



EUROPEAN MEDICINES AGENCY
SCIENCE MEDICINES HEALTH

12 January 2023
EMA/32202/2023 Human Medicines
Evaluation Division

List of nationally authorised medicinal products

Active substance: valsartan, hydrochlorothiazide / valsartan

Procedure no.: PSUSA/00010396/202204



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
Valsartan/Hydrochlorothiazide 160 mg/12.5 mg Film-coated Tablets	not available	PL 36687/0381	TORRENT PHARMA (UK) LTD.	XI
Valsartan/Hydrochlorothiazide 160 mg/25 mg Film-coated Tablets	not available	PL 36687/0382	TORRENT PHARMA (UK) LTD.	XI
Co-Vals 160 mg/12,5 mg comprimidos recubiertos con película	not available	65.797	ESTEVE PHARMACEUTICALS, S.A.	ES
Co-Vals Forte 320 mg/25 mg comprimidos recubiertos con película	not available	70.363	ESTEVE PHARMACEUTICALS, S.A.	ES
Diovan 160 mg filmdragerade tabletter	SE/H/0406/004	17495	NOVARTIS SVERIGE AB	SE
Diovan 3 mg/ml oral lösning	SE/H/0406/007	43158	NOVARTIS SVERIGE AB	SE
Diovan 320 mg filmdragerade tabletter	SE/H/0406/006	22840	NOVARTIS SVERIGE AB	SE
Diovan 40 mg filmdragerade tabletter	SE/H/0406/005	20464	NOVARTIS SVERIGE AB	SE
Diovan 80 mg filmdragerade tabletter	SE/H/0406/003	17494	NOVARTIS SVERIGE AB	SE
Valsocard HCT 80 mg/12.5 mg filmdragerade tabletter	not available	OGYI-T-21875/03	ACTAVIS GROUP PTC EHF.	HU
Valsocard HCT 160 mg/12.5 mg filmdragerade tabletter	not available	OGYI-T-21875/12	ACTAVIS GROUP PTC EHF.	HU
Valsocard HCT 80 mg/12.5 mg filmdragerade tabletter	not available	OGYI-T-21875/02	ACTAVIS GROUP PTC EHF.	HU
Valsocard HCT 160 mg/12.5 mg filmdragerade tabletter	not available	OGYI-T-21875/10	ACTAVIS GROUP PTC EHF.	HU
Valsocard HCT 80 mg/12.5 mg filmdragerade tabletter	not available	OGYI-T-21875/06	ACTAVIS GROUP PTC EHF.	HU
Valsocard HCT 80 mg/12.5 mg filmdragerade tabletter	not available	OGYI-T-21875/05	ACTAVIS GROUP PTC EHF.	HU
Valsocard HCT 80 mg/12.5 mg filmdragerade tabletter	not available	OGYI-T-21875/04	ACTAVIS GROUP PTC EHF.	HU
Valsocard HCT 160 mg/12.5 mg filmdragerade tabletter	not available	OGYI-T-21875/11	ACTAVIS GROUP PTC EHF.	HU
Valsocard HCT 160 mg/25 mg filmdragerade tabletter	not available	OGYI-T-21875/16	ACTAVIS GROUP PTC EHF.	HU
Valsocard HCT 160 mg/25 mg filmdragerade tabletter	not available	OGYI-T-21875/17	ACTAVIS GROUP PTC EHF.	HU
Valsocard HCT 160 mg/25 mg filmdragerade tabletter	not available	OGYI-T-21875/20	ACTAVIS GROUP PTC EHF.	HU

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filmtabletta				
Valsocard HCT 160 mg/25 mg filmtabletta	not available	OGYI-T-21875/21	ACTAVIS GROUP PTC EHF.	HU
Valsocard HCT 160 mg/25 mg filmtabletta	not available	OGYI-T-21875/18	ACTAVIS GROUP PTC EHF.	HU
Valsocard HCT 160 mg/25 mg filmtabletta	not available	OGYI-T-21875/19	ACTAVIS GROUP PTC EHF.	HU
Valsocard HCT 160 mg/12.5 mg filmtabletta	not available	OGYI-T-21875/13	ACTAVIS GROUP PTC EHF.	HU
Valsocard HCT 80 mg/12.5 mg filmtabletta	not available	OGYI-T-21875/07	ACTAVIS GROUP PTC EHF.	HU
Valsocard HCT 160 mg/12.5 mg filmtabletta	not available	OGYI-T-21875/09	ACTAVIS GROUP PTC EHF.	HU
Valsocard HCT 160 mg/12.5 mg filmtabletta	not available	OGYI-T-21875/14	ACTAVIS GROUP PTC EHF.	HU
Valsocard HCT 160 mg/12.5 mg filmtabletta	not available	OGYI-T-21875/08	ACTAVIS GROUP PTC EHF.	HU
Valsocard HCT 160 mg/25 mg filmtabletta	not available	OGYI-T-21875/15	ACTAVIS GROUP PTC EHF.	HU
Valsocard HCT 80 mg/12.5 mg filmtabletta	not available	OGYI-T-21875/01	ACTAVIS GROUP PTC EHF.	HU
Valsartan AbZ 120 mg Filmtabletten	SE/H/0746/003	70447.00.00	ABZ-PHARMA GMBH	DE
Valtsu 120 mg filmdragerade tabletter	SE/H/0746/003	26037	TEVA SWEDEN AB	SE
Valsartan-ratiopharm comp. 120 mg/12,5 mg Filmtabletten	AT/H/0514/001	90713.00.00	RATIOPHARM GMBH	DE
Valsartan / HCT ratiopharm 120mg/12,5mg Filmtabletten	AT/H/0514/001	135567	TEVA B.V	AT
Valsartan HCT Actavis 320 mg/25 mg plevele dengtos tabletės	FI/H/0857/002	LT/1/13/3193/012	ACTAVIS GROUP PTC EHF.	LT
Valsartan HCT Actavis 320 mg/25 mg plevele dengtos tabletės	FI/H/0857/002	LT/1/13/3193/013	ACTAVIS GROUP PTC EHF.	LT
Valsartan HCT Actavis 320 mg/25 mg plevele dengtos tabletės	FI/H/0857/002	LT/1/13/3193/015	ACTAVIS GROUP PTC EHF.	LT
Valsartan HCT Actavis 320 mg/25 mg	FI/H/0857/002	LT/1/13/3193/014	ACTAVIS GROUP PTC EHF.	LT

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plevele dengtos tabletės				
Valsartan HCT Actavis 320 mg/25 mg plevele dengtos tabletės	FI/H/0857/002	LT/1/13/3193/016	ACTAVIS GROUP PTC EHF.	LT
Valsartan HCT Actavis 320 mg/25 mg plevele dengtos tabletės	FI/H/0857/002	LT/1/13/3193/017	ACTAVIS GROUP PTC EHF.	LT
Valsartan HCT Actavis 320 mg/25 mg plevele dengtos tabletės	FI/H/0857/002	LT/1/13/3193/018	ACTAVIS GROUP PTC EHF.	LT
Valsartan HCT Actavis 320 mg/25 mg plevele dengtos tabletės	FI/H/0857/002	LT/1/13/3193/019	ACTAVIS GROUP PTC EHF.	LT
Valsartan HCT Actavis 320 mg/25 mg plevele dengtos tabletės	FI/H/0857/002	LT/1/13/3193/020	ACTAVIS GROUP PTC EHF.	LT
Valsartan HCT Actavis 320 mg/25 mg plevele dengtos tabletės	FI/H/0857/002	LT/1/13/3193/021	ACTAVIS GROUP PTC EHF.	LT
Valsartan HCT Actavis 320 mg/25 mg plevele dengtos tabletės	FI/H/0857/002	LT/1/13/3193/022	ACTAVIS GROUP PTC EHF.	LT
Valsartan HCT Actavis 320 mg/12,5 mg plevele dengtos tabletės	FI/H/0857/001	LT/1/13/3193/001	ACTAVIS GROUP PTC EHF.	LT
Valsartan HCT Actavis 320 mg/12,5 mg plevele dengtos tabletės	FI/H/0857/001	LT/1/13/3193/002	ACTAVIS GROUP PTC EHF.	LT
Valsartan HCT Actavis 320 mg/12,5 mg plevele dengtos tabletės	FI/H/0857/001	LT/1/13/3193/003	ACTAVIS GROUP PTC EHF.	LT
Valsartan HCT Actavis 320 mg/12,5 mg plevele dengtos tabletės	FI/H/0857/001	LT/1/13/3193/004	ACTAVIS GROUP PTC EHF.	LT
Valsartan HCT Actavis 320 mg/12,5 mg plevele dengtos tabletės	FI/H/0857/001	LT/1/13/3193/005	ACTAVIS GROUP PTC EHF.	LT
Valsartan HCT Actavis 320 mg/12,5 mg plevele dengtos tabletės	FI/H/0857/001	LT/1/13/3193/006	ACTAVIS GROUP PTC EHF.	LT
Valsartan HCT Actavis 320 mg/12,5 mg plevele dengtos tabletės	FI/H/0857/001	LT/1/13/3193/007	ACTAVIS GROUP PTC EHF.	LT
Valsartan HCT Actavis 320 mg/12,5 mg plevele dengtos tabletės	FI/H/0857/001	LT/1/13/3193/008	ACTAVIS GROUP PTC EHF.	LT
Valsartan HCT Actavis 320 mg/12,5 mg plevele dengtos tabletės	FI/H/0857/001	LT/1/13/3193/009	ACTAVIS GROUP PTC EHF.	LT
Valsartan HCT Actavis 320 mg/12,5 mg plevele dengtos tabletės	FI/H/0857/001	LT/1/13/3193/010	ACTAVIS GROUP PTC EHF.	LT

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plevele dengtos tabletės				
Valsartan HCT Actavis 320 mg/12,5 mg plevele dengtos tabletės	FI/H/0857/001	LT/1/13/3193/011	ACTAVIS GROUP PTC EHF.	LT
COTAREG 160 mg/12,5 mg, comprimé pelliculé	SE/H/0565/002	34009 363 107 2 8	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/12,5 mg, comprimé pelliculé	SE/H/0565/002	34009 371 254 0 6	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/12,5 mg, comprimé pelliculé	SE/H/0565/002	34009 372 013 7 7	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/12,5 mg, comprimé pelliculé	SE/H/0565/002	34009 372 014 3 8	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/12,5 mg, comprimé pelliculé	SE/H/0565/002	34009 381 565 9 1	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/12,5 mg, comprimé pelliculé	SE/H/0565/002	34009 381 566 5 2	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/12,5 mg, comprimé pelliculé	SE/H/0565/002	34009 381 567 1 3	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/12,5 mg, comprimé pelliculé	SE/H/0565/002	34009 381 568 8 1	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/12,5 mg, comprimé pelliculé	SE/H/0565/002	34009 381 569 4 2	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/12,5 mg, comprimé pelliculé	SE/H/0565/002	34009 381 570 2 4	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/12,5 mg, comprimé pelliculé	SE/H/0565/002	34009 381 571 9 2	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/12,5 mg, comprimé pelliculé	SE/H/0565/002	34009 564 976 8 3	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/12,5 mg, comprimé pelliculé	SE/H/0565/002	34009 564 977 4 4	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/12,5 mg, comprimé pelliculé	SE/H/0565/002	34009 564 978 0 5	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/25 mg, comprimé pelliculé	SE/H/0565/003	34009 359 232 0 2	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/25 mg, comprimé	SE/H/0565/003	34009 371 394 7 2	NOVARTIS PHARMA S.A.S.	FR

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pelliculé				
COTAREG 160 mg/25 mg, comprimé pelliculé	SE/H/0565/003	34009 371 395 3 3	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/25 mg, comprimé pelliculé	SE/H/0565/003	34009 371 397 6 2	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/25 mg, comprimé pelliculé	SE/H/0565/003	34009 381 572 5 3	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/25 mg, comprimé pelliculé	SE/H/0565/003	34009 381 573 1 4	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/25 mg, comprimé pelliculé	SE/H/0565/003	34009 381 574 8 2	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/25 mg, comprimé pelliculé	SE/H/0565/003	34009 381 575 4 3	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/25 mg, comprimé pelliculé	SE/H/0565/003	34009 381 576 0 4	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/25 mg, comprimé pelliculé	SE/H/0565/003	34009 381 577 7 2	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/25 mg, comprimé pelliculé	SE/H/0565/003	34009 381 578 3 3	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/25 mg, comprimé pelliculé	SE/H/0565/003	34009 563 683 7 2	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/25 mg, comprimé pelliculé	SE/H/0565/003	34009 563 684 3 3	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/25 mg, comprimé pelliculé	SE/H/0565/003	34009 563 686 6 2	NOVARTIS PHARMA S.A.S.	FR
COTAREG 80 mg/12,5 mg, comprimé pelliculé	SE/H/0565/001	34009 344 300 5 3	NOVARTIS PHARMA S.A.S.	FR
COTAREG 80 mg/12,5 mg, comprimé pelliculé	SE/H/0565/001	34009 344 301 1 4	NOVARTIS PHARMA S.A.S.	FR
COTAREG 80 mg/12,5 mg, comprimé pelliculé	SE/H/0565/001	34009 344 302 8 2	NOVARTIS PHARMA S.A.S.	FR
COTAREG 80 mg/12,5 mg, comprimé pelliculé	SE/H/0565/001	34009 344 303 4 3	NOVARTIS PHARMA S.A.S.	FR
COTAREG 80 mg/12,5 mg, comprimé	SE/H/0565/001	34009 371 961 9 2	NOVARTIS PHARMA S.A.S.	FR

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pelliculé				
COTAREG 80 mg/12,5 mg, comprimé pelliculé	SE/H/0565/001	34009 371 962 5 3	NOVARTIS PHARMA S.A.S.	FR
COTAREG 80 mg/12,5 mg, comprimé pelliculé	SE/H/0565/001	34009 371 963 1 4	NOVARTIS PHARMA S.A.S.	FR
COTAREG 80 mg/12,5 mg, comprimé pelliculé	SE/H/0565/001	34009 381 557 6 1	NOVARTIS PHARMA S.A.S.	FR
COTAREG 80 mg/12,5 mg, comprimé pelliculé	SE/H/0565/001	34009 381 558 2 2	NOVARTIS PHARMA S.A.S.	FR
COTAREG 80 mg/12,5 mg, comprimé pelliculé	SE/H/0565/001	34009 381 559 9 0	NOVARTIS PHARMA S.A.S.	FR
COTAREG 80 mg/12,5 mg, comprimé pelliculé	SE/H/0565/001	34009 381 560 7 2	NOVARTIS PHARMA S.A.S.	FR
COTAREG 80 mg/12,5 mg, comprimé pelliculé	SE/H/0565/001	34009 381 561 3 3	NOVARTIS PHARMA S.A.S.	FR
COTAREG 80 mg/12,5 mg, comprimé pelliculé	SE/H/0565/001	34009 381 563 6 2	NOVARTIS PHARMA S.A.S.	FR
COTAREG 80 mg/12,5 mg, comprimé pelliculé	SE/H/0565/001	34009 381 564 2 3	NOVARTIS PHARMA S.A.S.	FR
NISISCO 80 mg/12,5 mg, comprimé pelliculé	not available	34009 371 518 8 7	NOVARTIS PHARMA S.A.S.	FR
NISISCO 80 mg/12,5 mg, comprimé pelliculé	not available	34009 371 519 4 8	NOVARTIS PHARMA S.A.S.	FR
NISISCO 160 mg/25 mg, comprimé pelliculé	not available	34009 373 893 0 3	NOVARTIS PHARMA S.A.S.	FR
NISISCO 80 mg/12,5 mg, comprimé pelliculé	not available	34009 379 031 0 3	NOVARTIS PHARMA S.A.S.	FR
NISISCO 80 mg/12,5 mg, comprimé pelliculé	not available	34009 379 032 7 1	NOVARTIS PHARMA S.A.S.	FR
NISISCO 80 mg/12,5 mg, comprimé pelliculé	not available	34009 379 033 3 2	NOVARTIS PHARMA S.A.S.	FR
NISISCO 80 mg/12,5 mg, comprimé pelliculé	not available	34009 379 035 6 1	NOVARTIS PHARMA S.A.S.	FR
NISISCO 80 mg/12,5 mg, comprimé	not available	34009 379 036 2 2	NOVARTIS PHARMA S.A.S.	FR

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pelliculé				
NISISCO 80 mg/12,5 mg, comprimé pelliculé	not available	34009 379 037 9 0	NOVARTIS PHARMA S.A.S.	FR
NISISCO 160 mg/12,5 mg, comprimé pelliculé	not available	34009 379 039 1 2	NOVARTIS PHARMA S.A.S.	FR
NISISCO 160 mg/12,5 mg, comprimé pelliculé	not available	34009 379 041 6 2	NOVARTIS PHARMA S.A.S.	FR
NISISCO 160 mg/12,5 mg, comprimé pelliculé	not available	34009 379 042 2 3	NOVARTIS PHARMA S.A.S.	FR
NISISCO 160 mg/12,5 mg, comprimé pelliculé	not available	34009 379 043 9 1	NOVARTIS PHARMA S.A.S.	FR
NISISCO 160 mg/12,5 mg, comprimé pelliculé	not available	34009 379 044 5 2	NOVARTIS PHARMA S.A.S.	FR
NISISCO 160 mg/12,5 mg, comprimé pelliculé	not available	34009 379 045 1 3	NOVARTIS PHARMA S.A.S.	FR
NISISCO 160 mg/25 mg, comprimé pelliculé	not available	34009 379 047 4 2	NOVARTIS PHARMA S.A.S.	FR
NISISCO 160 mg/25 mg, comprimé pelliculé	not available	34009 379 048 0 3	NOVARTIS PHARMA S.A.S.	FR
NISISCO 160 mg/25 mg, comprimé pelliculé	not available	34009 379 049 7 1	NOVARTIS PHARMA S.A.S.	FR
NISISCO 160 mg/25 mg, comprimé pelliculé	not available	34009 379 050 5 3	NOVARTIS PHARMA S.A.S.	FR
NISISCO 160 mg/25 mg, comprimé pelliculé	not available	34009 379 051 1 4	NOVARTIS PHARMA S.A.S.	FR
NISISCO 160 mg/25 mg, comprimé pelliculé	not available	34009 379 052 8 2	NOVARTIS PHARMA S.A.S.	FR
NISISCO 80 mg/12,5 mg, comprimé pelliculé	not available	34009 345 414 4 5	NOVARTIS PHARMA S.A.S.	FR
NISISCO 80 mg/12,5 mg, comprimé pelliculé	not available	34009 345 415 0 6	NOVARTIS PHARMA S.A.S.	FR
NISISCO 80 mg/12,5 mg, comprimé pelliculé	not available	34009 345 416 7 4	NOVARTIS PHARMA S.A.S.	FR
NISISCO 80 mg/12,5 mg, comprimé	not available	34009 345 417 3 5	NOVARTIS PHARMA S.A.S.	FR

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pelliculé				
NISISCO 160 mg/25 mg, comprimé pelliculé	not available	34009 359 237 2 1	NOVARTIS PHARMA S.A.S.	FR
NISISCO 160 mg/12,5 mg, comprimé pelliculé	not available	34009 363 108 9 6	NOVARTIS PHARMA S.A.S.	FR
NISISCO 160 mg/25 mg, comprimé pelliculé	not available	34009 371 492 9 7	NOVARTIS PHARMA S.A.S.	FR
NISISCO 160 mg/25 mg, comprimé pelliculé	not available	34009 371 493 5 8	NOVARTIS PHARMA S.A.S.	FR
NISISCO 160 mg/12,5 mg, comprimé pelliculé	not available	34009 371 514 2 9	NOVARTIS PHARMA S.A.S.	FR
NISISCO 160 mg/12,5 mg, comprimé pelliculé	not available	34009 371 515 9 7	NOVARTIS PHARMA S.A.S.	FR
NISISCO 160 mg/12,5 mg, comprimé pelliculé	not available	34009 371 516 5 8	NOVARTIS PHARMA S.A.S.	FR
NISISCO 80 mg/12,5 mg, comprimé pelliculé	not available	34009 371 517 1 9	NOVARTIS PHARMA S.A.S.	FR
NISISCO 160 mg/25 mg, comprimé pelliculé	not available	34009 563 687 2 3	NOVARTIS PHARMA S.A.S.	FR
NISISCO 160 mg/25 mg, comprimé pelliculé	not available	34009 563 688 9 1	NOVARTIS PHARMA S.A.S.	FR
NISISCO 160 mg/25 mg, comprimé pelliculé	not available	34009 563 689 5 2	NOVARTIS PHARMA S.A.S.	FR
NISISCO 160 mg/12,5 mg, comprimé pelliculé	not available	34009 564 979 7 3	NOVARTIS PHARMA S.A.S.	FR
NISISCO 160 mg/12,5 mg, comprimé pelliculé	not available	34009 564 980 5 5	NOVARTIS PHARMA S.A.S.	FR
NISISCO 160 mg/12,5 mg, comprimé pelliculé	not available	34009 564 981 1 6	NOVARTIS PHARMA S.A.S.	FR
NISISCO 160 mg/12,5 mg, comprimé pelliculé	not available	34009 572 099 2 6	NOVARTIS PHARMA S.A.S.	FR
NISISCO 160 mg/25 mg, comprimé pelliculé	not available	34009 572 100 0 7	NOVARTIS PHARMA S.A.S.	FR
NISISCO 80 mg/12,5 mg, comprimé	not available	34009 572 101 7 5	NOVARTIS PHARMA S.A.S.	FR

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pelliculé				
Cotareg 80 mg/12,5 mg compresse rivestite con film	SE/H/0565/001	034114037	NOVARTIS FARMA S.P.A.	IT
Cotareg 80 mg/12,5 mg compresse rivestite con film	SE/H/0565/001	034114049	NOVARTIS FARMA S.P.A.	IT
Cotareg 160 mg/12,5 mg compresse rivestite con film	SE/H/0565/002	034114417	NOVARTIS FARMA S.P.A.	IT
Cotareg 160 mg/12,5 mg compresse rivestite con film	SE/H/0565/002	034114429	NOVARTIS FARMA S.P.A.	IT
Cotareg 160 mg/25 mg compresse rivestite con film	SE/H/0565/003	034114431	NOVARTIS FARMA S.P.A.	IT
Cotareg 160 mg/25 mg compresse rivestite con film	SE/H/0565/003	034114443	NOVARTIS FARMA S.P.A.	IT
Cotareg 80 mg/12,5 mg compresse rivestite con film	SE/H/0565/001	034114052	NOVARTIS FARMA S.P.A.	IT
Cotareg 80 mg/12,5 mg compresse rivestite con film	SE/H/0565/001	034114381	NOVARTIS FARMA S.P.A.	IT
Cotareg 80 mg/12,5 mg compresse rivestite con film	SE/H/0565/001	034114393	NOVARTIS FARMA S.P.A.	IT
Cotareg 80 mg/12,5 mg compresse rivestite con film	SE/H/0565/001	034114405	NOVARTIS FARMA S.P.A.	IT
Cotareg 160 mg/12,5 mg compresse rivestite con film	SE/H/0565/002	034114506	NOVARTIS FARMA S.P.A.	IT
Cotareg 160 mg/12,5 mg compresse rivestite con film	SE/H/0565/002	034114518	NOVARTIS FARMA S.P.A.	IT
Cotareg 160 mg/12,5 mg compresse rivestite con film	SE/H/0565/002	034114520	NOVARTIS FARMA S.P.A.	IT
Cotareg 160 mg/12,5 mg compresse rivestite con film	SE/H/0565/002	034114532	NOVARTIS FARMA S.P.A.	IT
Cotareg 160 mg/12,5 mg compresse rivestite con film	SE/H/0565/002	034114544	NOVARTIS FARMA S.P.A.	IT
Cotareg 80 mg/12,5 mg compresse rivestite con film	SE/H/0565/001	034114456	NOVARTIS FARMA S.P.A.	IT
Cotareg 80 mg/12,5 mg compresse	SE/H/0565/001	034114468	NOVARTIS FARMA S.P.A.	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
rivestite con film				
Cotareg 80 mg/12,5 mg compresse rivestite con film	SE/H/0565/001	034114470	NOVARTIS FARMA S.P.A.	IT
Cotareg 80 mg/12,5 mg compresse rivestite con film	SE/H/0565/001	034114482	NOVARTIS FARMA S.P.A.	IT
Cotareg 80 mg/12,5 mg compresse rivestite con film	SE/H/0565/001	034114494	NOVARTIS FARMA S.P.A.	IT
Cotareg 160 mg/12,5 mg compresse rivestite con film	SE/H/0565/002	034114557	NOVARTIS FARMA S.P.A.	IT
Cotareg 160 mg/12,5 mg compresse rivestite con film	SE/H/0565/002	034114569	NOVARTIS FARMA S.P.A.	IT
Cotareg 160 mg/12,5 mg compresse rivestite con film	SE/H/0565/002	034114571	NOVARTIS FARMA S.P.A.	IT
Cotareg 160 mg/25 mg compresse rivestite con film	SE/H/0565/003	034114583	NOVARTIS FARMA S.P.A.	IT
Cotareg 160 mg/25 mg compresse rivestite con film	SE/H/0565/003	034114595	NOVARTIS FARMA S.P.A.	IT
Cotareg 160 mg/25 mg compresse rivestite con film	SE/H/0565/003	034114607	NOVARTIS FARMA S.P.A.	IT
Cotareg 160 mg/25 mg compresse rivestite con film	SE/H/0565/003	034114619	NOVARTIS FARMA S.P.A.	IT
Cotareg 160 mg/25 mg compresse rivestite con film	SE/H/0565/003	034114621	NOVARTIS FARMA S.P.A.	IT
Cotareg 160 mg/25 mg compresse rivestite con film	SE/H/0565/003	034114633	NOVARTIS FARMA S.P.A.	IT
Cotareg 160 mg/25 mg compresse rivestite con film	SE/H/0565/003	034114645	NOVARTIS FARMA S.P.A.	IT
Cotareg 160 mg/25 mg compresse rivestite con film	SE/H/0565/003	034114658	NOVARTIS FARMA S.P.A.	IT
Cotareg 80 mg/12,5 mg compresse rivestite con film	SE/H/0565/001	034114660	NOVARTIS FARMA S.P.A.	IT
Cotareg 80 mg/12,5 mg compresse rivestite con film	SE/H/0565/001	034114672	NOVARTIS FARMA S.P.A.	IT
Cotareg 80 mg/12,5 mg compresse	SE/H/0565/001	034114684	NOVARTIS FARMA S.P.A.	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
rivestite con film				
Cotareg 80 mg/12,5 mg compresse rivestite con film	SE/H/0565/001	034114696	NOVARTIS FARMA S.P.A.	IT
Cotareg 80 mg/12,5 mg compresse rivestite con film	SE/H/0565/001	034114708	NOVARTIS FARMA S.P.A.	IT
Cotareg 160 mg/12,5 mg compresse rivestite con film	SE/H/0565/002	034114064	NOVARTIS FARMA S.P.A.	IT
Cotareg 160 mg/12,5 mg compresse rivestite con film	SE/H/0565/002	034114076	NOVARTIS FARMA S.P.A.	IT
Cotareg 160 mg/12,5 mg compresse rivestite con film	SE/H/0565/002	034114088	NOVARTIS FARMA S.P.A.	IT
Cotareg 160 mg/12,5 mg compresse rivestite con film	SE/H/0565/002	034114090	NOVARTIS FARMA S.P.A.	IT
Cotareg 160 mg/12,5 mg compresse rivestite con film	SE/H/0565/002	034114102	NOVARTIS FARMA S.P.A.	IT
Cotareg 160 mg/12,5 mg compresse rivestite con film	SE/H/0565/002	034114114	NOVARTIS FARMA S.P.A.	IT
Cotareg 160 mg/12,5 mg compresse rivestite con film	SE/H/0565/002	034114126	NOVARTIS FARMA S.P.A.	IT
Cotareg 160 mg/25 mg compresse rivestite con film	SE/H/0565/003	034114138	NOVARTIS FARMA S.P.A.	IT
Cotareg 160 mg/25 mg compresse rivestite con film	SE/H/0565/003	034114140	NOVARTIS FARMA S.P.A.	IT
Cotareg 160 mg/25 mg compresse rivestite con film	SE/H/0565/003	034114153	NOVARTIS FARMA S.P.A.	IT
Cotareg 160 mg/25 mg compresse rivestite con film	SE/H/0565/003	034114165	NOVARTIS FARMA S.P.A.	IT
Cotareg 160 mg/25 mg compresse rivestite con film	SE/H/0565/003	034114177	NOVARTIS FARMA S.P.A.	IT
Cotareg 160 mg/25 mg compresse rivestite con film	SE/H/0565/003	034114189	NOVARTIS FARMA S.P.A.	IT
Cotareg 160 mg/25 mg compresse rivestite con film	SE/H/0565/003	034114191	NOVARTIS FARMA S.P.A.	IT
Cotareg 320 mg/12,5 mg compresse	SE/H/0565/004	034114203	NOVARTIS FARMA S.P.A.	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
rivestite con film				
Cotareg 320 mg/12,5 mg compresse rivestite con film	SE/H/0565/004	034114215	NOVARTIS FARMA S.P.A.	IT
Cotareg 320 mg/12,5 mg compresse rivestite con film	SE/H/0565/004	034114227	NOVARTIS FARMA S.P.A.	IT
Cotareg 320 mg/12,5 mg compresse rivestite con film	SE/H/0565/004	034114239	NOVARTIS FARMA S.P.A.	IT
Cotareg 320 mg/12,5 mg compresse rivestite con film	SE/H/0565/004	034114241	NOVARTIS FARMA S.P.A.	IT
Cotareg 320 mg/12,5 mg compresse rivestite con film	SE/H/0565/004	034114254	NOVARTIS FARMA S.P.A.	IT
Cotareg 320 mg/12,5 mg compresse rivestite con film	SE/H/0565/004	034114266	NOVARTIS FARMA S.P.A.	IT
Cotareg 320 mg/12,5 mg compresse rivestite con film	SE/H/0565/004	034114278	NOVARTIS FARMA S.P.A.	IT
Cotareg 320 mg/12,5 mg compresse rivestite con film	SE/H/0565/004	034114367	NOVARTIS FARMA S.P.A.	IT
Cotareg 320 mg/25 mg compresse rivestite con film	SE/H/0565/005	034114280	NOVARTIS FARMA S.P.A.	IT
Cotareg 320 mg/25 mg compresse rivestite con film	SE/H/0565/005	034114292	NOVARTIS FARMA S.P.A.	IT
Cotareg 320 mg/25 mg compresse rivestite con film	SE/H/0565/005	034114304	NOVARTIS FARMA S.P.A.	IT
Cotareg 320 mg/25 mg compresse rivestite con film	SE/H/0565/005	034114316	NOVARTIS FARMA S.P.A.	IT
Cotareg 320 mg/25 mg compresse rivestite con film	SE/H/0565/005	034114328	NOVARTIS FARMA S.P.A.	IT
Cotareg 320 mg/25 mg compresse rivestite con film	SE/H/0565/005	034114330	NOVARTIS FARMA S.P.A.	IT
Cotareg 320 mg/25 mg compresse rivestite con film	SE/H/0565/005	034114342	NOVARTIS FARMA S.P.A.	IT
Cotareg 320 mg/25 mg compresse rivestite con film	SE/H/0565/005	034114355	NOVARTIS FARMA S.P.A.	IT
Cotareg 320 mg/25 mg compresse	SE/H/0565/005	034114379	NOVARTIS FARMA S.P.A.	IT

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rivestite con film				
Cotareg 80 mg/12,5 mg compresse rivestite con film	SE/H/0565/001	034114013	NOVARTIS FARMA S.P.A.	IT
Cotareg 80 mg/12,5 mg compresse rivestite con film	SE/H/0565/001	034114025	NOVARTIS FARMA S.P.A.	IT
Valsartan Teva 320 mg compresse rivestite con film	DK/H/1517/004	040149496	TEVA ITALIA S.R.L.	IT
Valsartan Teva 320 mg compresse rivestite con film	DK/H/1517/004	040149609	TEVA ITALIA S.R.L.	IT
Valsartan Teva 320 mg compresse rivestite con film	DK/H/1517/004	040149534	TEVA ITALIA S.R.L.	IT
Valsartan Teva 320 mg compresse rivestite con film	DK/H/1517/004	040149561	TEVA ITALIA S.R.L.	IT
Valsartan Teva 320 mg compresse rivestite con film	DK/H/1517/004	040149611	TEVA ITALIA S.R.L.	IT
Valsartan Teva 320 mg compresse rivestite con film	DK/H/1517/004	040149662	TEVA ITALIA S.R.L.	IT
Valsartan Teva 320 mg compresse rivestite con film	DK/H/1517/004	040149508	TEVA ITALIA S.R.L.	IT
Valsartan Teva 320 mg compresse rivestite con film	DK/H/1517/004	040149559	TEVA ITALIA S.R.L.	IT
Valsartan Teva 320 mg compresse rivestite con film	DK/H/1517/004	040149585	TEVA ITALIA S.R.L.	IT
Valsartan Teva 320 mg compresse rivestite con film	DK/H/1517/004	040149484	TEVA ITALIA S.R.L.	IT
Valsartan Teva 320 mg compresse rivestite con film	DK/H/1517/004	040149647	TEVA ITALIA S.R.L.	IT
Valsartan Teva 320 mg compresse rivestite con film	DK/H/1517/004	040149635	TEVA ITALIA S.R.L.	IT
Valsartan Teva 320 mg compresse rivestite con film	DK/H/1517/004	040149623	TEVA ITALIA S.R.L.	IT
Valsartan Teva 320 mg compresse rivestite con film	DK/H/1517/004	040149510	TEVA ITALIA S.R.L.	IT
Valsartan Teva 320 mg compresse	DK/H/1517/004	040149522	TEVA ITALIA 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
rivestite con film				
Valsartan Teva 320 mg compresse rivestite con film	DK/H/1517/004	040149546	TEVA ITALIA S.R.L.	IT
Valsartan Teva 320 mg compresse rivestite con film	DK/H/1517/004	040149597	TEVA ITALIA S.R.L.	IT
Valsartan Teva 320 mg compresse rivestite con film	DK/H/1517/004	040149573	TEVA ITALIA S.R.L.	IT
Valsartan Teva 320 mg compresse rivestite con film	DK/H/1517/004	040149650	TEVA ITALIA S.R.L.	IT
Valsartan ratiopharm 120 mg filmdragerade tabletter	SE/H/0747/003	26041	RATIOPHARM GMBH	SE
Valsartan-ratiopharm 120 mg Filmtabletten	SE/H/0747/003	70451.00.00	RATIOPHARM GMBH	DE
Valsocard 80 mg fimtabletta	not available	OGYI-T-21874/12	ACTAVIS GROUP PTC EHF.	HU
Valsocard 80 mg fimtabletta	not available	OGYI-T-21874/11	ACTAVIS GROUP PTC EHF.	HU
Valsocard 80 mg fimtabletta	not available	OGYI-T-21874/10	ACTAVIS GROUP PTC EHF.	HU
Valsocard 80 mg fimtabletta	not available	OGYI-T-21874/09	ACTAVIS GROUP PTC EHF.	HU
Valsocard 80 mg fimtabletta	not available	OGYI-T-21874/14	ACTAVIS GROUP PTC EHF.	HU
Valsocard 80 mg fimtabletta	not available	OGYI-T-21874/13	ACTAVIS GROUP PTC EHF.	HU
Valsocard 160 mg filmtabletta	not available	OGYI-T-21874/16	ACTAVIS GROUP PTC EHF.	HU
Valsocard 160 mg filmtabletta	not available	OGYI-T-21874/17	ACTAVIS GROUP PTC EHF.	HU
Valsocard 160 mg filmtabletta	not available	OGYI-T-21874/19	ACTAVIS GROUP PTC EHF.	HU
Valsocard 160 mg filmtabletta	not available	OGYI-T-21874/18	ACTAVIS GROUP PTC EHF.	HU
Valsocard 160 mg filmtabletta	not available	OGYI-T-21874/20	ACTAVIS GROUP PTC EHF.	HU
Valsocard 160 mg filmtabletta	not available	OGYI-T-21874/21	ACTAVIS GROUP PTC EHF.	HU
Valsocard 80 mg fimtabletta	not available	OGYI-T-21874/08	ACTAVIS GROUP PTC EHF.	HU
Valsocard 160 mg fimtabletta	not available	OGYI-T-21874/15	ACTAVIS GROUP PTC EHF.	HU
TAREG 160 mg, comprimé pelliculé	SE/H/0406/004	34009 356 905 4 8	NOVARTIS PHARMA S.A.S.	FR
TAREG 160 mg, comprimé pelliculé	SE/H/0406/004	34009 356 906 0 9	NOVARTIS PHARMA S.A.S.	FR
TAREG 160 mg, comprimé pelliculé	SE/H/0406/004	34009 356 907 7 7	NOVARTIS PHARMA S.A.S.	FR
TAREG 160 mg, comprimé pelliculé	SE/H/0406/004	34009 371 390 1 4	NOVARTIS PHARMA S.A.S.	FR
TAREG 160 mg, comprimé pelliculé	SE/H/0406/004	34009 371 391 8 2	NOVARTIS PHARMA S.A.S.	FR
TAREG 160 mg, comprimé pelliculé	SE/H/0406/004	34009 381 550 1 3	NOVARTIS PHARMA S.A.S.	FR

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TAREG 160 mg, comprimé pelliculé	SE/H/0406/004	34009 381 551 8 1	NOVARTIS PHARMA S.A.S.	FR
TAREG 160 mg, comprimé pelliculé	SE/H/0406/004	34009 381 552 4 2	NOVARTIS PHARMA S.A.S.	FR
TAREG 160 mg, comprimé pelliculé	SE/H/0406/004	34009 381 553 0 3	NOVARTIS PHARMA S.A.S.	FR
TAREG 160 mg, comprimé pelliculé	SE/H/0406/004	34009 381 555 3 2	NOVARTIS PHARMA S.A.S.	FR
TAREG 40 mg, comprimé pelliculé sécable	SE/H/0406/005	34009 369 583 0 2	NOVARTIS PHARMA S.A.S.	FR
TAREG 40 mg, comprimé pelliculé sécable	SE/H/0406/005	34009 369 584 7 0	NOVARTIS PHARMA S.A.S.	FR
TAREG 40 mg, comprimé pelliculé sécable	SE/H/0406/005	34009 369 585 3 1	NOVARTIS PHARMA S.A.S.	FR
TAREG 40 mg, comprimé pelliculé sécable	SE/H/0406/005	34009 371 380 6 2	NOVARTIS PHARMA S.A.S.	FR
TAREG 40 mg, comprimé pelliculé sécable	SE/H/0406/005	34009 371 381 2 3	NOVARTIS PHARMA S.A.S.	FR
TAREG 40 mg, comprimé pelliculé sécable	SE/H/0406/005	34009 381 538 1 1	NOVARTIS PHARMA S.A.S.	FR
TAREG 40 mg, comprimé pelliculé sécable	SE/H/0406/005	34009 381 539 8 9	NOVARTIS PHARMA S.A.S.	FR
TAREG 40 mg, comprimé pelliculé sécable	SE/H/0406/005	34009 381 540 6 1	NOVARTIS PHARMA S.A.S.	FR
TAREG 40 mg, comprimé pelliculé sécable	SE/H/0406/005	34009 381 541 2 2	NOVARTIS PHARMA S.A.S.	FR
TAREG 40 mg, comprimé pelliculé sécable	SE/H/0406/005	34009 381 543 5 1	NOVARTIS PHARMA S.A.S.	FR
TAREG 80 mg, comprimé pelliculé	SE/H/0406/003	34009 356 901 9 7	NOVARTIS PHARMA S.A.S.	FR
TAREG 80 mg, comprimé pelliculé	SE/H/0406/003	34009 356 902 5 8	NOVARTIS PHARMA S.A.S.	FR
TAREG 80 mg, comprimé pelliculé	SE/H/0406/003	34009 356 903 1 9	NOVARTIS PHARMA S.A.S.	FR
TAREG 80 mg, comprimé pelliculé	SE/H/0406/003	34009 371 385 8 1	NOVARTIS PHARMA S.A.S.	FR
TAREG 80 mg, comprimé pelliculé	SE/H/0406/003	34009 371 386 4 2	NOVARTIS PHARMA S.A.S.	FR
TAREG 80 mg, comprimé pelliculé	SE/H/0406/003	34009 381 544 1 2	NOVARTIS PHARMA S.A.S.	FR
TAREG 80 mg, comprimé pelliculé	SE/H/0406/003	34009 381 545 8 0	NOVARTIS PHARMA S.A.S.	FR
TAREG 80 mg, comprimé pelliculé	SE/H/0406/003	34009 381 546 4 1	NOVARTIS PHARMA S.A.S.	FR
TAREG 80 mg, comprimé pelliculé	SE/H/0406/003	34009 381 547 0 2	NOVARTIS PHARMA S.A.S.	FR

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TAREG 80 mg, comprimé pelliculé	SE/H/0406/003	34009 381 549 3 1	NOVARTIS PHARMA S.A.S.	FR
Tareg 3 mg/ml, solution buvable	SE/H/0406/007	34009 491 474 8 9	NOVARTIS PHARMA S.A.S.	FR
NISIS 80 mg, comprimé pelliculé	not available	34009 372 290 0 5	NOVARTIS PHARMA S.A.S.	FR
NISIS 80 mg, comprimé pelliculé	not available	34009 372 291 7 3	NOVARTIS PHARMA S.A.S.	FR
NISIS 80 mg, comprimé pelliculé	not available	34009 373 891 8 1	NOVARTIS PHARMA S.A.S.	FR
NISIS 160 mg, comprimé pelliculé	not available	34009 373 892 4 2	NOVARTIS PHARMA S.A.S.	FR
NISIS 80 mg, comprimé pelliculé	not available	34009 379 061 7 3	NOVARTIS PHARMA S.A.S.	FR
NISIS 80 mg, comprimé pelliculé	not available	34009 379 062 3 4	NOVARTIS PHARMA S.A.S.	FR
NISIS 80 mg, comprimé pelliculé	not available	34009 379 064 6 3	NOVARTIS PHARMA S.A.S.	FR
NISIS 80 mg, comprimé pelliculé	not available	34009 379 065 2 4	NOVARTIS PHARMA S.A.S.	FR
NISIS 80 mg, comprimé pelliculé	not available	34009 379 066 9 2	NOVARTIS PHARMA S.A.S.	FR
NISIS 80 mg, comprimé pelliculé	not available	34009 379 067 5 3	NOVARTIS PHARMA S.A.S.	FR
NISIS 160 mg, comprimé pelliculé	not available	34009 379 068 1 4	NOVARTIS PHARMA S.A.S.	FR
NISIS 160 mg, comprimé pelliculé	not available	34009 379 069 8 2	NOVARTIS PHARMA S.A.S.	FR
NISIS 80 mg, comprimé pelliculé	not available	34009 356 937 3 0	NOVARTIS PHARMA S.A.S.	FR
NISIS 80 mg, comprimé pelliculé	not available	34009 356 939 6 9	NOVARTIS PHARMA S.A.S.	FR
NISIS 80 mg, comprimé pelliculé	not available	34009 356 940 4 1	NOVARTIS PHARMA S.A.S.	FR
NISIS 80 mg, comprimé pelliculé	not available	34009 356 941 0 2	NOVARTIS PHARMA S.A.S.	FR
NISIS 80 mg, comprimé pelliculé	not available	34009 356 942 7 0	NOVARTIS PHARMA S.A.S.	FR
NISIS 80 mg, comprimé pelliculé	not available	34009 356 943 3 1	NOVARTIS PHARMA S.A.S.	FR
NISIS 160 mg, comprimé pelliculé	not available	34009 356 945 6 0	NOVARTIS PHARMA S.A.S.	FR
NISIS 160 mg, comprimé pelliculé	not available	34009 356 946 2 1	NOVARTIS PHARMA S.A.S.	FR
NISIS 160 mg, comprimé pelliculé	not available	34009 356 947 9 9	NOVARTIS PHARMA S.A.S.	FR
NISIS 160 mg, comprimé pelliculé	not available	34009 356 948 5 0	NOVARTIS PHARMA S.A.S.	FR
NISIS 160 mg, comprimé pelliculé	not available	34009 356 958 0 2	NOVARTIS PHARMA S.A.S.	FR
NISIS 160 mg, comprimé pelliculé	not available	34009 356 959 7 0	NOVARTIS PHARMA S.A.S.	FR
NISIS 160 mg, comprimé pelliculé	not available	34009 371 494 1 9	NOVARTIS PHARMA S.A.S.	FR
NISIS 160 mg, comprimé pelliculé	not available	34009 371 495 8 7	NOVARTIS PHARMA S.A.S.	FR
NISIS 160 mg, comprimé pelliculé	not available	34009 371 496 4 8	NOVARTIS PHARMA S.A.S.	FR
NISIS 160 mg, comprimé pelliculé	not available	34009 371 497 0 9	NOVARTIS PHARMA S.A.S.	FR
NISIS 160 mg, comprimé pelliculé	not available	34009 371 498 7 7	NOVARTIS PHARMA S.A.S.	FR
NISIS 80 mg, comprimé pelliculé	not available	34009 371 510 7 8	NOVARTIS PHARMA S.A.S.	FR
NISIS 80 mg, comprimé pelliculé	not available	34009 371 511 3 9	NOVARTIS PHARMA S.A.S.	FR

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NISIS 80 mg, comprimé pelliculé	not available	34009 371 513 6 8	NOVARTIS PHARMA S.A.S.	FR
NISIS 160 mg, comprimé pelliculé	not available	34009 379 070 6 4	NOVARTIS PHARMA S.A.S.	FR
NISIS 160 mg, comprimé pelliculé	not available	34009 379 071 2 5	NOVARTIS PHARMA S.A.S.	FR
NISIS 160 mg, comprimé pelliculé	not available	34009 379 072 9 3	NOVARTIS PHARMA S.A.S.	FR
NISIS 160 mg, comprimé pelliculé	not available	34009 379 073 5 4	NOVARTIS PHARMA S.A.S.	FR
NISIS 40 mg, comprimé pelliculé sécable	not available	34009 356 931 5 0	NOVARTIS PHARMA S.A.S.	FR
NISIS 40 mg, comprimé pelliculé sécable	not available	34009 356 932 1 1	NOVARTIS PHARMA S.A.S.	FR
NISIS 40 mg, comprimé pelliculé sécable	not available	34009 356 933 8 9	NOVARTIS PHARMA S.A.S.	FR
NISIS 40 mg, comprimé pelliculé sécable	not available	34009 356 934 4 0	NOVARTIS PHARMA S.A.S.	FR
NISIS 40 mg, comprimé pelliculé sécable	not available	34009 356 935 0 1	NOVARTIS PHARMA S.A.S.	FR
NISIS 40 mg, comprimé pelliculé sécable	not available	34009 356 936 7 9	NOVARTIS PHARMA S.A.S.	FR
NISIS 40 mg, comprimé pelliculé sécable	not available	34009 371 509 9 6	NOVARTIS PHARMA S.A.S.	FR
NISIS 40 mg, comprimé pelliculé sécable	not available	34009 372 292 3 4	NOVARTIS PHARMA S.A.S.	FR
NISIS 40 mg, comprimé pelliculé sécable	not available	34009 372 294 6 3	NOVARTIS PHARMA S.A.S.	FR
NISIS 40 mg, comprimé pelliculé sécable	not available	34009 372 295 2 4	NOVARTIS PHARMA S.A.S.	FR
NISIS 40 mg, comprimé pelliculé sécable	not available	34009 372 296 9 2	NOVARTIS PHARMA S.A.S.	FR
NISIS 40 mg, comprimé pelliculé sécable	not available	34009 373 890 1 3	NOVARTIS PHARMA S.A.S.	FR
NISIS 40 mg, comprimé pelliculé sécable	not available	34009 379 054 0 4	NOVARTIS PHARMA S.A.S.	FR
NISIS 40 mg, comprimé pelliculé sécable	not available	34009 379 055 7 2	NOVARTIS PHARMA S.A.S.	FR
NISIS 40 mg, comprimé pelliculé	not available	34009 379 056 3 3	NOVARTIS PHARMA S.A.S.	FR

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
sécable				
NISIS 40 mg, comprimé pelliculé sécable	not available	34009 379 058 6 2	NOVARTIS PHARMA S.A.S.	FR
NISIS 40 mg, comprimé pelliculé sécable	not available	34009 379 059 2 3	NOVARTIS PHARMA S.A.S.	FR
NISIS 40 mg, comprimé pelliculé sécable	not available	34009 379 060 0 5	NOVARTIS PHARMA S.A.S.	FR
Valsocard HCT 80 mg/12.5 mg filmdoublette	not available	OGYI-T-21875/03	ACTAVIS GROUP PTC EHF.	HU
Valsocard HCT 160 mg/12.5 mg filmdoublette	not available	OGYI-T-21875/12	ACTAVIS GROUP PTC EHF.	HU
Valsocard HCT 80 mg/12.5 mg filmdoublette	not available	OGYI-T-21875/02	ACTAVIS GROUP PTC EHF.	HU
Valsocard HCT 160 mg/12.5 mg filmdoublette	not available	OGYI-T-21875/10	ACTAVIS GROUP PTC EHF.	HU
Valsocard HCT 80 mg/12.5 mg filmdoublette	not available	OGYI-T-21875/06	ACTAVIS GROUP PTC EHF.	HU
Valsocard HCT 80 mg/12.5 mg filmdoublette	not available	OGYI-T-21875/05	ACTAVIS GROUP PTC EHF.	HU
Valsocard HCT 80 mg/12.5 mg filmdoublette	not available	OGYI-T-21875/04	ACTAVIS GROUP PTC EHF.	HU
Valsocard HCT 160 mg/12.5 mg filmdoublette	not available	OGYI-T-21875/11	ACTAVIS GROUP PTC EHF.	HU
Valsocard HCT 160 mg/25 mg filmdoublette	not available	OGYI-T-21875/16	ACTAVIS GROUP PTC EHF.	HU
Valsocard HCT 160 mg/25 mg filmdoublette	not available	OGYI-T-21875/17	ACTAVIS GROUP PTC EHF.	HU
Valsocard HCT 160 mg/25 mg filmdoublette	not available	OGYI-T-21875/20	ACTAVIS GROUP PTC EHF.	HU
Valsocard HCT 160 mg/25 mg filmdoublette	not available	OGYI-T-21875/21	ACTAVIS GROUP PTC EHF.	HU
Valsocard HCT 160 mg/25 mg filmdoublette	not available	OGYI-T-21875/18	ACTAVIS GROUP PTC EHF.	HU
Valsocard HCT 160 mg/25 mg	not available	OGYI-T-21875/19	ACTAVIS GROUP PTC EHF.	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
filmtabletta				
Valsocard HCT 160 mg/12.5 mg filmtabletta	not available	OGYI-T-21875/13	ACTAVIS GROUP PTC EHF.	HU
Valsocard HCT 80 mg/12.5 mg filmtabletta	not available	OGYI-T-21875/07	ACTAVIS GROUP PTC EHF.	HU
Valsocard HCT 160 mg/12.5 mg filmtabletta	not available	OGYI-T-21875/09	ACTAVIS GROUP PTC EHF.	HU
Valsocard HCT 160 mg/12.5 mg filmtabletta	not available	OGYI-T-21875/14	ACTAVIS GROUP PTC EHF.	HU
Valsocard HCT 160 mg/12.5 mg filmtabletta	not available	OGYI-T-21875/08	ACTAVIS GROUP PTC EHF.	HU
Valsocard HCT 160 mg/25 mg filmtabletta	not available	OGYI-T-21875/15	ACTAVIS GROUP PTC EHF.	HU
Valsocard HCT 80 mg/12.5 mg filmtabletta	not available	OGYI-T-21875/01	ACTAVIS GROUP PTC EHF.	HU
Co-Vals 320 mg/12,5 mg comprimidos recubiertos con película	not available	70.361	ESTEVE PHARMACEUTICALS, S.A.	ES
Co-Vals Forte 160 mg/25 mg comprimidos recubiertos con película	not available	66.676	ESTEVE PHARMACEUTICALS, S.A.	ES
Vals 320 mg comprimidos recubiertos con película.	not available	69.345	ESTEVE PHARMACEUTICALS, S.A.	ES
Valsartan-CT 120 mg Filmtabletten	SE/H/0748/003	70455.00.00	ABZ-PHARMA GMBH	DE
Valsartan Teva Pharma 120 mg filmdragerad tablett	SE/H/0748/003	26045	TEVA PHARMA B.V.	SE
Troval 80 mg Film-coated tablets	not available	ML 21333	DELORBIS PHARMACEUTICALS LTD	CY
Troval 160 mg Film-coated tablets	not available	ML 21334	DELORBIS PHARMACEUTICALS LTD	CY
Vals 80 mg comprimidos recubiertos con película.	not available	64.495	ESTEVE PHARMACEUTICALS, S.A.	ES
Diovan Comp 160 mg/12,5 mg filmdragerade tabletter	SE/H/0565/002	19525	NOVARTIS SVERIGE AB	SE
Diovan Comp 160 mg/25 mg filmdragerade tabletter	SE/H/0565/003	21204	NOVARTIS SVERIGE AB	SE

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
Diovan Comp 320 mg/12,5 mg filmdragerade tabletter	SE/H/0565/004	23641	NOVARTIS SVERIGE AB	SE
Diovan Comp 320 mg/25 mg filmdragerade tabletter	SE/H/0565/005	23642	NOVARTIS SVERIGE AB	SE
Diovan Comp 80 mg/12,5 mg filmdragerade tabletter	SE/H/0565/001	14098	NOVARTIS SVERIGE AB	SE
Valsartan comp. AbZ 120 mg/12,5 mg Filmtabletten	AT/H/0515/001	90714.00.00	ABZ-PHARMA GMBH	DE
Valsartan/HCT TEVA 120 mg/12,5 mg Filmtabletten	AT/H/0515/001	135568	TEVA B.V	AT
Valpression 40 mg compresse rivestite con film	not available	033119239	A. MENARINI - INDUSTRIE FARMACEUTICHE RIUNITE - S.R.L.	IT
Valpression 40 mg compresse rivestite con film	not available	033119203	A. MENARINI - INDUSTRIE FARMACEUTICHE RIUNITE - S.R.L.	IT
Valpression 40 mg compresse rivestite con film	not available	033119215	A. MENARINI - INDUSTRIE FARMACEUTICHE RIUNITE - S.R.L.	IT
Valpression 40 mg compresse rivestite con film	not available	033119254	A. MENARINI - INDUSTRIE FARMACEUTICHE RIUNITE - S.R.L.	IT
Valpression 320 mg compresse rivestite con film	not available	033119292	A. MENARINI - INDUSTRIE FARMACEUTICHE RIUNITE - S.R.L.	IT
Valpression 40 mg compresse rivestite con film	not available	033119266	A. MENARINI - INDUSTRIE FARMACEUTICHE RIUNITE - S.R.L.	IT
Valpression 40 mg compresse rivestite con film	not available	033119227	A. MENARINI - INDUSTRIE FARMACEUTICHE RIUNITE - S.R.L.	IT
Valpression 40 mg compresse rivestite con film	not available	033119241	A. MENARINI - INDUSTRIE FARMACEUTICHE RIUNITE - 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
Valpression 320 mg compresse rivestite con film	not available	033119280	A. MENARINI - INDUSTRIE FARMACEUTICHE RIUNITE - S.R.L.	IT
Valpression 320 mg compresse rivestite con film	not available	033119278	A. MENARINI - INDUSTRIE FARMACEUTICHE RIUNITE - S.R.L.	IT
Valpression 320 mg compresse rivestite con film	not available	033119330	A. MENARINI - INDUSTRIE FARMACEUTICHE RIUNITE - S.R.L.	IT
Valpression 320 mg compresse rivestite con film	not available	033119316	A. MENARINI - INDUSTRIE FARMACEUTICHE RIUNITE - S.R.L.	IT
Valpression 320 mg compresse rivestite con film	not available	033119304	A. MENARINI - INDUSTRIE FARMACEUTICHE RIUNITE - S.R.L.	IT
Valpression 40 mg compresse rivestite con film	not available	033119153	A. MENARINI - INDUSTRIE FARMACEUTICHE RIUNITE - S.R.L.	IT
Valpression 40 mg compresse rivestite con film	not available	033119177	A. MENARINI - INDUSTRIE FARMACEUTICHE RIUNITE - S.R.L.	IT
Valpression 320 mg compresse rivestite con film	not available	033119328	A. MENARINI - INDUSTRIE FARMACEUTICHE RIUNITE - S.R.L.	IT
Valpression 320 mg compresse rivestite con film	not available	033119342	A. MENARINI - INDUSTRIE FARMACEUTICHE RIUNITE - S.R.L.	IT
Valpression 40 mg compresse rivestite con film	not available	033119140	A. MENARINI - INDUSTRIE FARMACEUTICHE RIUNITE - S.R.L.	IT
Valpression 40 mg compresse rivestite con film	not available	033119138	A. MENARINI - INDUSTRIE FARMACEUTICHE RIUNITE - S.R.L.	IT
Valpression 40 mg compresse rivestite	not available	033119191	A. MENARINI - INDUSTRIE	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
con film			FARMACEUTICHE RIUNITE - S.R.L.	
Valpression 40 mg compresse rivestite con film	not available	033119165	A. MENARINI - INDUSTRIE FARMACEUTICHE RIUNITE - S.R.L.	IT
Valpression 40 mg compresse rivestite con film	not available	033119189	A. MENARINI - INDUSTRIE FARMACEUTICHE RIUNITE - S.R.L.	IT
Valpression 80 mg capsule	not available	033119013	A. MENARINI - INDUSTRIE FARMACEUTICHE RIUNITE - S.R.L.	IT
Valpression 160 mg capsule	not available	033119025	A. MENARINI - INDUSTRIE FARMACEUTICHE RIUNITE - S.R.L.	IT
Valpression 320 mg compresse rivestite con film	not available	033119355	A. MENARINI - INDUSTRIE FARMACEUTICHE RIUNITE - S.R.L.	IT
Valsartan 40 mg film-coated tablets	not available	PL 17907/0360	BRISTOL LABORATORIES LIMITED	XI
Valsartan 80 mg film-coated tablets	not available	PL 17907/0361	BRISTOL LABORATORIES LIMITED	XI
Valsartan 160 mg film-coated tablets	not available	PL 17907/0362	BRISTOL LABORATORIES LIMITED	XI
Valsartan 320 mg film-coated tablets	not available	PL 17907/0363	BRISTOL LABORATORIES LIMITED	XI
Angiosan 160 mg filmdragerade tabletter	SE/H/0407/002	19956	NOVARTIS SVERIGE AB	SE
Angiosan 320 mg	SE/H/0407/004	22841	NOVARTIS SVERIGE AB	SE
Angiosan 40 mg filmdragerade tabletter	SE/H/0407/003	20465	NOVARTIS SVERIGE AB	SE
Angiosan 80 mg filmdragerade tabletter	SE/H/0407/001	19955	NOVARTIS SVERIGE AB	SE
Valsartan comp. AbZ 120 mg/12,5 mg Filmtabletten	AT/H/0515/001	90714.00.00	ABZ-PHARMA GMBH	DE

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
Valsartan/HCT TEVA 120 mg/12,5 mg Filmtabletten	AT/H/0515/001	135568	TEVA B.V	AT
Co-Vals 160 mg/12,5 mg comprimidos recubiertos con película	not available	65.797	ESTEVE PHARMACEUTICALS, S.A.	ES
Co-Vals Forte 320 mg/25 mg comprimidos recubiertos con película	not available	70.363	ESTEVE PHARMACEUTICALS, S.A.	ES
Co-Vals 320 mg/12,5 mg comprimidos recubiertos con película	not available	70.361	ESTEVE PHARMACEUTICALS, S.A.	ES
Co-Vals Forte 160 mg/25 mg comprimidos recubiertos con película	not available	66.676	ESTEVE PHARMACEUTICALS, S.A.	ES
Vals 320 mg comprimidos recubiertos con película.	not available	69.345	ESTEVE PHARMACEUTICALS, S.A.	ES
AuroValsart HCT, 80 mg + 12,5 mg, tabletki powlekane	PT/H/1594/001/DC	23887	AUROVITAS PHARMA POLSKA SP. Z O.O	PL
AuroValsart HCT, 160 mg + 12,5 mg, tabletki powlekane	PT/H/1594/002/DC	23888	AUROVITAS PHARMA POLSKA SP. Z O.O	PL
AuroValsart HCT, 320 mg + 12,5 mg, tabletki powlekane	PT/H/1594/004/DC	23890	AUROVITAS PHARMA POLSKA SP. Z O.O	PL
Angiosan Comp 160 mg/12,5 mg filmdragerade tabletter	SE/H/0566/002	22458	NOVARTIS SVERIGE AB	SE
Angiosan Comp 160 mg/25 mg filmdragerade tabletter	SE/H/0566/003	22459	NOVARTIS SVERIGE AB	SE
Angiosan Comp 320 mg/12.5 mg film-coated tablets	SE/H/0566/004	23643	NOVARTIS SVERIGE AB	SE
Angiosan Comp 320 mg/25 mg filmdragerade tabletter	SE/H/0566/005	23644	NOVARTIS SVERIGE AB	SE
Angiosan Comp 80 mg /12,5 mg filmdragerade tabletter	SE/H/0566/001	22457	NOVARTIS SVERIGE AB	SE
Valsartan 40mg capsules	not available	PL 20416/503	CRESCENT PHARMA LIMITED	XI
Valsartan 80mg capsules	not available	PL 20416/504	CRESCENT PHARMA LIMITED	XI
Valsartan 160mg capsules	not available	PL 20416/505	CRESCENT PHARMA LIMITED	XI

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
Vals 160 mg comprimidos recubiertos con película.	not available	64.494	ESTEVE PHARMACEUTICALS, S.A.	ES
Co-Vals 320 mg/12,5 mg comprimidos recubiertos con película	not available	70.361	ESTEVE PHARMACEUTICALS, S.A.	ES
Co-Vals Forte 160 mg/25 mg comprimidos recubiertos con película	not available	66.676	ESTEVE PHARMACEUTICALS, S.A.	ES
Vals 320 mg comprimidos recubiertos con película.	not available	69.345	ESTEVE PHARMACEUTICALS, S.A.	ES
Valsocard 80 mg fimtableta	not available	OGYI-T-21874/12	ACTAVIS GROUP PTC EHF.	HU
Valsocard 80 mg fimtableta	not available	OGYI-T-21874/11	ACTAVIS GROUP PTC EHF.	HU
Valsocard 80 mg fimtableta	not available	OGYI-T-21874/10	ACTAVIS GROUP PTC EHF.	HU
Valsocard 80 mg fimtableta	not available	OGYI-T-21874/09	ACTAVIS GROUP PTC EHF.	HU
Valsocard 80 mg fimtableta	not available	OGYI-T-21874/14	ACTAVIS GROUP PTC EHF.	HU
Valsocard 80 mg fimtableta	not available	OGYI-T-21874/13	ACTAVIS GROUP PTC EHF.	HU
Valsocard 160 mg filmdragerade tabletter	not available	OGYI-T-21874/16	ACTAVIS GROUP PTC EHF.	HU
Valsocard 160 mg filmdragerade tabletter	not available	OGYI-T-21874/17	ACTAVIS GROUP PTC EHF.	HU
Valsocard 160 mg filmdragerade tabletter	not available	OGYI-T-21874/19	ACTAVIS GROUP PTC EHF.	HU
Valsocard 160 mg filmdragerade tabletter	not available	OGYI-T-21874/18	ACTAVIS GROUP PTC EHF.	HU
Valsocard 160 mg filmdragerade tabletter	not available	OGYI-T-21874/20	ACTAVIS GROUP PTC EHF.	HU
Valsocard 160 mg filmdragerade tabletter	not available	OGYI-T-21874/21	ACTAVIS GROUP PTC EHF.	HU
Valsocard 80 mg fimtableta	not available	OGYI-T-21874/08	ACTAVIS GROUP PTC EHF.	HU
Valsocard 160 mg fimtableta	not available	OGYI-T-21874/15	ACTAVIS GROUP PTC EHF.	HU
Valsartan//Hydroklortiazid Novartis 160 mg/25 mg filmdragerade tabletter	SE/H/0567/002	22461	NOVARTIS SVERIGE AB	SE
Valsartan/Hydroklortiazid Novartis 160 mg/12,5 mg filmdragerade tabletter	SE/H/0567/001	22460	NOVARTIS SVERIGE AB	SE
Valsartan/Hydroklortiazid Novartis 320 mg/12,5 mg filmdragerade tabletter	SE/H/0567/004	23645	NOVARTIS SVERIGE AB	SE
Valsartan/Hydroklortiazid Novartis 320 mg/25 mg filmdragerade tabletter	SE/H/0567/005	23646	NOVARTIS SVERIGE AB	SE
Valsartan/Hydroklortiazid Novartis 80 mg/12,5 mg filmdragerade tabletter	SE/H/0567/003	22462	NOVARTIS SVERIGE AB	SE
Corixil 80 mg/12,5 mg compresse	IT/H/0244/001	034774024	SANDOZ S.P.A.	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
rivestite con film				
Corixil 160 mg/12,5 mg compresse rivestite con film	IT/H/0244/002	034774036	SANDOZ S.P.A.	IT
Corixil 160 mg/12,5 mg compresse rivestite con film	IT/H/0244/002	034774048	SANDOZ S.P.A.	IT
Corixil 160 mg/12,5 mg compresse rivestite con film	IT/H/0244/002	034774075	SANDOZ S.P.A.	IT
Corixil 160 mg/12,5 mg compresse rivestite con film	IT/H/0244/002	034774099	SANDOZ S.P.A.	IT
Corixil 160 mg/12,5 mg compresse rivestite con film	IT/H/0244/002	034774087	SANDOZ S.P.A.	IT
Corixil 160 mg/25 mg compresse rivestite con film	IT/H/0244/003	034774125	SANDOZ S.P.A.	IT
Corixil 160 mg/25 mg compresse rivestite con film	IT/H/0244/003	034774137	SANDOZ S.P.A.	IT
Corixil 160 mg/25 mg compresse rivestite con film	IT/H/0244/003	034774149	SANDOZ S.P.A.	IT
Corixil 320 mg/12,5 mg compresse rivestite con film	IT/H/0244/004	034774202	SANDOZ S.P.A.	IT
Corixil 320 mg/12,5 mg compresse rivestite con film	IT/H/0244/004	034774214	SANDOZ S.P.A.	IT
Corixil 320 mg/12,5 mg compresse rivestite con film	IT/H/0244/004	034774238	SANDOZ S.P.A.	IT
Corixil 320 mg/12,5 mg compresse rivestite con film	IT/H/0244/004	034774341	SANDOZ S.P.A.	IT
Corixil 80 mg/12,5 mg compresse rivestite con film	IT/H/0244/001	034774012	SANDOZ S.P.A.	IT
Corixil 160 mg/12,5 mg compresse rivestite con film	IT/H/0244/002	034774051	SANDOZ S.P.A.	IT
Corixil 160 mg/25 mg compresse rivestite con film	IT/H/0244/003	034774101	SANDOZ S.P.A.	IT
Corixil 160 mg/12,5 mg compresse rivestite con film	IT/H/0244/002	034774063	SANDOZ S.P.A.	IT
Corixil 160 mg/25 mg compresse	IT/H/0244/003	034774113	SANDOZ S.P.A.	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
rivestite con film				
Corixil 160 mg/25 mg compresse rivestite con film	IT/H/0244/003	034774152	SANDOZ S.P.A.	IT
Corixil 160 mg/25 mg compresse rivestite con film	IT/H/0244/003	034774164	SANDOZ S.P.A.	IT
Corixil 320 mg/12,5 mg compresse rivestite con film	IT/H/0244/004	034774176	SANDOZ S.P.A.	IT
Corixil 320 mg/12,5 mg compresse rivestite con film	IT/H/0244/004	034774190	SANDOZ S.P.A.	IT
Corixil 320 mg/12,5 mg compresse rivestite con film	IT/H/0244/004	034774226	SANDOZ S.P.A.	IT
Corixil 320 mg/12,5 mg compresse rivestite con film	IT/H/0244/004	034774188	SANDOZ S.P.A.	IT
Corixil 320 mg/12,5 mg compresse rivestite con film	IT/H/0244/004	034774240	SANDOZ S.P.A.	IT
Corixil 320 mg/25 mg compresse rivestite con film	IT/H/0244/005	034774265	SANDOZ S.P.A.	IT
Corixil 320 mg/25 mg compresse rivestite con film	IT/H/0244/005	034774253	SANDOZ S.P.A.	IT
Corixil 320 mg/25 mg compresse rivestite con film	IT/H/0244/005	034774277	SANDOZ S.P.A.	IT
Corixil 320 mg/25 mg compresse rivestite con film	IT/H/0244/005	034774289	SANDOZ S.P.A.	IT
Corixil 320 mg/25 mg compresse rivestite con film	IT/H/0244/005	034774303	SANDOZ S.P.A.	IT
Corixil 320 mg/25 mg compresse rivestite con film	IT/H/0244/005	034774315	SANDOZ S.P.A.	IT
Corixil 320 mg/25 mg compresse rivestite con film	IT/H/0244/005	034774327	SANDOZ S.P.A.	IT
Corixil 320 mg/25 mg compresse rivestite con film	IT/H/0244/005	034774339	SANDOZ S.P.A.	IT
Corixil 320 mg/25 mg compresse rivestite con film	IT/H/0244/005	034774354	SANDOZ S.P.A.	IT
COMBISARTAN 160 mg/12,5 mg	not available	034134078	A. MENARINI - INDUSTRIE	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
compresse rivestite con film			FARMACEUTICHE RIUNITE - S.R.L.	
COMBISARTAN 160 mg/25 mg compresse rivestite con film	not available	034134155	A. MENARINI - INDUSTRIE FARMACEUTICHE RIUNITE - S.R.L.	IT
COMBISARTAN 160 mg/12,5 mg compresse rivestite con film	not available	034134054	A. MENARINI - INDUSTRIE FARMACEUTICHE RIUNITE - S.R.L.	IT
COMBISARTAN 160 mg/12,5 mg compresse rivestite con film	not available	034134039	A. MENARINI - INDUSTRIE FARMACEUTICHE RIUNITE - S.R.L.	IT
COMBISARTAN 160 mg/12,5 mg compresse rivestite con film	not available	034134092	A. MENARINI - INDUSTRIE FARMACEUTICHE RIUNITE - S.R.L.	IT
COMBISARTAN 160 mg/12,5 mg compresse rivestite con film	not available	034134080	A. MENARINI - INDUSTRIE FARMACEUTICHE RIUNITE - S.R.L.	IT
COMBISARTAN 160 mg/12,5 mg compresse rivestite con film	not available	034134041	A. MENARINI - INDUSTRIE FARMACEUTICHE RIUNITE - S.R.L.	IT
COMBISARTAN 160 mg/25 mg compresse rivestite con film	not available	034134167	A. MENARINI - INDUSTRIE FARMACEUTICHE RIUNITE - S.R.L.	IT
COMBISARTAN 160 mg/25 mg compresse rivestite con film	not available	034134116	A. MENARINI - INDUSTRIE FARMACEUTICHE RIUNITE - S.R.L.	IT
COMBISARTAN 160 mg/25 mg compresse rivestite con film	not available	034134104	A. MENARINI - INDUSTRIE FARMACEUTICHE RIUNITE - S.R.L.	IT
COMBISARTAN 160 mg/25 mg compresse rivestite con film	not available	034134128	A. MENARINI - INDUSTRIE FARMACEUTICHE RIUNITE - S.R.L.	IT
COMBISARTAN 80 mg/12,5 mg compresse rivestite con film	not available	034134015	A. MENARINI - INDUSTRIE FARMACEUTICHE RIUNITE	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
			- S.R.L.	
COMBISARTAN 160 mg/25 mg compresse rivestite con film	not available	034134142	A. MENARINI - INDUSTRIE FARMACEUTICHE RIUNITE - S.R.L.	IT
COMBISARTAN 80 mg/12,5 mg compresse rivestite con film	not available	034134027	A. MENARINI - INDUSTRIE FARMACEUTICHE RIUNITE - S.R.L.	IT
COMBISARTAN 160 mg/12,5 mg compresse rivestite con film	not available	034134066	A. MENARINI - INDUSTRIE FARMACEUTICHE RIUNITE - S.R.L.	IT
COMBISARTAN 160 mg/25 mg compresse rivestite con film	not available	034134130	A. MENARINI - INDUSTRIE FARMACEUTICHE RIUNITE - S.R.L.	IT
COMBISARTAN 320 mg/12,5 mg compresse rivestite con film	not available	034134332	A. MENARINI - INDUSTRIE FARMACEUTICHE RIUNITE - S.R.L.	IT
COMBISARTAN 320 mg/12,5 mg compresse rivestite con film	not available	034134229	A. MENARINI - INDUSTRIE FARMACEUTICHE RIUNITE - S.R.L.	IT
COMBISARTAN 320 mg/12,5 mg compresse rivestite con film	not available	034134205	A. MENARINI - INDUSTRIE FARMACEUTICHE RIUNITE - S.R.L.	IT
COMBISARTAN 320 mg/12,5 mg compresse rivestite con film	not available	034134217	A. MENARINI - INDUSTRIE FARMACEUTICHE RIUNITE - S.R.L.	IT
COMBISARTAN 320 mg/12,5 mg compresse rivestite con film	not available	034134179	A. MENARINI - INDUSTRIE FARMACEUTICHE RIUNITE - S.R.L.	IT
COMBISARTAN 320 mg/12,5 mg compresse rivestite con film	not available	034134243	A. MENARINI - INDUSTRIE FARMACEUTICHE RIUNITE - S.R.L.	IT
COMBISARTAN 320 mg/12,5 mg compresse rivestite con film	not available	034134181	A. MENARINI - INDUSTRIE FARMACEUTICHE RIUNITE - 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
COMBISARTAN 320 mg/12,5 mg compresse rivestite con film	not available	034134231	A. MENARINI - INDUSTRIE FARMACEUTICHE RIUNITE - S.R.L.	IT
COMBISARTAN 320 mg/12,5 mg compresse rivestite con film	not available	034134193	A. MENARINI - INDUSTRIE FARMACEUTICHE RIUNITE - S.R.L.	IT
COMBISARTAN 320 mg/25 mg compresse rivestite con film	not available	034134268	A. MENARINI - INDUSTRIE FARMACEUTICHE RIUNITE - S.R.L.	IT
COMBISARTAN 320 mg/25 mg compresse rivestite con film	not available	034134256	A. MENARINI - INDUSTRIE FARMACEUTICHE RIUNITE - S.R.L.	IT
COMBISARTAN 320 mg/25 mg compresse rivestite con film	not available	034134294	A. MENARINI - INDUSTRIE FARMACEUTICHE RIUNITE - S.R.L.	IT
COMBISARTAN 320 mg/25 mg compresse rivestite con film	not available	034134320	A. MENARINI - INDUSTRIE FARMACEUTICHE RIUNITE - S.R.L.	IT
COMBISARTAN 320 mg/25 mg compresse rivestite con film	not available	034134344	A. MENARINI - INDUSTRIE FARMACEUTICHE RIUNITE - S.R.L.	IT
COMBISARTAN 320 mg/25 mg compresse rivestite con film	not available	034134306	A. MENARINI - INDUSTRIE FARMACEUTICHE RIUNITE - S.R.L.	IT
COMBISARTAN 320 mg/25 mg compresse rivestite con film	not available	034134270	A. MENARINI - INDUSTRIE FARMACEUTICHE RIUNITE - S.R.L.	IT
COMBISARTAN 320 mg/25 mg compresse rivestite con film	not available	034134318	A. MENARINI - INDUSTRIE FARMACEUTICHE RIUNITE - S.R.L.	IT
COMBISARTAN 320 mg/25 mg compresse rivestite con film	not available	034134282	A. MENARINI - INDUSTRIE FARMACEUTICHE RIUNITE - S.R.L.	IT
Valsartan Novartis 320 mg	SE/H/0408/004	22842	NOVARTIS SVERIGE AB	SE

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
Valsartan Novartis 40 mg filmdragerad tablett	SE/H/0408/003	20466	NOVARTIS SVERIGE AB	SE
Valsartan Novartis 80 mg filmdragerade tabletter	SE/H/0408/001	19957	NOVARTIS SVERIGE AB	SE
Valsartan-Novartis 160 mg filmdragerade tabletter	SE/H/0408/002	19958	NOVARTIS SVERIGE AB	SE