



EUROPEAN MEDICINES AGENCY
SCIENCE MEDICINES HEALTH

27 October 2016
EMA/735253/2016
Procedure Management and Committees Support

List of nationally authorised medicinal products

Active substance: eletriptan

Procedure no.: PSUSA/00001204/201602



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
RELERT 20 mg comprimidos recubiertos con película	UK/H/0434/001	64.122	PFIZER, S.L.	ES
Relert 20 mg comprimidos revestidos por película	UK/H/0432/001	3686789	LABORATÓRIOS PFIZER, LDA.	PT
RELERT 20 mg filmomhulde tabletten	UK/H/0432/001	BE227123	PFIZER S.A. (BELGIUM)	BE
RELERT 20 mg filmomhulde tabletten	UK/H/0432/001	BE227114	PFIZER S.A. (BELGIUM)	BE
RELERT 20 mg Filmtabletten	UK/H/0432/001	2007069300	PFIZER S.A. (BELGIUM)	LU
RELERT 20 mg, comprimé pelliculé	UK/H/0434/001	34009 357 877 4 3	PFIZER HOLDING FRANCE (S.C.A.)	FR
RELERT 20 mg, comprimé pelliculé	UK/H/0434/001	34009 357 876 8 2	PFIZER HOLDING FRANCE (S.C.A.)	FR
RELERT 20 mg, comprimé pelliculé	UK/H/0434/001	34009 357 878 0 4	PFIZER HOLDING FRANCE (S.C.A.)	FR
RELERT 20 mg, comprimé pelliculé	UK/H/0434/001	34009 357 874 5 3	PFIZER HOLDING FRANCE (S.C.A.)	FR
RELERT 20 mg, comprimé pelliculé	UK/H/0434/001	34009 357 875 1 4	PFIZER HOLDING FRANCE (S.C.A.)	FR
RELERT 20 mg, comprimé pelliculé	UK/H/0434/001	34009 357 873 9 2	PFIZER HOLDING FRANCE (S.C.A.)	FR
RELERT 40 mg comprimidos recubiertos con película	UK/H/0434/002	64.123	PFIZER, S.L.	ES
Relert 40 mg comprimidos revestidos por película	UK/H/0432/002	3688389	LABORATÓRIOS PFIZER, LDA.	PT
RELERT 40 mg filmdragerade tabletter	UK/H/0432/002	16425	PFIZER OY	FI
RELERT 40 mg filmomhulde tabletten	UK/H/0432/002	BE227157	PFIZER S.A. (BELGIUM)	BE
RELERT 40 mg	UK/H/0432/002	BE227166	PFIZER S.A. (BELGIUM)	BE

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Filmdragerade tabletter				
RELERT 40 mg Filmdragerade tabletter	UK/H/0432/002	2007069301	PFIZER S.A. (BELGIUM)	LU
RELERT 40 mg, comprimé pelliculé	UK/H/0434/002	34009 357 883 4 4	PFIZER HOLDING FRANCE (S.C.A.)	FR
RELERT 40 mg, comprimé pelliculé	UK/H/0434/002	34009 357 880 5 4	PFIZER HOLDING FRANCE (S.C.A.)	FR
RELERT 40 mg, comprimé pelliculé	UK/H/0434/002	34009 357 881 1 5	PFIZER HOLDING FRANCE (S.C.A.)	FR
RELERT 40 mg, comprimé pelliculé	UK/H/0434/002	34009 357 882 8 3	PFIZER HOLDING FRANCE (S.C.A.)	FR
RELERT 40 mg, comprimé pelliculé	UK/H/0434/002	34009 357 879 7 2	PFIZER HOLDING FRANCE (S.C.A.)	FR
RELERT 40 mg, comprimé pelliculé	UK/H/0434/002	34009 357 884 0 5	PFIZER HOLDING FRANCE (S.C.A.)	FR
RELERT® 40 mg film- coated tablets	UK/H/0434/002	PL 00057/0456	PFIZER LIMITED	UK
RELERT® 20 mg film- coated tablets	UK/H/0434/001	PL 00057/0455	PFIZER LIMITED	UK
RELPAK 20 mg comprimidos recubiertos con película	UK/H/0432/001	64.124	PFIZER, S.L.	ES
RELPAK 20 mg film- coated tablets	UK/H/0432/001	PA 822/6/1	PFIZER HEALTHCARE IRELAND	IE
Relpax 20 mg filmtablettor	UK/H/0432/001	17176	PFIZER AB	SE
Relpax 20 mg filmtablettor	UK/H/0432/001	01-2002	PFIZER AS	NO
RELPAK 20 mg filmom obalené tablety	not available	33/0177/01-S	PFIZER EUROPE MA EEIG	SK
RELPAK 20 mg filmsko obložene tablete	not available	H/02/01330/001	PFIZER LUXEMBOURG SARL	SI
RELPAK 20 mg filmtablettor	not available	OGYI-T-8249/02	PFIZER KFT.	HU

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
RELPAx 20 mg filmtabletta	not available	OGYI-T-8249/03	PFIZER KFT.	HU
RELPAx 20 mg filmtabletta	not available	OGYI-T-8249/01	PFIZER KFT.	HU
Relpax 20 mg, comprimate filmate	not available	664/2008/01	PFIZER EUROPE MA EEIG	RO
Relpax 20 mg, comprimate filmate	not available	664/2008/03	PFIZER EUROPE MA EEIG	RO
Relpax 20 mg, comprimate filmate	not available	664/2008/02	PFIZER EUROPE MA EEIG	RO
RELPAx 20 mg, comprimé pelliculé	UK/H/0432/001	34009 357 860 4 3	PFIZER HOLDING FRANCE (S.C.A.)	FR
RELPAx 20 mg, comprimé pelliculé	UK/H/0432/001	34009 357 863 3 3	PFIZER HOLDING FRANCE (S.C.A.)	FR
RELPAx 20 mg, comprimé pelliculé	UK/H/0432/001	34009 357 861 0 4	PFIZER HOLDING FRANCE (S.C.A.)	FR
RELPAx 20 mg, comprimé pelliculé	UK/H/0432/001	34009 357 859 6 1	PFIZER HOLDING FRANCE (S.C.A.)	FR
RELPAx 20 mg, comprimé pelliculé	UK/H/0432/001	34009 357 862 7 2	PFIZER HOLDING FRANCE (S.C.A.)	FR
RELPAx 20 mg, comprimé pelliculé	UK/H/0432/001	34009 357 865 6 2	PFIZER HOLDING FRANCE (S.C.A.)	FR
Relpax 20 mg, filmuhúðaðar töflur	UK/H/0432/001	IS/1/01/013/01	PFIZER APS	IS
Relpax 20, filmomhulde tabletten 20 mg	UK/H/0432/001	RVG 26578	PFIZER B.V.	NL
RELPAx 40 mg apvalkotās tabletes	not available	04-0253	PFIZER LIMITED	LV
RELPAx 40 mg comprimidos recubiertos con película	UK/H/0432/002	64.125	PFIZER, S.L.	ES
RELPAx 40 mg film-coated tablets	UK/H/0432/002	PA 822/6/2	PFIZER HEALTHCARE IRELAND	IE

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Relpax 40 mg filmdragerade tabletter	UK/H/0432/002	17177	PFIZER AB	SE
Relpax 40 mg filmdrasjerte tabletter	UK/H/0432/002	01-2003	PFIZER AS	NO
RELPAK 40 mg filmom obalené tablety	not available	33/0178/01-S	PFIZER EUROPE MA EEIG	SK
RELPAK 40 mg filmsko obložene tablete	not available	H/02/01330/002	PFIZER LUXEMBOURG SARL	SI
RELPAK 40 mg filmtabletta	not available	OGYI-T-8249/04	PFIZER KFT.	HU
RELPAK 40 mg filmtabletta	not available	OGYI-T-8249/06	PFIZER KFT.	HU
RELPAK 40 mg filmtabletta	not available	OGYI-T-8249/05	PFIZER KFT.	HU
RELPAK 40 mg potahované tablety	not available	33/304/00-C	PFIZER, SPOL. S R.O.	CZ
Relpax 40 mg, comprimate filmate	not available	665/2008/03	PFIZER EUROPE MA EEIG	RO
Relpax 40 mg, comprimate filmate	not available	665/2008/04	PFIZER EUROPE MA EEIG	RO
Relpax 40 mg, comprimate filmate	not available	665/2008/02	PFIZER EUROPE MA EEIG	RO
Relpax 40 mg, comprimate filmate	not available	665/2008/01	PFIZER EUROPE MA EEIG	RO
RELPAK 40 mg, comprimé pelliculé	UK/H/0432/002	34009 357 866 2 3	PFIZER HOLDING FRANCE (S.C.A.)	FR
RELPAK 40 mg, comprimé pelliculé	UK/H/0432/002	34009 357 871 6 3	PFIZER HOLDING FRANCE (S.C.A.)	FR
RELPAK 40 mg, comprimé pelliculé	UK/H/0432/002	34009 357 867 9 1	PFIZER HOLDING FRANCE (S.C.A.)	FR
RELPAK 40 mg, comprimé pelliculé	UK/H/0432/002	34009 357 869 1 3	PFIZER HOLDING FRANCE (S.C.A.)	FR
RELPAK 40 mg, comprimé pelliculé	UK/H/0432/002	34009 357 872 2 4	PFIZER HOLDING FRANCE (S.C.A.)	FR
RELPAK 40 mg, comprimé pelliculé	UK/H/0432/002	34009 357 868 5 2	PFIZER HOLDING FRANCE (S.C.A.)	FR

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Relpax 40 mg, filmuhúðaðar töflur	UK/H/0432/002	IS/1/01/013/02	PFIZER APS	IS
Relpax 40, filmomhulde tabletten 40 mg	UK/H/0432/002	RVG 26579	PFIZER B.V.	NL
RELPAK 40® mg film-coated tablets	UK/H/0432/002	PL 00057/0453	PFIZER LIMITED	UK
RELPAK 80 mg filmom obalené tablety	not available	33/0179/01-S	PFIZER EUROPE MA EEIG	SK
RELPAK 80 mg potahované tablety	not available	33/305/00-C	PFIZER, SPOL. S R.O.	CZ
RELPAK Επικαλυμμένα με λεπτό υμένιο δισκία 20 mg	UK/H/0432/001	58334/12-09-2008	PFIZER HELLAS, A.E.	GR
RELPAK Επικαλυμμένα με λεπτό υμένιο δισκία 40 mg	UK/H/0432/002	58335/12-09-2008	PFIZER HELLAS, A.E.	GR
Relpax, 20 mg, tabletki powlekane	not available	9499	PFIZER EUROPE MA EEIG	PL
Relpax, 40 mg, tabletki powlekane	not available	9500	PFIZER EUROPE MA EEIG	PL
Relpax, filmovertrukne tabletter	UK/H/0432/001	32308	PFIZER APS	DK
Relpax, filmovertrukne tabletter	UK/H/0432/002	32309	PFIZER APS	DK
RELPAK® 20 mg - Filmtabletten	UK/H/0432/001	1-24154	PFIZER CORPORATION AUSTRIA GESELLSCHAFT M.B.H.	AT
RELPAK® 20 mg compresse rivestite con film	UK/H/0432/001	035307103	PFIZER ITALIA S.R.L.	IT
RELPAK® 20 mg compresse rivestite con film	UK/H/0432/001	035307053	PFIZER ITALIA S.R.L.	IT
RELPAK® 20 mg compresse rivestite	UK/H/0432/001	035307089	PFIZER ITALIA S.R.L.	IT

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con film				
RELPAK® 20 mg comprese rivestite con film	UK/H/0432/001	035307180	PFIZER ITALIA S.R.L.	IT
RELPAK® 20 mg Filmdabletten	UK/H/0432/001	51508.00.00	PFIZER PHARMA GMBH	DE
RELPAK® 40 mg - Filmdabletten	UK/H/0432/002	1-24155	PFIZER CORPORATION AUSTRIA GESELLSCHAFT M.B.H.	AT
RELPAK® 40 mg comprese rivestite con film	UK/H/0432/002	035307204	PFIZER ITALIA S.R.L.	IT
RELPAK® 40 mg comprese rivestite con film	UK/H/0432/002	035307368	PFIZER ITALIA S.R.L.	IT
RELPAK® 40 mg Filmdabletten	UK/H/0432/002	51508.01.00	PFIZER PHARMA GMBH	DE
RELPAK® 20 mg film- coated tablets	UK/H/0432/001	PL 00057/0452	PFIZER LIMITED	UK
РЕЛПАКС 20 mg филмирани таблетки	not available	20020756	PFIZER EUROPE MA EEIG	BG
РЕЛПАКС 40 mg филмирани таблетки	not available	20020755	PFIZER EUROPE MA EEIG	BG