	UK
Sudafed Sinus Pressure & Pain 200mg/30mg film-coated tablets	DE/H/4183/001	PL 15513/0396	MCNEIL PRODUCTS LIMITED	UK
Sudafed Sinus Pressure & Pain Tablets	not available	PL 15513/0126	MCNEIL PRODUCTS LIMITED	UK
Sudafed Sinus Pressure & Pain Tablets	not available	PL 15513/0126	MCNEIL PRODUCTS LIMITED	UK
Sudafed Sinus Pressure & Pain Tablets	not available	PL 15513/0126	MCNEIL PRODUCTS LIMITED	UK
Sudafed Sinus Pressure & Pain Tablets	not available	PL 15513/0126	MCNEIL PRODUCTS LIMITED	UK