



EUROPEAN MEDICINES AGENCY
SCIENCE MEDICINES HEALTH

11 February 2021
EMA/172058/2021
Human Medicines Evaluation Division

List of nationally authorised medicinal products

Active substance: lidocaine_tetracaine

Procedure no.: PSUSA/00001868/202006



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
PLIAGLIS 70 mg / 70 mg par gramme, crème	DE/H/3278/001	34009 224 112 7 9	GALDERMA INTERNATIONAL	FR
PLIAGLIS 70 mg / 70 mg par gramme, crème	DE/H/3278/001	34009 224 113 3 0	GALDERMA INTERNATIONAL	FR
Pliaglis 70 mg/g + 70 mg/g Crema	DE/H/3278/001	041546019	DIFA COOPER S.P.A.	IT
Pliaglis 70 mg/g + 70 mg/g Crema	DE/H/3278/001	041546021	DIFA COOPER S.P.A.	IT
Pliaglis 70 mg/g + 70 mg/g Creme	DE/H/3278/001	BE434436	GALDERMA BENELUX B.V.	BE
Pliaglis 70 mg/g + 70 mg/g crème	DE/H/3278/001	RVG 109762	GALDERMA BENELUX B.V.	NL
Pliaglis 70 mg/g + 70 mg/g κρέμα	DE/H/3278/001	30937/27.03.2019	GALDERMA INTERNATIONAL	GR
Pliaglis 70 mg/g + 70 mg/g, crème	DE/H/3278/001	2013060200	GALDERMA BENELUX B.V.	LU
Pliaglis 70 mg/g + 70 mg/g, crème	DE/H/3278/001	BE434436	GALDERMA BENELUX B.V.	BE
Pliaglis 70 mg/g + 70 mg/g, crème	DE/H/3278/001	BE434436	GALDERMA BENELUX B.V.	BE
Pliaglis 70mg/g + 70mg/g cream	DE/H/3278/001	PA1717/002/001	ADOH B.V.	IE
Pliaglis 70mg/g + 70mg/g cream	DE/H/3278/001	PL 39560/0012	ADOH B.V.	UK
Pliaglis 70mg/g + 70mg/g crema	DE/H/3278/001	76.730	INDUSTRIAL FARMACEUTICA CANTABRIA, S.A.	ES
Pliaglis 70mg/g + 70mg/g Creme	DE/H/3278/001	1-31484	ADOH B.V.	AT

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Pliaglis 70mg/g + 70mg/g Creme	DE/H/3278/001	84728.00.00	ADOH B.V.	DE
Pliaglis 70mg/g + 70mg/g Creme	DE/H/3278/001	2013060200	GALDERMA BENELUX B.V.	LU
Pliaglis 70mg/g + 70mg/g creme	DE/H/3278/001	5483508	LABORATORIOS GALDERMA SA – SUCURSAL EM PORTUGAL	PT
Pliaglis, 70 mg/g + 70 mg/g, krem	DE/H/3278/001	20451	ADOH B.V.	PL
Pliapel 70 mg/g + 70 mg/g krem	DE/H/3278/001	11-8378	ADOH B.V.	NO
Pliapel, creme	DE/H/3278/001	48871	ADOH B.V.	DK
Ralydan	SE/H/0762/001	41559	EUROCEPT INTERNATIONAL BV	DK
Rapydan 70 mg/70 mg emplâtre médicamenteux	SE/H/0762/001	BE 325832	EUROCEPT INTERNATIONAL BV	BE
Rapydan 70 mg/70 mg emplâtre médicamenteux	SE/H/0762/001	2008010017 (NAT: 0473933; 0473947; 0473951; 0473964; 0473978; 0473981)	EUROCEPT INTERNATIONAL BV	LU
Rapydan 70 mg/70 mg medicated plaster	SE/H/0762/001	PA 1591/001/001	EUROCEPT INTERNATIONAL BV	IE
Rapydan 70 mg/70 mg medicated plaster	SE/H/0762/001	PL 35068/0001	EUROCEPT INTERNATIONAL BV	UK
Rapydan 70 mg/70 mg medicinale pleister	SE/H/0762/001	BE 325832	EUROCEPT INTERNATIONAL BV	BE
Rapydan 70 mg/70 mg medicinale pleister	SE/H/0762/001	RVG 100315	EUROCEPT INTERNATIONAL BV	NL
Rapydan 70 mg/70 mg medicinskt plåster	SE/H/0762/001	22468	EUROCEPT INTERNATIONAL BV	SE

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Rapydan 70 mg/70 mg medisinert plaster	SE/H/0762/001	07-5016	EUROCEPT INTERNATIONAL BV	NO
Rapydan 70 mg/70 mg wirkstoffhaltiges Pflaster	SE/H/0762/001	BE325832	EUROCEPT INTERNATIONAL BV	BE
Rapydan 70 mg/70 mg wirkstoffhaltiges Pflaster	SE/H/0762/001	70270.00.00	EUROCEPT INTERNATIONAL BV	DE
Rapydan 70 mg/70 mg wirkstoffhaltiges Pflaster	SE/H/0762/001	1637/08010017	EUROCEPT INTERNATIONAL BV	LU
Rapydan 70 mg/70 mg wirkstoffhaltiges Pflaster Wirkstoffe: Lidocain / Tetracain	SE/H/0762/001	1-27348	EUROCEPT INTERNATIONAL BV	AT
Rapydan 70 mg/70 mg φαρμακούχο έμπλαστρο	SE/H/0762/001	022501	EUROCEPT INTERNATIONAL BV	CY
Rapydan 70 mg/70 mg φαρμακούχο έμπλαστρο	SE/H/0762/001	277880101	EUROCEPT INTERNATIONAL BV	GR
Rapydan 70 mg/70 mg φαρμακούχο έμπλαστρο	SE/H/0762/001	277880102	EUROCEPT INTERNATIONAL BV	GR
Rapydan 70 mg/70 mg φαρμακούχο έμπλαστρο	SE/H/0762/001	277880103	EUROCEPT INTERNATIONAL BV	GR
Rapydan 70 mg/70 mg φαρμακούχο έμπλαστρο	SE/H/0762/001	277880104	EUROCEPT INTERNATIONAL BV	GR
Rapydan 70 mg/70 mg φαρμακούχο έμπλαστρο	SE/H/0762/001	277880105	EUROCEPT INTERNATIONAL BV	GR
Rapydan 70 mg/70 mg φαρμακούχο έμπλαστρο	SE/H/0762/001	277880106	EUROCEPT INTERNATIONAL BV	GR
Rapydan 70 mg/70 mg, emplastro medicamentoso	SE/H/0762/001/MR	69581	EUROCEPT INTERNATIONAL BV	PT

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Rapydan, 70 mg + 70 mg, plaster leczniczy	SE/H/0762/001	14748	EUROCEPT INTERNATIONAL BV	PL
Velocaine 70 mg/70 mg gyógyszerezes tapasz	SE/H/0762/001/MR	OGYI-T-20742/01-06. (1X, 2X, 5X, 10X, 25X, 50X)	EUROCEPT INTERNATIONAL BV	HU