



EUROPEAN MEDICINES AGENCY
SCIENCE MEDICINES HEALTH

8 March 2018
EMA/182668/2018
Human Medicines Evaluation Division

List of nationally authorised medicinal products

Active substance(s): triamcinolone (topical and nasal formulations)

Procedure No.: PSUSA/00003017/201707



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
AFTAB 0,025 mg Hafttabletten	not available	29856.00.00	MEDA PHARMA GMBH & CO. KG	DE
AFTAB 25 microgrammi compresse buccali mucoadesive	not available	028478	ROTTAPHARM S.P.A.	IT
Aftab 25 mikrog bukkaalitabletti	not available	12764	MEDA OY	FI
Aftab 25 mikrogram buckaltablett	not available	12764	MEDA OY	FI
Aftach	not available	8771816	ANGELINI FARMACÊUTICA, LDA	PT
Aftach	not available	8771808	ANGELINI FARMACÊUTICA, LDA	PT
Allegra nasal 55 microgrammes/dose suspension pour pulvérisation nasale	UK/H/4484/001	BE400671	SANOFI BELGIUM	BE
Allegra nasal 55 microgrammes/dose suspension pour pulvérisation nasale	UK/H/4484/001	BE400662	SANOFI BELGIUM	BE
Allegra nasal 55 microgrammes/dose suspension pour pulvérisation nasale	UK/H/4484/001	0641166	SANOFI BELGIUM	LU
Allegra nasal 55 microgrammes/dose suspension pour pulvérisation nasale	UK/H/4484/001	0641152	SANOFI BELGIUM	LU
Allegra nasal, 55 microgram/dosis neusspray, suspensie	UK/H/4484/001	BE400671	SANOFI BELGIUM	BE
Allegra nasal, 55 microgram/dosis neusspray, suspensie	UK/H/4484/001	BE400662	SANOFI BELGIUM	BE

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FTOROCORT 1 mg/g ziede	not available	96-0589	GEDEON RICHTER PLC.	LV
FTOROCORT 0,1 % tepalas	NULL	LT/1/94/0978/001	GEDEON RICHTER PLC.	LT
FTOROCORT, 1 mg/g salv	not available	093494	GEDEON RICHTER PLC.	EE
NASACORT	not available	19193	SANOFI-AVENTIS CYPRUS LTD	CY
NASACORT	not available	19193	SANOFI-AVENTIS CYPRUS LTD	CY
NASACORT	UK/H/0189/001	19048	SANOFI-AVENTIS DENMARK A/S	DK
NASACORT	UK/H/0189/001	19048	SANOFI-AVENTIS DENMARK A/S	DK
NASACORT	UK/H/0189/001	237440102	SANOFI-AVENTIS AEBE	GR
NASACORT	UK/H/0189/001	237440101	SANOFI-AVENTIS AEBE	GR
NASACORT	not available	MA082/01701	SANOFI MALTA LTD	MT
NASACORT	not available	MA082/01701	SANOFI MALTA LTD	MT
NASACORT	UK/H/0189/001	2589489	SANOFI - PRODUTOS FARMACEUTICOS LDA	PT
NASACORT 55 microgram/dosis, neusspray, suspensie	UK/H/0189/001	BE190215	SANOFI BELGIUM	BE
NASACORT 55 microgram/dosis, neusspray, suspensie	UK/H/0189/001	BE190215	SANOFI BELGIUM	BE
Nasacort 55 micrograme/doză spray nazal suspensie	UK/H/0189/001	7952/2015/01	SANOFI-AVENTIS ROMANIA SRL	RO
Nasacort 55 micrograme/doză spray nazal suspensie	UK/H/0189/001	7952/2015/02	SANOFI-AVENTIS ROMANIA SRL	RO
NASACORT 55 MICROGRAMMES/DOSE, SUSPENSION POUR	not available	342 922-9	SANOFI-AVENTIS FRANCE	FR

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PULVERISATION NASALE				
NASACORT 55 microgrammes/dose, suspension pour pulvérisation nasale	UK/H/0189/001	BE190215	SANOFI BELGIUM	BE
NASACORT 55 microgrammes/dose, suspension pour pulvérisation nasale	UK/H/0189/001	BE190215	SANOFI BELGIUM	BE
Nasacort 55 microgrammes/dose, suspension pour pulvérisation nasale	UK/H/0189/001	0247171	SANOFI BELGIUM	LU
NASACORT 55 microgrammi/dose, spray nasale, sospensione	UK/H/0189/001	033938010	SANOFI SPA	IT
NASACORT 55 microgrammi/dose, spray nasale, sospensione	UK/H/0189/001	033938022	SANOFI SPA	IT
NASACORT 55 microgramos/dosis, suspensión para pulverización nasal	UK/H/0189/001	61970	SANOFI-AVENTIS, S.A.	ES
NASACORT 55 microgramos/dosis, suspensión para pulverización nasal	UK/H/0189/001	61970	SANOFI-AVENTIS, S.A.	ES
NASACORT 55 MICROGRAMS/DOSE, NASAL SPRAY, SUSPENSION	not available	960219	SANOFI-AVENTIS NORGE AS	IS
NASACORT 55 MICROGRAMS/DOSE,	not available	960219	SANOFI-AVENTIS NORGE AS	IS

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NASAL SPRAY, SUSPENSION				
NASACORT 55 micrograms/dose, nasal spray, suspension	UK/H/0189/001	PL 04425/0287	AVENTIS PHARMA LTD	UK
NASACORT 55 micrograms/dose, nasal spray, suspension	UK/H/0189/001	PL 04425/0287	AVENTIS PHARMA LTD	UK
NASACORT 55 MIKROGRAM/DOS, NASSPRAY, SUSPENSION	UK/H/0189/001	13781	SANOFI AB	SE
NASACORT 55 MIKROGRAM/DOS, NASSPRAY, SUSPENSION	UK/H/0189/001	13781	SANOFI AB	SE
NASACORT 55 mikrogrami/devā deguna aerosols, suspensija	UK/H/0189/001	15-0135	SANOFI-AVENTIS LATVIA SIA	LV
NASACORT 55 mikrogrami/devā deguna aerosols, suspensija	UK/H/0189/001	15-0135	SANOFI-AVENTIS LATVIA SIA	LV
Nasacort 55 mikrogramm/adag szuszpenziós orrspray	UK/H/0189/001	OGYI-T-22852/02	SANOFI-AVENTIS ZRT	HU
Nasacort 55 mikrogramm/adag szuszpenziós orrspray	UK/H/0189/001	OGYI-T-22852/01	SANOFI-AVENTIS ZRT	HU
NASACORT 55 MIKROGRAMM/DOSIS	UK/H/0189/001	20150255	SANOFI BULGARIA EOOD	BG
NASACORT 55 MIKROGRAMM/DOSIS	UK/H/0189/001	20150255	SANOFI BULGARIA EOOD	BG
NASACORT 55 MIKROGRAMM/DOSIS	UK/H/0189/001	41503.00.00	SANOFI-AVENTIS DEUTSCHLAND GMBH	DE
NASACORT 55 MIKROGRAMM/DOSIS	UK/H/0189/001	41503.00.00	SANOFI-AVENTIS DEUTSCHLAND GMBH	DE
Nasacort 55	UK/H/0189/001	12965	SANOFI OY	FI

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mikrogrammaa/annos nenäsumute, suspensio				
Nasacort 55 mikrogrammaa/annos nenäsumute, suspensio	UK/H/0189/001	12965	SANOFI OY	FI
NASACORT 55 mikrogramov /odmerek pršilo za nos	UK/H/0189/001	H/16/02186/001	SANOFI-AVENTIS D.O.O.	SI
NASACORT 55 mikrogramov /odmerek pršilo za nos	UK/H/0189/001	H/16/02186/002	SANOFI-AVENTIS D.O.O.	SI
Nasacort 55 mikrogramov/dávka	UK/H/0189/001	69/0207/15-S	SANOFI-AVENTIS SLOVAKIA SRO	SK
Nasacort 55 mikrogramov/dávka	UK/H/0189/001	69/0207/15-S	SANOFI-AVENTIS SLOVAKIA SRO	SK
NASACORT 55 micrograms/dose, nasal spray, suspension	UK/H/0189/001	PA 540/11/1	SANOFI-AVENTIS IRELAND LTD. T/A SANOFI	IE
NASACORT 55 micrograms/dose, nasal spray, suspension	UK/H/0189/001	PA 540/11/1	SANOFI-AVENTIS IRELAND LTD. T/A SANOFI	IE
NASACORT ALLERGY	UK/H/4484/001	23710/22-3-2013	SANOFI-AVENTIS AEBE	GR
NASACORT ALLERGY	UK/H/4484/001	23710/22-3-2013	SANOFI-AVENTIS AEBE	GR
NASACORT ALLERGY	UK/H/4484/001	PA 540/11/2	SANOFI-AVENTIS IRELAND LTD. T/A SANOFI	IE
NASACORT ALLERGY	UK/H/4484/001	PA 540/11/2	SANOFI-AVENTIS IRELAND LTD. T/A SANOFI	IE
NASACORT ALLERGY	not available	PL 04425/0605	AVENTIS PHARMA LTD	UK
Nasacort nes spray, suspensjon	not available	95-3932	SANOFI-AVENTIS NORGE AS	NO
Nasacort nes spray, suspensjon	not available	95-3932	SANOFI-AVENTIS NORGE AS	NO
NASACORT SANOFI-AVENTIS 55 MIKROGRAMM/SPRUHST	UK/H/4484/001	82818.00.00	SANOFI-AVENTIS DEUTSCHLAND GMBH	DE

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OSS NASENSPRAY, SUSPENSION				
NASACORT SANOFI-AVENTIS 55 MIKROGRAMM/SPRUHST OSS NASENSPRAY, SUSPENSION	UK/H/4484/001	82818.00.00	SANOFI-AVENTIS DEUTSCHLAND GMBH	DE
Nasacort, 55 microgram/verstuiving neusspray, suspensie	UK/H/0189/001	RGV 21837	SANOFI-AVENTIS NETHERLANDS B.V.	NL
Nasacort, 55 mikrogrammi/pihustuses , ninasprei, suspensioon	UK/H/0189/001	875715	SANOFI-AVENTIS ESTONIA OÜ	EE
Nasacort, 55 mikrogrammi/pihustuses , ninasprei, suspensioon	UK/H/0189/001	875715	SANOFI-AVENTIS ESTONIA OÜ	EE
Nasacort, 55 mikrogramów/dawkę, aerozol do nosa, zawiesina	UK/H/0189/001	22738	AVENTIS PHARMA LTD	PL
Nasacort, 55 mikrogramów/dawkę, aerozol do nosa, zawiesina	UK/H/0189/001	22738	AVENTIS PHARMA LTD	PL
RHINISAN 55 MIKROGRAMM/DOSIS	not available	41504.00.00	SANOFI-AVENTIS DEUTSCHLAND GMBH	DE
RHINISAN 55 MIKROGRAMM/DOSIS	not available	41504.00.00	SANOFI-AVENTIS DEUTSCHLAND GMBH	DE
Triamcinolon Teva 1 mg/g kožní emulze	not available	46/219/74-C	TEVA CZECH INDUSTRIES S.R.O	CZ
Triamcinolon/acetonide zalf	not available	RVG 52672	BASIC PHARMA MANUFACTURING BV	NL
Triamcinolona Sanofi 55 microgramas/dose, suspensão para	UK/H/4484/001	5404579	SANOFI - PRODUTOS FARMACEUTICOS LDA	PT

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pulverização nasal				
Triamcinolona Sanofi 55 microgramas/dose, suspensão para pulverização nasal	UK/H/4484/001	5404603	SANOFI - PRODUTOS FARMACEUTICOS LDA	PT
Triamcinolonacetamid "sanofi-aventis"	UK/H/4484/001	47352	SANOFI-AVENTIS DENMARK A/S	DK
Triamcinolonacetamid "sanofi-aventis"	UK/H/4484/001	47352	SANOFI-AVENTIS DENMARK A/S	DK
Triamcinolonacetamida hidrofílica creme 1 mg/g	not available	RVG 56785	BASIC PHARMA MANUFACTURING BV	NL
Triamcinolonacetamida hidrofílica creme Basic Pharma 1 mg/g	not available	RVG 19042=51893	APOTEX EUROPE B.V.	NL
Triamcinolonacetamida hidrofílica creme Basic Pharma 1 mg/g	not available	RVG 51893	BASIC PHARMA MANUFACTURING BV	NL
Triamcinolone acetamida 55 microgramas/dose, spray nasal, suspensão	UK/H/4484/001	PL 04425/0669	AVENTIS PHARMA LTD	UK
Triamcinolone acetamida 55 microgramas/dose, spray nasal, suspensão	UK/H/4484/001	PL 04425/0669	AVENTIS PHARMA LTD	UK
Triamcinolone acetamida Sanofi 55 microgramas/dose, spray nasal, suspensão	UK/H/0189/001	LT/1/15/3760/001	UAB SANOFI-AVENTIS LIETUVA	LT
Triamcinolone acetamida Sanofi 55 microgramas/dose, spray nasal, suspensão	UK/H/0189/001	LT/1/15/3760/002	UAB SANOFI-AVENTIS LIETUVA	LT
TRIAMCINOLONE SANOFI 55 MIKGROGRAMŮ/DÁVKA	UK/H/0189/001	69/324/15-C	SANOFI-AVENTIS SRO	CZ
TRIAMCINOLONE	UK/H/0189/001	69/324/15-C	SANOFI-AVENTIS SRO	CZ

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SANOFI 55 MIKGROGRAMŮ/DÁVKA				