



EUROPEAN MEDICINES AGENCY
SCIENCE MEDICINES HEALTH

11 January 2018
EMA/32396/2018
Human Medicines Evaluation Division

List of nationally authorised medicinal products

Active substance(s): eprosartan / hydrochlorothiazide

Procedure No.: PSUSA/00001244/201704



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
Teveten Plus 600 mg/12,5 mg, επικαλυμμένα με λεπτό υμένιο δισκία	DE/H/0539/001	20235	MYLAN HEALTHCARE GMBH	CY
Teveten® Plus HCT 600 mg/12,5 mg, Filmtabletten	DE/H/0539/001	57956.00.00	MYLAN HEALTHCARE GMBH	DE
Teveten Plus 600 mg/12.5 mg, film-coated tablets	DE/H/0539/001	MA 1064/00602	BGP PRODUCTS LTD.	MT
CoTeveten 600 mg/12,5 mg filmsko obložene tablete	DE/H/0539/001	H/06/00427/010	MYLAN HEALTHCARE GMBH	SI
CoTeveten 600 mg/12,5 mg filmsko obložene tablete	DE/H/0539/001	H/06/00427/001	MYLAN HEALTHCARE GMBH	SI
CoTeveten 600 mg/12,5 mg filmsko obložene tablete	DE/H/0539/001	H/06/00427/002	MYLAN HEALTHCARE GMBH	SI
CoTeveten 600 mg/12,5 mg filmsko obložene tablete	DE/H/0539/001	H/06/00427/003	MYLAN HEALTHCARE GMBH	SI
CoTeveten 600 mg/12,5 mg filmsko obložene tablete	DE/H/0539/001	H/06/00427/004	MYLAN HEALTHCARE GMBH	SI
CoTeveten 600 mg/12,5 mg filmsko obložene tablete	DE/H/0539/001	H/06/00427/005	MYLAN HEALTHCARE GMBH	SI
CoTeveten 600 mg/12,5 mg filmsko obložene tablete	DE/H/0539/001	H/06/00427/006	MYLAN HEALTHCARE GMBH	SI
CoTeveten 600 mg/12,5 mg filmsko obložene tablete	DE/H/0539/001	H/06/00427/007	MYLAN HEALTHCARE GMBH	SI
CoTeveten 600 mg/12,5 mg filmsko obložene tablete	DE/H/0539/001	H/06/00427/008	MYLAN HEALTHCARE GMBH	SI
CoTeveten 600 mg/12,5 mg filmsko obložene tablete	DE/H/0539/001	H/06/00427/009	MYLAN HEALTHCARE GMBH	SI
Naviten Combi 600 mg/12,5 mg filmom obalené tablety	DE/H/0539/001	58/0328/08-S	TS PHARMA S.R.O.	SK
Теветен Плюс 600 mg/12,5 mg филмирани таблетки	not available	20040339	MYLAN HEALTHCARE GMBH	BG
Futuran PLUS 600 mg/12,5 mg, comprimidos recubiertos con película	not available	66.568	BGP PRODUCTS OPERATIONS SL	ES
Teveten Plus, filmomhulde tabletten 600/12,5 mg	not available	RVG 25926	BGP PRODUCTS B.V.	NL
REGULATEN PLUS 600 mg/12,5	not available	66.528	BGP PRODUCTS OPERATIONS SL	ES

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mg, comprimidos recubiertos con película				
NAVIXEN PLUS 600 mg/12,5 mg, comprimidos recubiertos con película	not available	66512	BGP PRODUCTS OPERATIONS SL	ES
Eprosartan comp.-CT 600 mg/12,5 mg Filmtabletten	not available	74095.00.00	ABZ-PHARMA GMBH	DE
Eprosartan-ratiopharm® comp. 600 mg/12,5 mg Filmtabletten	not available	57957.00.00	RATIOPHARM GMBH	DE
Teveten Plus 600 mg/12,5 mg Filmtabletten	DE/H/0382/001	1-25513	BGP PRODUCTS GES. M. B. H.	AT
Teveten Plus 600 mg/12,5 mg, Filmtabletten	DE/H/0382/001	BE269001	MYLAN EPD SPRL	BE
Teveten Plus 600 mg/12,5 mg, filmomhulde tabletten	DE/H/0382/001	BE269001	MYLAN EPD SPRL	BE
Teveten Plus 600 mg/12,5 mg, comprimés pelliculés	DE/H/0382/001	BE269001	MYLAN EPD SPRL	BE
Teveten Plus 600 mg/12,5 mg, comprimés pelliculés	DE/H/0382/001	2005010013	MYLAN EPD SPRL	LU
Teveten® Plus HCT 600 mg/12,5 mg, Filmtabletten	DE/H/0382/001	50260.00.00	MYLAN HEALTHCARE GMBH	DE
TEVETENS PLUS 600 mg/12,5 mg, comprimidos recubiertos con película	DE/H/0382/001	66.206	BGP PRODUCTS OPERATIONS SL	ES
Teveten Comp 600 mg/12,5 mg kalvopäällysteiset tabletit	DE/H/0382/001	19072	MYLAN FINLAND OY	FI
TEVETEN PLUS 600 mg/12,5 mg, επικαλυμμένα με λεπτό υμένιο δισκία	DE/H/0382/001	41060/24-6-2008	BGP PRODUCTS LTD (GREECE)	GR
TIARTAN 600 mg/12,5 mg, compresse rivestite con film	DE/H/0382/001	036772022	BGP PRODUCTS S.R.L.	IT
TIARTAN 600 mg/12,5 mg, compresse rivestite con film	DE/H/0382/001	036772034	BGP PRODUCTS S.R.L.	IT
TIARTAN 600 mg/12,5 mg, compresse rivestite con film	DE/H/0382/001	036772046	BGP PRODUCTS S.R.L.	IT
TIARTAN 600 mg/12,5 mg,	DE/H/0382/001	036772010	BGP PRODUCTS S.R.L.	IT

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comprime rivestite con film				
Teveten Comp 600 mg/12,5 mg, filmdrasjerte tabletter	DE/H/0382/001	03-2357	BGP PRODUCTS AS	NO
TEVETEN Plus 600 mg + 12,5 mg comprimidos revestidos por película	DE/H/0382/001	5175088	BGP PRODUCTS UNIPessoal, LDA.	PT
TEVETEN Plus 600 mg + 12,5 mg comprimidos revestidos por película	DE/H/0382/001	5175187	BGP PRODUCTS UNIPessoal, LDA.	PT
TEVETEN Plus 600 mg + 12,5 mg comprimidos revestidos por película	DE/H/0382/001	5174982	BGP PRODUCTS UNIPessoal, LDA.	PT
Teveten Comp 600 mg/12,5 mg filmdragerad tablett	DE/H/0382/001	20378	BGP PRODUCTS AB	SE
Teveten Comp 600 mg/ 12,5 mg filmdragerade tabletter	DE/H/0382/001	19072	MYLAN FINLAND OY	FI
Teveten Plus 600 mg/12,5 mg, Filmtabletten	DE/H/0382/001	2005010013	MYLAN EPD SPRL	LU
Teveten Plus 600 mg/12.5 mg, film-coated tablets	DE/H/0382/001	PA 2010/18/1	MYLAN IRE HEALTHCARE LIMITED	IE
Teveten® Plus HCT 600 mg/12,5 mg, Filmtabletten	DE/H/0538/001	57955.00.00	MYLAN HEALTHCARE GMBH	DE
COTEVETEN 600 mg/12,5 mg, comprimé pelliculé	DE/H/0538/001	34009 379 211 9 0	MYLAN MEDICAL SAS	FR
COTEVETEN 600 mg/12,5 mg, comprimé pelliculé	DE/H/0538/001	34009 376 103 0 8	MYLAN MEDICAL SAS	FR
COTEVETEN 600 mg/12,5 mg, comprimé pelliculé	DE/H/0538/001	34009 379 212 5 1	MYLAN MEDICAL SAS	FR
COTEVETEN 600 mg/12,5 mg, comprimé pelliculé	DE/H/0538/001	34009 376 104 7 6	MYLAN MEDICAL SAS	FR
COTEVETEN 600 mg/12,5 mg, comprimé pelliculé	DE/H/0538/001	34009 569 646 6 6	MYLAN MEDICAL SAS	FR
COTEVETEN 600 mg/12,5 mg, comprimé pelliculé	DE/H/0538/001	34009 376 102 4 7	MYLAN MEDICAL SAS	FR