



EUROPEAN MEDICINES AGENCY
SCIENCE MEDICINES HEALTH

11 January 2018
EMA/34118/2018
Human Medicines Evaluation Division

List of nationally authorised medicinal products

Active substance(s): valsartan, hydrochlorothiazide/valsartan

Procedure No.: PSUSA/00010396/201704



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
Angiosan 160 mg filmdragerade tabletter	SE/H/0407/002	19956	NOVARTIS SVERIGE AB	SE
Angiosan 160 mg filmdragerade tabletter	SE/H/0407/002	19956	NOVARTIS SVERIGE AB	SE
Angiosan 160 mg filmdragerade tabletter	SE/H/0407/002	19956	NOVARTIS SVERIGE AB	SE
Angiosan 160 mg filmdragerade tabletter	SE/H/0407/002	19956	NOVARTIS SVERIGE AB	SE
Angiosan 160 mg filmdragerade tabletter	SE/H/0407/002	19956	NOVARTIS SVERIGE AB	SE
Angiosan 160 mg filmdragerade tabletter	SE/H/0407/002	19956	NOVARTIS SVERIGE AB	SE
Angiosan 3 mg/ml oral lösning	SE/H/0407/005	44277	NOVARTIS SVERIGE AB	SE
Angiosan 3 mg/ml oral lösning	SE/H/0407/005	44277	NOVARTIS SVERIGE AB	SE
Angiosan 3 mg/ml oral lösning	SE/H/0407/005	44277	NOVARTIS SVERIGE AB	SE
Angiosan 3 mg/ml oral lösning	SE/H/0407/005	44277	NOVARTIS SVERIGE AB	SE
Angiosan 3 mg/ml oral lösning	SE/H/0407/005	44277	NOVARTIS SVERIGE AB	SE
Angiosan 3 mg/ml oral lösning	SE/H/0407/005	44277	NOVARTIS SVERIGE AB	SE
Angiosan 320 mg	SE/H/0407/004	22841	NOVARTIS SVERIGE AB	SE
Angiosan 320 mg	SE/H/0407/004	22841	NOVARTIS SVERIGE AB	SE
Angiosan 320 mg	SE/H/0407/004	22841	NOVARTIS SVERIGE AB	SE
Angiosan 320 mg	SE/H/0407/004	22841	NOVARTIS SVERIGE AB	SE
Angiosan 320 mg	SE/H/0407/004	22841	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 40 mg filmdragerade tabletter	SE/H/0407/003	20465	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
Angiosan 40 mg filmdragerade tabletter	SE/H/0407/003	20465	NOVARTIS SVERIGE AB	SE
Angiosan 40 mg filmdragerade tabletter	SE/H/0407/003	20465	NOVARTIS SVERIGE AB	SE
Angiosan 40 mg filmdragerade tabletter	SE/H/0407/003	20465	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
Angiosan 80 mg filmdragerade tabletter	SE/H/0407/001	19955	NOVARTIS SVERIGE AB	SE
Angiosan 80 mg filmdragerade tabletter	SE/H/0407/001	19955	NOVARTIS SVERIGE AB	SE
Angiosan 80 mg filmdragerade tabletter	SE/H/0407/001	19955	NOVARTIS SVERIGE AB	SE
Angiosan 80 mg filmdragerade tabletter	SE/H/0407/001	19955	NOVARTIS SVERIGE AB	SE
Angiosan 80 mg filmdragerade tabletter	SE/H/0407/001	19955	NOVARTIS SVERIGE AB	SE
Angiosan Comp 160 mg/12,5 mg filmdragerade tabletter	SE/H/0566/002	22458	NOVARTIS SVERIGE AB	SE
Angiosan Comp 160 mg/12,5 mg filmdragerade tabletter	SE/H/0566/002	22458	NOVARTIS SVERIGE AB	SE
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 160	SE/H/0566/003	22459	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
mg/25 mg filmdragerade tabletter				
Angiosan Comp 160 mg/25 mg filmdragerade tabletter	SE/H/0566/003	22459	NOVARTIS SVERIGE AB	SE
Angiosan Comp 320 mg/12,5 mg filmdragerade tabletter	SE/H/0566/004	23643	NOVARTIS SVERIGE AB	SE
Angiosan Comp 320 mg/12,5 mg filmdragerade tabletter	SE/H/0566/004	23643	NOVARTIS SVERIGE AB	SE
Angiosan Comp 320 mg/12,5 mg filmdragerade tabletter	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 320 mg/25 mg filmdragerade tabletter	SE/H/0566/005	23644	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
Angiosan Comp 80 mg /12,5 mg filmdragerade tabletter	SE/H/0566/001	22457	NOVARTIS SVERIGE AB	SE
Angiosan Comp 80 mg /12,5 mg filmdragerade tabletter	SE/H/0566/001	22457	NOVARTIS SVERIGE AB	SE
Awalten, 80 mg + 12,5	not available	22739	S-LAB SP.ZO.O.	PL

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
mg, tabletki powlekane				
CO-DALZAD (160+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/002	247200201	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (160+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/002	247200202	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (160+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/002	247200203	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (160+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/002	247200204	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (160+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/002	247200205	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (160+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/002	247200206	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (160+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/002	247200207	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (160+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/002	247200208	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (160+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/002	247200209	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (160+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/002	247200201	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (160+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ	SE/H/0566/002	247200202	NOVARTIS (HELLAS) S.A.C.I.	GR

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
ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ				
CO-DALZAD (160+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/002	247200203	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (160+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/002	247200204	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (160+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/002	247200205	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (160+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/002	247200206	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (160+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/002	247200207	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (160+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/002	247200208	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (160+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/002	247200209	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (160+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/002	247200201	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (160+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/002	247200202	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (160+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/002	247200203	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (160+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ	SE/H/0566/002	247200204	NOVARTIS (HELLAS) S.A.C.I.	GR

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
ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ				
CO-DALZAD (160+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/002	247200205	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (160+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/002	247200206	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (160+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/002	247200207	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (160+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/002	247200208	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (160+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/002	247200209	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (160+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/003	247200301	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (160+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/003	247200302	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (160+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/003	247200303	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (160+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/003	247200304	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (160+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/003	247200305	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (160+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ	SE/H/0566/003	247200306	NOVARTIS (HELLAS) S.A.C.I.	GR

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
ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ				
CO-DALZAD (160+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/003	247200307	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (160+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/003	247200308	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (160+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/003	247200309	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (160+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/003	247200310	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (160+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/003	247200301	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (160+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/003	247200302	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (160+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/003	247200303	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (160+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/003	247200304	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (160+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/003	247200305	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (160+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/003	247200306	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (160+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ	SE/H/0566/003	247200307	NOVARTIS (HELLAS) S.A.C.I.	GR

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
ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ				
CO-DALZAD (160+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/003	247200308	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (160+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/003	247200309	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (160+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/003	247200310	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (160+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/003	247200301	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (160+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/003	247200302	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (160+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/003	247200303	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (160+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/003	247200304	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (160+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/003	247200305	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (160+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/003	247200306	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (160+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/003	247200307	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (160+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ	SE/H/0566/003	247200308	NOVARTIS (HELLAS) S.A.C.I.	GR

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
ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ				
CO-DALZAD (160+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/003	247200309	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (160+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/003	247200310	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (320+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/004	247200401	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (320+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/004	247200402	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (320+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/004	247200403	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (320+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/004	247200404	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (320+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/004	247200405	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (320+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/004	247200406	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (320+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/004	247200407	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (320+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/004	247200408	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (320+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ	SE/H/0566/004	247200409	NOVARTIS (HELLAS) S.A.C.I.	GR

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
ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ				
CO-DALZAD (320+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/004	247200401	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (320+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/004	247200402	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (320+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/004	247200403	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (320+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/004	247200404	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (320+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/004	247200405	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (320+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/004	247200406	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (320+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/004	247200407	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (320+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/004	247200408	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (320+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/004	247200409	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (320+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/004	247200401	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (320+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ	SE/H/0566/004	247200402	NOVARTIS (HELLAS) S.A.C.I.	GR

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
ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ				
CO-DALZAD (320+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/004	247200403	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (320+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/004	247200404	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (320+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/004	247200405	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (320+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/004	247200406	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (320+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/004	247200407	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (320+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/004	247200408	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (320+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/004	247200409	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (320+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/005	247200501	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (320+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/005	247200502	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (320+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/005	247200503	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (320+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ	SE/H/0566/005	247200504	NOVARTIS (HELLAS) S.A.C.I.	GR

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
ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ				
CO-DALZAD (320+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/005	247200505	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (320+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/005	247200506	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (320+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/005	247200507	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (320+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/005	247200508	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (320+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/005	247200509	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (320+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/005	247200501	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (320+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/005	247200502	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (320+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/005	247200503	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (320+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/005	247200504	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (320+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/005	247200505	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (320+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ	SE/H/0566/005	247200506	NOVARTIS (HELLAS) S.A.C.I.	GR

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
ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ				
CO-DALZAD (320+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/005	247200507	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (320+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/005	247200508	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (320+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/005	247200509	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (320+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/005	247200501	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (320+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/005	247200502	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (320+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/005	247200503	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (320+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/005	247200504	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (320+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/005	247200505	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (320+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/005	247200506	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (320+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/005	247200507	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (320+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ	SE/H/0566/005	247200508	NOVARTIS (HELLAS) S.A.C.I.	GR

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
ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ				
CO-DALZAD (320+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/005	247200509	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (80+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/001	247200102	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (80+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/001	247200103	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (80+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/001	247200104	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (80+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/001	247200105	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (80+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/001	247200106	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (80+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/001	247200107	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (80+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/001	247200108	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (80+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/001	247200109	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (80+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/001	247200102	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (80+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ	SE/H/0566/001	247200103	NOVARTIS (HELLAS) S.A.C.I.	GR

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
ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ				
CO-DALZAD (80+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/001	247200104	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (80+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/001	247200105	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (80+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/001	247200106	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (80+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/001	247200107	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (80+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/001	247200108	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (80+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/001	247200109	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (80+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/001	247200102	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (80+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/001	247200103	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (80+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/001	247200104	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (80+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/001	247200105	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (80+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ	SE/H/0566/001	247200106	NOVARTIS (HELLAS) S.A.C.I.	GR

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
ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ				
CO-DALZAD (80+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/001	247200107	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (80+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/001	247200108	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DALZAD (80+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0566/001	247200109	NOVARTIS (HELLAS) S.A.C.I.	GR
Co-Dalzad (80+12,5) mg επικαλυμμένα με λεπτό υμένιο δισκία	SE/H/0566/001	247200101	NOVARTIS (HELLAS) S.A.C.I.	GR
Co-Dalzad (80+12,5) mg επικαλυμμένα με λεπτό υμένιο δισκία	SE/H/0566/001	247200101	NOVARTIS (HELLAS) S.A.C.I.	GR
Co-Dalzad (80+12,5) mg επικαλυμμένα με λεπτό υμένιο δισκία	SE/H/0566/001	247200101	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (160+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/002	238890201	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (160+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/002	238890202	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (160+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/002	238890203	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (160+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/002	238890204	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (160+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ	SE/H/0565/002	238890205	NOVARTIS (HELLAS) S.A.C.I.	GR

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
ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ				
CO-DIOVAN (160+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/002	238890206	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (160+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/002	238890207	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (160+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/002	238890201	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (160+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/002	238890202	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (160+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/002	238890203	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (160+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/002	238890204	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (160+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/002	238890205	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (160+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/002	238890206	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (160+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/002	238890207	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (160+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/002	238890201	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (160+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ	SE/H/0565/002	238890202	NOVARTIS (HELLAS) S.A.C.I.	GR

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
ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ				
CO-DIOVAN (160+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/002	238890203	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (160+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/002	238890204	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (160+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/002	238890205	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (160+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/002	238890206	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (160+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/002	238890207	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (160+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/002	238890201	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (160+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/002	238890202	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (160+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/002	238890203	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (160+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/002	238890204	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (160+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/002	238890205	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (160+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ	SE/H/0565/002	238890206	NOVARTIS (HELLAS) S.A.C.I.	GR

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
ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ				
CO-DIOVAN (160+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/002	238890207	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (160+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/003	238890301	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (160+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/003	238890302	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (160+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/003	238890303	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (160+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/003	238890304	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (160+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/003	238890305	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (160+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/003	238890306	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (160+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/003	238890307	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (160+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/003	238890308	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (160+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/003	238890309	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (160+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ	SE/H/0565/003	238890310	NOVARTIS (HELLAS) S.A.C.I.	GR

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
ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ				
CO-DIOVAN (160+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/003	238890301	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (160+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/003	238890302	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (160+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/003	238890303	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (160+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/003	238890304	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (160+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/003	238890305	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (160+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/003	238890306	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (160+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/003	238890307	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (160+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/003	238890308	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (160+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/003	238890309	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (160+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/003	238890310	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (160+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ	SE/H/0565/003	238890301	NOVARTIS (HELLAS) S.A.C.I.	GR

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
ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ				
CO-DIOVAN (160+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/003	238890302	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (160+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/003	238890303	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (160+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/003	238890304	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (160+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/003	238890305	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (160+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/003	238890306	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (160+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/003	238890307	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (160+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/003	238890308	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (160+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/003	238890309	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (160+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/003	238890310	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (160+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/003	238890301	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (160+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ	SE/H/0565/003	238890302	NOVARTIS (HELLAS) S.A.C.I.	GR

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
ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ				
CO-DIOVAN (160+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/003	238890303	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (160+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/003	238890304	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (160+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/003	238890305	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (160+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/003	238890306	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (160+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/003	238890307	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (160+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/003	238890308	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (160+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/003	238890309	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (160+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/003	238890310	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (320+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/004	238890401	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (320+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/004	238890402	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (320+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ	SE/H/0565/004	238890403	NOVARTIS (HELLAS) S.A.C.I.	GR

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
ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ				
CO-DIOVAN (320+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/004	238890404	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (320+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/004	238890405	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (320+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/004	238890406	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (320+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/004	238890407	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (320+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/004	238890408	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (320+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/004	238890409	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (320+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/004	238890401	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (320+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/004	238890402	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (320+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/004	238890403	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (320+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/004	238890404	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (320+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ	SE/H/0565/004	238890405	NOVARTIS (HELLAS) S.A.C.I.	GR

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
ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ				
CO-DIOVAN (320+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/004	238890406	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (320+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/004	238890407	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (320+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/004	238890408	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (320+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/004	238890409	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (320+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/004	238890401	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (320+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/004	238890402	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (320+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/004	238890403	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (320+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/004	238890404	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (320+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/004	238890405	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (320+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/004	238890406	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (320+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ	SE/H/0565/004	238890407	NOVARTIS (HELLAS) S.A.C.I.	GR

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
ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ				
CO-DIOVAN (320+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/004	238890408	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (320+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/004	238890409	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (320+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/004	238890401	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (320+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/004	238890402	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (320+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/004	238890403	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (320+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/004	238890404	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (320+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/004	238890405	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (320+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/004	238890406	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (320+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/004	238890407	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (320+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/004	238890408	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (320+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ	SE/H/0565/004	238890409	NOVARTIS (HELLAS) S.A.C.I.	GR

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
ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ				
CO-DIOVAN (320+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/005	238890501	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (320+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/005	238890502	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (320+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/005	238890503	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (320+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/005	238890504	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (320+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/005	238890505	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (320+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/005	238890506	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (320+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/005	238890507	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (320+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/005	238890508	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (320+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/005	238890509	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (320+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/005	238890501	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (320+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ	SE/H/0565/005	238890502	NOVARTIS (HELLAS) S.A.C.I.	GR

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
ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ				
CO-DIOVAN (320+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/005	238890503	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (320+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/005	238890504	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (320+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/005	238890505	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (320+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/005	238890506	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (320+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/005	238890507	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (320+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/005	238890508	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (320+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/005	238890509	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (320+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/005	238890501	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (320+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/005	238890502	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (320+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/005	238890503	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (320+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ	SE/H/0565/005	238890504	NOVARTIS (HELLAS) S.A.C.I.	GR

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
ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ				
CO-DIOVAN (320+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/005	238890505	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (320+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/005	238890506	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (320+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/005	238890507	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (320+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/005	238890508	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (320+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/005	238890509	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (320+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/005	238890501	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (320+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/005	238890502	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (320+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/005	238890503	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (320+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/005	238890504	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (320+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/005	238890505	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (320+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ	SE/H/0565/005	238890506	NOVARTIS (HELLAS) S.A.C.I.	GR

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
ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ				
CO-DIOVAN (320+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/005	238890507	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (320+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/005	238890508	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (320+25) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/005	238890509	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (80+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/001	238890101	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (80+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/001	238890102	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (80+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/001	238890103	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (80+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/001	238890104	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (80+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/001	238890105	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (80+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/001	238890106	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (80+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/001	238890107	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (80+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ	SE/H/0565/001	238890108	NOVARTIS (HELLAS) S.A.C.I.	GR

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
ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ				
CO-DIOVAN (80+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/001	238890101	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (80+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/001	238890102	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (80+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/001	238890103	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (80+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/001	238890104	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (80+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/001	238890105	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (80+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/001	238890106	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (80+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/001	238890107	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (80+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/001	238890108	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (80+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/001	238890101	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (80+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/001	238890102	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (80+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ	SE/H/0565/001	238890103	NOVARTIS (HELLAS) S.A.C.I.	GR

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
ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ				
CO-DIOVAN (80+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/001	238890104	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (80+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/001	238890105	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (80+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/001	238890106	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (80+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/001	238890107	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (80+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/001	238890108	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (80+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/001	238890101	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (80+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/001	238890102	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (80+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/001	238890103	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (80+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/001	238890104	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (80+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/001	238890105	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (80+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ	SE/H/0565/001	238890106	NOVARTIS (HELLAS) S.A.C.I.	GR

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
ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ				
CO-DIOVAN (80+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/001	238890107	NOVARTIS (HELLAS) S.A.C.I.	GR
CO-DIOVAN (80+12,5) MG ΕΠΗΘΑΙΠΚΚ'ΕΛΑ ΚΕ ΙΕΠΗΦ ΠΚ'ΕΛΗΝ ΔΗΖΘ'ΙΑ	SE/H/0565/001	238890108	NOVARTIS (HELLAS) S.A.C.I.	GR
Co-Diovan 160 mg + 25 mg tabletki powlekane	SE/H/0565/003	10629	NOVARTIS POLAND SP. Z O. O.	PL
Co-Diovan 160 mg + 25 mg tabletki powlekane	SE/H/0565/003	10629	NOVARTIS POLAND SP. Z O. O.	PL
Co-Diovan 160 mg + 25 mg tabletki powlekane	SE/H/0565/003	10629	NOVARTIS POLAND SP. Z O. O.	PL
Co-Diovan 160 mg + 25 mg tabletki powlekane	SE/H/0565/003	10629	NOVARTIS POLAND SP. Z O. O.	PL
CO-DIOVAN 160 MG/12,5 MG APVALKOTĀS TABLETES	SE/H/0565/002	02-0393	NOVARTIS FINLAND OY	LV
CO-DIOVAN 160 MG/12,5 MG APVALKOTĀS TABLETES	SE/H/0565/002	02-0393	NOVARTIS FINLAND OY	LV
CO-DIOVAN 160 MG/12,5 MG APVALKOTĀS TABLETES	SE/H/0565/002	02-0393	NOVARTIS FINLAND OY	LV
CO-DIOVAN 160 MG/12,5 MG APVALKOTĀS TABLETES	SE/H/0565/002	02-0393	NOVARTIS FINLAND OY	LV
Co-Diován 160 mg/12,5 mg comprimidos recubiertos con película	SE/H/0565/002	65.794	NOVARTIS FARMACÉUTICA S.A.	ES
Co-Diován 160 mg/12,5 mg comprimidos recubiertos con película	SE/H/0565/002	65.794	NOVARTIS FARMACÉUTICA S.A.	ES

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
Co-Diován 160 mg/12,5 mg comprimidos recubiertos con película	SE/H/0565/002	65.794	NOVARTIS FARMACÉUTICA S.A.	ES
Co-Diován 160 mg/12,5 mg comprimidos recubiertos con película	SE/H/0565/002	65.794	NOVARTIS FARMACÉUTICA S.A.	ES
Co-Diovan 160 mg/12,5 mg comprimidos revestidos por película	SE/H/0565/002	4804688	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 160 mg/12,5 mg comprimidos revestidos por película	SE/H/0565/002	4804787	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 160 mg/12,5 mg comprimidos revestidos por película	SE/H/0565/002	4804886	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 160 mg/12,5 mg comprimidos revestidos por película	SE/H/0565/002	4804985	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 160 mg/12,5 mg comprimidos revestidos por película	SE/H/0565/002	4805081	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 160 mg/12,5 mg comprimidos revestidos por película	SE/H/0565/002	4805180	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 160 mg/12,5 mg comprimidos revestidos por película	SE/H/0565/002	4805289	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 160 mg/12,5 mg comprimidos revestidos por película	SE/H/0565/002	5022769	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 160 mg/12,5 mg comprimidos revestidos por película	SE/H/0565/002	5022777	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT

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
Co-Diovan 160 mg/12,5 mg comprimidos revestidos por película	SE/H/0565/002	4804688	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 160 mg/12,5 mg comprimidos revestidos por película	SE/H/0565/002	4804787	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 160 mg/12,5 mg comprimidos revestidos por película	SE/H/0565/002	4804886	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 160 mg/12,5 mg comprimidos revestidos por película	SE/H/0565/002	4804985	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 160 mg/12,5 mg comprimidos revestidos por película	SE/H/0565/002	4805081	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 160 mg/12,5 mg comprimidos revestidos por película	SE/H/0565/002	4805180	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 160 mg/12,5 mg comprimidos revestidos por película	SE/H/0565/002	4805289	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 160 mg/12,5 mg comprimidos revestidos por película	SE/H/0565/002	5022769	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 160 mg/12,5 mg comprimidos revestidos por película	SE/H/0565/002	5022777	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 160 mg/12,5 mg comprimidos revestidos por película	SE/H/0565/002	4804688	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 160 mg/12,5 mg comprimidos revestidos por película	SE/H/0565/002	4804787	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT

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
Co-Diovan 160 mg/12,5 mg comprimidos revestidos por película	SE/H/0565/002	4804886	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 160 mg/12,5 mg comprimidos revestidos por película	SE/H/0565/002	4804985	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 160 mg/12,5 mg comprimidos revestidos por película	SE/H/0565/002	4805081	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 160 mg/12,5 mg comprimidos revestidos por película	SE/H/0565/002	4805180	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 160 mg/12,5 mg comprimidos revestidos por película	SE/H/0565/002	4805289	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 160 mg/12,5 mg comprimidos revestidos por película	SE/H/0565/002	5022769	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 160 mg/12,5 mg comprimidos revestidos por película	SE/H/0565/002	5022777	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 160 mg/12,5 mg comprimidos revestidos por película	SE/H/0565/002	4804688	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 160 mg/12,5 mg comprimidos revestidos por película	SE/H/0565/002	4804787	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 160 mg/12,5 mg comprimidos revestidos por película	SE/H/0565/002	4804886	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 160 mg/12,5 mg comprimidos revestidos por película	SE/H/0565/002	4804985	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT

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
Co-Diovan 160 mg/12,5 mg comprimidos revestidos por película	SE/H/0565/002	4805081	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 160 mg/12,5 mg comprimidos revestidos por película	SE/H/0565/002	4805180	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 160 mg/12,5 mg comprimidos revestidos por película	SE/H/0565/002	4805289	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 160 mg/12,5 mg comprimidos revestidos por película	SE/H/0565/002	5022769	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 160 mg/12,5 mg comprimidos revestidos por película	SE/H/0565/002	5022777	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
CO-DIOVAN 160 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/002	H/03/00406/019	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/002	H/03/00406/020	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/002	H/03/00406/021	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/002	H/03/00406/022	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/002	H/03/00406/023	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/002	H/03/00406/024	NOVARTIS PHARMA GMBH	SI

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
CO-DIOVAN 160 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/002	H/03/00406/025	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/002	H/03/00406/026	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/002	H/03/00406/027	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/002	H/03/00406/028	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/002	H/03/00406/029	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/002	H/03/00406/030	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/002	H/03/00406/031	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/002	H/03/00406/032	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/002	H/03/00406/033	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/002	H/03/00406/034	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/002	H/03/00406/019	NOVARTIS PHARMA GMBH	SI

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
CO-DIOVAN 160 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/002	H/03/00406/020	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/002	H/03/00406/021	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/002	H/03/00406/022	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/002	H/03/00406/023	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/002	H/03/00406/024	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/002	H/03/00406/025	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/002	H/03/00406/026	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/002	H/03/00406/027	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/002	H/03/00406/028	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/002	H/03/00406/029	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/002	H/03/00406/030	NOVARTIS PHARMA GMBH	SI

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
CO-DIOVAN 160 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/002	H/03/00406/031	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/002	H/03/00406/032	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/002	H/03/00406/033	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/002	H/03/00406/034	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/002	H/03/00406/019	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/002	H/03/00406/020	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/002	H/03/00406/021	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/002	H/03/00406/022	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/002	H/03/00406/023	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/002	H/03/00406/024	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/002	H/03/00406/025	NOVARTIS PHARMA GMBH	SI

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
CO-DIOVAN 160 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/002	H/03/00406/026	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/002	H/03/00406/027	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/002	H/03/00406/028	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/002	H/03/00406/029	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/002	H/03/00406/030	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/002	H/03/00406/031	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/002	H/03/00406/032	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/002	H/03/00406/033	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/002	H/03/00406/034	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/002	H/03/00406/019	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/002	H/03/00406/020	NOVARTIS PHARMA GMBH	SI

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
CO-DIOVAN 160 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/002	H/03/00406/021	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/002	H/03/00406/022	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/002	H/03/00406/023	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/002	H/03/00406/024	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/002	H/03/00406/025	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/002	H/03/00406/026	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/002	H/03/00406/027	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/002	H/03/00406/028	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/002	H/03/00406/029	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/002	H/03/00406/030	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/002	H/03/00406/031	NOVARTIS PHARMA GMBH	SI

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
CO-DIOVAN 160 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/002	H/03/00406/032	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/002	H/03/00406/033	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/002	H/03/00406/034	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/12,5 MG PLĚVELE DENGTOŠ TABLETĚS	SE/H/0565/002	LT/1/01/1235/002	NOVARTIS FINLAND OY	LT
CO-DIOVAN 160 MG/12,5 MG PLĚVELE DENGTOŠ TABLETĚS	SE/H/0565/002	LT/1/01/1235/002	NOVARTIS FINLAND OY	LT
CO-DIOVAN 160 MG/12,5 MG PLĚVELE DENGTOŠ TABLETĚS	SE/H/0565/002	LT/1/01/1235/002	NOVARTIS FINLAND OY	LT
CO-DIOVAN 160 MG/12,5 MG PLĚVELE DENGTOŠ TABLETĚS	SE/H/0565/002	LT/1/01/1235/002	NOVARTIS FINLAND OY	LT
Co-Diovan 160 mg/12,5 mg επικαλυμμένα με λεπτό υμένιο δισκία	SE/H/0565/002	18977	NOVARTIS PHARMACEUTICALS UK LIMITED	CY
Co-Diovan 160 mg/12,5 mg επικαλυμμένα με λεπτό υμένιο δισκία	SE/H/0565/002	18977	NOVARTIS PHARMACEUTICALS UK LIMITED	CY
Co-Diovan 160 mg/12,5 mg επικαλυμμένα με λεπτό υμένιο δισκία	SE/H/0565/002	18977	NOVARTIS PHARMACEUTICALS UK LIMITED	CY
Co-Diovan 160 mg/12,5 mg επικαλυμμένα με λεπτό υμένιο δισκία	SE/H/0565/002	18977	NOVARTIS PHARMACEUTICALS UK LIMITED	CY

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
Co-Diovan 160 mg/12.5 mg film-coated tablets	SE/H/0565/002	PA 13/91/2	NOVARTIS PHARMACEUTICALS UK LIMITED	IE
Co-Diovan 160 mg/12.5 mg film-coated tablets	SE/H/0565/002	PA 13/91/2	NOVARTIS PHARMACEUTICALS UK LIMITED	IE
Co-Diovan 160 mg/12.5 mg film-coated tablets	SE/H/0565/002	PA 13/91/2	NOVARTIS PHARMACEUTICALS UK LIMITED	IE
Co-Diovan 160 mg/12.5 mg film-coated tablets	SE/H/0565/002	PA 13/91/2	NOVARTIS PHARMACEUTICALS UK LIMITED	IE
CO-DIOVAN 160 MG/12.5 MG FILM-COATED TABLETS	SE/H/0565/002	088/00402	NOVARTIS PHARMACEUTICALS UK LIMITED	MT
CO-DIOVAN 160 MG/12.5 MG FILM-COATED TABLETS	SE/H/0565/002	088/00402	NOVARTIS PHARMACEUTICALS UK LIMITED	MT
CO-DIOVAN 160 MG/12.5 MG FILM-COATED TABLETS	SE/H/0565/002	088/00402	NOVARTIS PHARMACEUTICALS UK LIMITED	MT
CO-DIOVAN 160 MG/12.5 MG FILM-COATED TABLETS	SE/H/0565/002	088/00402	NOVARTIS PHARMACEUTICALS UK LIMITED	MT
CO-DIOVAN 160 MG/25 MG APVALKOTĀS TABLETES	SE/H/0565/003	02-0394	NOVARTIS FINLAND OY	LV
CO-DIOVAN 160 MG/25 MG APVALKOTĀS TABLETES	SE/H/0565/003	02-0394	NOVARTIS FINLAND OY	LV
CO-DIOVAN 160 MG/25 MG APVALKOTĀS TABLETES	SE/H/0565/003	02-0394	NOVARTIS FINLAND OY	LV

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
CO-DIOVAN 160 MG/25 MG APVALKOTĀS TABLETES	SE/H/0565/003	02-0394	NOVARTIS FINLAND OY	LV
Co-Diovan 160 mg/25 mg comprimidos revestidos por película	SE/H/0565/003	5022801	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 160 mg/25 mg comprimidos revestidos por película	SE/H/0565/003	5022819	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 160 mg/25 mg comprimidos revestidos por película	SE/H/0565/003	5273388	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 160 mg/25 mg comprimidos revestidos por película	SE/H/0565/003	5273487	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 160 mg/25 mg comprimidos revestidos por película	SE/H/0565/003	5273586	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 160 mg/25 mg comprimidos revestidos por película	SE/H/0565/003	5273685	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 160 mg/25 mg comprimidos revestidos por película	SE/H/0565/003	5022801	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 160 mg/25 mg comprimidos revestidos por película	SE/H/0565/003	5022819	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 160 mg/25 mg comprimidos revestidos por película	SE/H/0565/003	5273388	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 160 mg/25 mg comprimidos revestidos por película	SE/H/0565/003	5273487	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT

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
Co-Diovan 160 mg/25 mg comprimidos revestidos por película	SE/H/0565/003	5273586	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 160 mg/25 mg comprimidos revestidos por película	SE/H/0565/003	5273685	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 160 mg/25 mg comprimidos revestidos por película	SE/H/0565/003	5022801	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 160 mg/25 mg comprimidos revestidos por película	SE/H/0565/003	5022819	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 160 mg/25 mg comprimidos revestidos por película	SE/H/0565/003	5273388	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 160 mg/25 mg comprimidos revestidos por película	SE/H/0565/003	5273487	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 160 mg/25 mg comprimidos revestidos por película	SE/H/0565/003	5273586	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 160 mg/25 mg comprimidos revestidos por película	SE/H/0565/003	5273685	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 160 mg/25 mg comprimidos revestidos por película	SE/H/0565/003	5022801	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 160 mg/25 mg comprimidos revestidos por película	SE/H/0565/003	5022819	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 160 mg/25 mg comprimidos revestidos por película	SE/H/0565/003	5273388	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT

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
Co-Diovan 160 mg/25 mg comprimidos revestidos por película	SE/H/0565/003	5273487	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 160 mg/25 mg comprimidos revestidos por película	SE/H/0565/003	5273586	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 160 mg/25 mg comprimidos revestidos por película	SE/H/0565/003	5273685	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 160 mg/25 mg film-coated tablets	SE/H/0565/003	PA 13/91/3	NOVARTIS PHARMACEUTICALS UK LIMITED	IE
Co-Diovan 160 mg/25 mg film-coated tablets	SE/H/0565/003	PA 13/91/3	NOVARTIS PHARMACEUTICALS UK LIMITED	IE
Co-Diovan 160 mg/25 mg film-coated tablets	SE/H/0565/003	PA 13/91/3	NOVARTIS PHARMACEUTICALS UK LIMITED	IE
Co-Diovan 160 mg/25 mg film-coated tablets	SE/H/0565/003	PA 13/91/3	NOVARTIS PHARMACEUTICALS UK LIMITED	IE
CO-DIOVAN 160 MG/25 MG FILM-COATED TABLETS	SE/H/0565/003	088/00403	NOVARTIS PHARMACEUTICALS UK LIMITED	MT
CO-DIOVAN 160 MG/25 MG FILM-COATED TABLETS	SE/H/0565/003	088/00403	NOVARTIS PHARMACEUTICALS UK LIMITED	MT
CO-DIOVAN 160 MG/25 MG FILM-COATED TABLETS	SE/H/0565/003	088/00403	NOVARTIS PHARMACEUTICALS UK LIMITED	MT
CO-DIOVAN 160 MG/25 MG FILM-COATED TABLETS	SE/H/0565/003	088/00403	NOVARTIS PHARMACEUTICALS UK LIMITED	MT

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
CO-DIOVAN 160 MG/25 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/003	H/03/00406/035	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/25 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/003	H/03/00406/036	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/25 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/003	H/03/00406/037	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/25 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/003	H/03/00406/038	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/25 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/003	H/03/00406/039	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/25 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/003	H/03/00406/040	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/25 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/003	H/03/00406/041	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/25 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/003	H/03/00406/042	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/25 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/003	H/03/00406/043	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/25 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/003	H/03/00406/044	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/25 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/003	H/03/00406/045	NOVARTIS PHARMA GMBH	SI

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
CO-DIOVAN 160 MG/25 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/003	H/03/00406/046	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/25 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/003	H/03/00406/047	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/25 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/003	H/03/00406/048	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/25 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/003	H/03/00406/049	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/25 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/003	H/03/00406/050	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/25 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/003	H/03/00406/035	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/25 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/003	H/03/00406/036	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/25 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/003	H/03/00406/037	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/25 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/003	H/03/00406/038	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/25 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/003	H/03/00406/039	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/25 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/003	H/03/00406/040	NOVARTIS PHARMA GMBH	SI

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
CO-DIOVAN 160 MG/25 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/003	H/03/00406/041	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/25 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/003	H/03/00406/042	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/25 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/003	H/03/00406/043	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/25 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/003	H/03/00406/044	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/25 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/003	H/03/00406/045	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/25 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/003	H/03/00406/046	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/25 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/003	H/03/00406/047	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/25 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/003	H/03/00406/048	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/25 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/003	H/03/00406/049	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/25 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/003	H/03/00406/050	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/25 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/003	H/03/00406/035	NOVARTIS PHARMA GMBH	SI

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
CO-DIOVAN 160 MG/25 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/003	H/03/00406/036	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/25 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/003	H/03/00406/037	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/25 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/003	H/03/00406/038	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/25 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/003	H/03/00406/039	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/25 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/003	H/03/00406/040	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/25 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/003	H/03/00406/041	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/25 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/003	H/03/00406/042	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/25 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/003	H/03/00406/043	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/25 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/003	H/03/00406/044	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/25 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/003	H/03/00406/045	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/25 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/003	H/03/00406/046	NOVARTIS PHARMA GMBH	SI

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
CO-DIOVAN 160 MG/25 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/003	H/03/00406/047	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/25 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/003	H/03/00406/048	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/25 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/003	H/03/00406/049	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/25 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/003	H/03/00406/050	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/25 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/003	H/03/00406/035	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/25 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/003	H/03/00406/036	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/25 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/003	H/03/00406/037	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/25 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/003	H/03/00406/038	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/25 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/003	H/03/00406/039	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/25 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/003	H/03/00406/040	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/25 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/003	H/03/00406/041	NOVARTIS PHARMA GMBH	SI

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
CO-DIOVAN 160 MG/25 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/003	H/03/00406/042	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/25 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/003	H/03/00406/043	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/25 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/003	H/03/00406/044	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/25 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/003	H/03/00406/045	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/25 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/003	H/03/00406/046	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/25 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/003	H/03/00406/047	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/25 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/003	H/03/00406/048	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/25 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/003	H/03/00406/049	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/25 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/003	H/03/00406/050	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 160 MG/25 MG PLĚVELE DENGTOŠ TABLETŠ	SE/H/0565/003	LT/1/01/1235/003	NOVARTIS FINLAND OY	LT
CO-DIOVAN 160 MG/25 MG PLĚVELE DENGTOŠ TABLETŠ	SE/H/0565/003	LT/1/01/1235/003	NOVARTIS FINLAND OY	LT

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
CO-DIOVAN 160 MG/25 MG PLÉVELE DENGTOΣ TABLETÉS	SE/H/0565/003	LT/1/01/1235/003	NOVARTIS FINLAND OY	LT
CO-DIOVAN 160 MG/25 MG PLÉVELE DENGTOΣ TABLETÉS	SE/H/0565/003	LT/1/01/1235/003	NOVARTIS FINLAND OY	LT
Co-Diovan 160 mg/25 mg επικαλυμμένα με λεπτό υμένιο δισκία	SE/H/0565/003	19477	NOVARTIS PHARMACEUTICALS UK LIMITED	CY
Co-Diovan 160 mg/25 mg επικαλυμμένα με λεπτό υμένιο δισκία	SE/H/0565/003	19477	NOVARTIS PHARMACEUTICALS UK LIMITED	CY
Co-Diovan 160 mg/25 mg επικαλυμμένα με λεπτό υμένιο δισκία	SE/H/0565/003	19477	NOVARTIS PHARMACEUTICALS UK LIMITED	CY
Co-Diovan 160 mg/25 mg επικαλυμμένα με λεπτό υμένιο δισκία	SE/H/0565/003	19477	NOVARTIS PHARMACEUTICALS UK LIMITED	CY
Co-Diovan 160/12,5 mg õhukese polümeerikattega tabletid	SE/H/0565/002	380602	NOVARTIS FINLAND OY	EE
Co-Diovan 160/12,5 mg õhukese polümeerikattega tabletid	SE/H/0565/002	380602	NOVARTIS FINLAND OY	EE
Co-Diovan 160/12,5 mg õhukese polümeerikattega tabletid	SE/H/0565/002	380602	NOVARTIS FINLAND OY	EE
Co-Diovan 160/12,5 mg õhukese polümeerikattega tabletid	SE/H/0565/002	380602	NOVARTIS FINLAND OY	EE

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
Co-Diovan 160/12,5, 160 mg/12,5 mg filmomhulde tabletten	SE/H/0565/002	RVG 29491	NOVARTIS PHARMA B.V.	NL
Co-Diovan 160/12,5, 160 mg/12,5 mg filmomhulde tabletten	SE/H/0565/002	RVG 29491	NOVARTIS PHARMA B.V.	NL
Co-Diovan 160/12,5, 160 mg/12,5 mg filmomhulde tabletten	SE/H/0565/002	RVG 29491	NOVARTIS PHARMA B.V.	NL
Co-Diovan 160/12,5, 160 mg/12,5 mg filmomhulde tabletten	SE/H/0565/002	RVG 29491	NOVARTIS PHARMA B.V.	NL
Co-Diovan 160/12.5 mg filmom oblozene tablete	SE/H/0565/002	HR-H-819796318-01	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 160/12.5 mg filmom oblozene tablete	SE/H/0565/002	HR-H-819796318-02	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 160/12.5 mg filmom oblozene tablete	SE/H/0565/002	HR-H-819796318-03	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 160/12.5 mg filmom oblozene tablete	SE/H/0565/002	HR-H-819796318-04	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 160/12.5 mg filmom oblozene tablete	SE/H/0565/002	HR-H-819796318-05	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 160/12.5 mg filmom oblozene tablete	SE/H/0565/002	HR-H-819796318-06	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 160/12.5 mg filmom oblozene tablete	SE/H/0565/002	HR-H-819796318-07	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 160/12.5 mg filmom oblozene tablete	SE/H/0565/002	HR-H-819796318-08	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 160/12.5 mg filmom oblozene tablete	SE/H/0565/002	HR-H-819796318-01	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 160/12.5 mg filmom oblozene tablete	SE/H/0565/002	HR-H-819796318-02	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 160/12.5 mg	SE/H/0565/002	HR-H-819796318-03	NOVARTIS HRVATSKA	HR

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
filmom obložene tablete			D.O.O.	
Co-Diovan 160/12.5 mg filmom obložene tablete	SE/H/0565/002	HR-H-819796318-04	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 160/12.5 mg filmom obložene tablete	SE/H/0565/002	HR-H-819796318-05	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 160/12.5 mg filmom obložene tablete	SE/H/0565/002	HR-H-819796318-06	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 160/12.5 mg filmom obložene tablete	SE/H/0565/002	HR-H-819796318-07	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 160/12.5 mg filmom obložene tablete	SE/H/0565/002	HR-H-819796318-08	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 160/12.5 mg filmom obložene tablete	SE/H/0565/002	HR-H-819796318-01	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 160/12.5 mg filmom obložene tablete	SE/H/0565/002	HR-H-819796318-02	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 160/12.5 mg filmom obložene tablete	SE/H/0565/002	HR-H-819796318-03	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 160/12.5 mg filmom obložene tablete	SE/H/0565/002	HR-H-819796318-04	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 160/12.5 mg filmom obložene tablete	SE/H/0565/002	HR-H-819796318-05	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 160/12.5 mg filmom obložene tablete	SE/H/0565/002	HR-H-819796318-06	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 160/12.5 mg filmom obložene tablete	SE/H/0565/002	HR-H-819796318-07	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 160/12.5 mg filmom obložene tablete	SE/H/0565/002	HR-H-819796318-08	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 160/12.5 mg filmom obložene tablete	SE/H/0565/002	HR-H-819796318-01	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 160/12.5 mg filmom obložene tablete	SE/H/0565/002	HR-H-819796318-02	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 160/12.5 mg filmom obložene tablete	SE/H/0565/002	HR-H-819796318-03	NOVARTIS HRVATSKA D.O.O.	HR

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
Co-Diovan 160/12.5 mg filmom obložene tablete	SE/H/0565/002	HR-H-819796318-04	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 160/12.5 mg filmom obložene tablete	SE/H/0565/002	HR-H-819796318-05	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 160/12.5 mg filmom obložene tablete	SE/H/0565/002	HR-H-819796318-06	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 160/12.5 mg filmom obložene tablete	SE/H/0565/002	HR-H-819796318-07	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 160/12.5 mg filmom obložene tablete	SE/H/0565/002	HR-H-819796318-08	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 160/25 mg filmom obložene tablete	SE/H/0565/003	HR-H-133986302-01	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 160/25 mg filmom obložene tablete	SE/H/0565/003	HR-H-133986302-02	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 160/25 mg filmom obložene tablete	SE/H/0565/003	HR-H-133986302-03	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 160/25 mg filmom obložene tablete	SE/H/0565/003	HR-H-133986302-04	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 160/25 mg filmom obložene tablete	SE/H/0565/003	HR-H-133986302-05	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 160/25 mg filmom obložene tablete	SE/H/0565/003	HR-H-133986302-06	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 160/25 mg filmom obložene tablete	SE/H/0565/003	HR-H-133986302-07	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 160/25 mg filmom obložene tablete	SE/H/0565/003	HR-H-133986302-08	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 160/25 mg filmom obložene tablete	SE/H/0565/003	HR-H-133986302-01	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 160/25 mg filmom obložene tablete	SE/H/0565/003	HR-H-133986302-02	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 160/25 mg filmom obložene tablete	SE/H/0565/003	HR-H-133986302-03	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 160/25 mg	SE/H/0565/003	HR-H-133986302-04	NOVARTIS HRVATSKA	HR

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
filmom obložene tablete			D.O.O.	
Co-Diovan 160/25 mg filmom obložene tablete	SE/H/0565/003	HR-H-133986302-05	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 160/25 mg filmom obložene tablete	SE/H/0565/003	HR-H-133986302-06	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 160/25 mg filmom obložene tablete	SE/H/0565/003	HR-H-133986302-07	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 160/25 mg filmom obložene tablete	SE/H/0565/003	HR-H-133986302-08	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 160/25 mg filmom obložene tablete	SE/H/0565/003	HR-H-133986302-01	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 160/25 mg filmom obložene tablete	SE/H/0565/003	HR-H-133986302-02	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 160/25 mg filmom obložene tablete	SE/H/0565/003	HR-H-133986302-03	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 160/25 mg filmom obložene tablete	SE/H/0565/003	HR-H-133986302-04	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 160/25 mg filmom obložene tablete	SE/H/0565/003	HR-H-133986302-05	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 160/25 mg filmom obložene tablete	SE/H/0565/003	HR-H-133986302-06	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 160/25 mg filmom obložene tablete	SE/H/0565/003	HR-H-133986302-07	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 160/25 mg filmom obložene tablete	SE/H/0565/003	HR-H-133986302-08	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 160/25 mg filmom obložene tablete	SE/H/0565/003	HR-H-133986302-01	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 160/25 mg filmom obložene tablete	SE/H/0565/003	HR-H-133986302-02	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 160/25 mg filmom obložene tablete	SE/H/0565/003	HR-H-133986302-03	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 160/25 mg filmom obložene tablete	SE/H/0565/003	HR-H-133986302-04	NOVARTIS HRVATSKA D.O.O.	HR

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
Co-Diovan 160/25 mg filmom obložene tablete	SE/H/0565/003	HR-H-133986302-05	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 160/25 mg filmom obložene tablete	SE/H/0565/003	HR-H-133986302-06	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 160/25 mg filmom obložene tablete	SE/H/0565/003	HR-H-133986302-07	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 160/25 mg filmom obložene tablete	SE/H/0565/003	HR-H-133986302-08	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 160/25, 160 mg/25 mg filmomhulde tabletten	SE/H/0565/003	RVG 31122	NOVARTIS PHARMA B.V.	NL
Co-Diovan 160/25, 160 mg/25 mg filmomhulde tabletten	SE/H/0565/003	RVG 31122	NOVARTIS PHARMA B.V.	NL
Co-Diovan 160/25, 160 mg/25 mg filmomhulde tabletten	SE/H/0565/003	RVG 31122	NOVARTIS PHARMA B.V.	NL
Co-Diovan 160/25, 160 mg/25 mg filmomhulde tabletten	SE/H/0565/003	RVG 31122	NOVARTIS PHARMA B.V.	NL
Co-Diovan 320 mg/12,5 mg – Filmtabletten	SE/H/0565/004	1-27297	NOVARTIS PHARMA GMBH	AT
Co-Diovan 320 mg/12,5 mg – Filmtabletten	SE/H/0565/004	1-27297	NOVARTIS PHARMA GMBH	AT
Co-Diovan 320 mg/12,5 mg – Filmtabletten	SE/H/0565/004	1-27297	NOVARTIS PHARMA GMBH	AT
Co-Diovan 320 mg/12,5 mg – Filmtabletten	SE/H/0565/004	1-27297	NOVARTIS PHARMA GMBH	AT
Co-Diován 320 mg/12,5 mg comprimidos recubiertos con película	SE/H/0565/004	69.930	NOVARTIS FARMACÉUTICA S.A.	ES
Co-Diován 320 mg/12,5 mg comprimidos	SE/H/0565/004	69.930	NOVARTIS FARMACÉUTICA S.A.	ES

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
recubiertos con película				
Co-Diován 320 mg/12,5 mg comprimidos recubiertos con película	SE/H/0565/004	69.930	NOVARTIS FARMACÉUTICA S.A.	ES
Co-Diován 320 mg/12,5 mg comprimidos recubiertos con película	SE/H/0565/004	69.930	NOVARTIS FARMACÉUTICA S.A.	ES
Co-Diovan 320 mg/12,5 mg comprimidos revestidos por película	SE/H/0565/004	SE/H/0565/004	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 320 mg/12,5 mg comprimidos revestidos por película	SE/H/0565/004	SE/H/0565/004	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 320 mg/12,5 mg comprimidos revestidos por película	SE/H/0565/004	SE/H/0565/004	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 320 mg/12,5 mg comprimidos revestidos por película	SE/H/0565/004	SE/H/0565/004	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 320 mg/12,5 mg comprimidos revestidos por película	SE/H/0565/004	SE/H/0565/004	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 320 mg/12,5 mg comprimidos revestidos por película	SE/H/0565/004	5072400	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 320 mg/12,5 mg comprimidos revestidos por película	SE/H/0565/004	5072418	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 320 mg/12,5 mg comprimidos revestidos por película	SE/H/0565/004	5072426	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 320 mg/12,5 mg comprimidos	SE/H/0565/004	SE/H/0565/004	NOVARTIS FARMA - PRODUTOS	PT

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
revestidos por película			FARMACÊUTICOS S.A.	
Co-Diovan 320 mg/12,5 mg comprimidos revestidos por película	SE/H/0565/004	SE/H/0565/004	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 320 mg/12,5 mg comprimidos revestidos por película	SE/H/0565/004	SE/H/0565/004	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 320 mg/12,5 mg comprimidos revestidos por película	SE/H/0565/004	SE/H/0565/004	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 320 mg/12,5 mg comprimidos revestidos por película	SE/H/0565/004	SE/H/0565/004	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 320 mg/12,5 mg comprimidos revestidos por película	SE/H/0565/004	5072400	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 320 mg/12,5 mg comprimidos revestidos por película	SE/H/0565/004	5072418	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 320 mg/12,5 mg comprimidos revestidos por película	SE/H/0565/004	5072426	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 320 mg/12,5 mg comprimidos revestidos por película	SE/H/0565/004	SE/H/0565/004	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 320 mg/12,5 mg comprimidos revestidos por película	SE/H/0565/004	SE/H/0565/004	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 320 mg/12,5 mg comprimidos revestidos por película	SE/H/0565/004	SE/H/0565/004	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 320 mg/12,5 mg comprimidos	SE/H/0565/004	SE/H/0565/004	NOVARTIS FARMA - PRODUTOS	PT

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
revestidos por película			FARMACÊUTICOS S.A.	
Co-Diovan 320 mg/12,5 mg comprimidos revestidos por película	SE/H/0565/004	SE/H/0565/004	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 320 mg/12,5 mg comprimidos revestidos por película	SE/H/0565/004	5072400	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 320 mg/12,5 mg comprimidos revestidos por película	SE/H/0565/004	5072418	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 320 mg/12,5 mg comprimidos revestidos por película	SE/H/0565/004	5072426	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 320 mg/12,5 mg comprimidos revestidos por película	SE/H/0565/004	SE/H/0565/004	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 320 mg/12,5 mg comprimidos revestidos por película	SE/H/0565/004	SE/H/0565/004	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 320 mg/12,5 mg comprimidos revestidos por película	SE/H/0565/004	SE/H/0565/004	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 320 mg/12,5 mg comprimidos revestidos por película	SE/H/0565/004	SE/H/0565/004	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 320 mg/12,5 mg comprimidos revestidos por película	SE/H/0565/004	SE/H/0565/004	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 320 mg/12,5 mg comprimidos revestidos por película	SE/H/0565/004	5072400	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 320 mg/12,5 mg comprimidos	SE/H/0565/004	5072418	NOVARTIS FARMA - PRODUTOS	PT

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
revestidos por película			FARMACÊUTICOS S.A.	
Co-Diovan 320 mg/12,5 mg comprimidos revestidos por película	SE/H/0565/004	5072426	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
CO-DIOVAN 320 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/004	H/03/00406/051	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 320 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/004	H/03/00406/052	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 320 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/004	H/03/00406/053	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 320 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/004	H/03/00406/054	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 320 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/004	H/03/00406/055	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 320 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/004	H/03/00406/056	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 320 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/004	H/03/00406/057	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 320 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/004	H/03/00406/058	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 320 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/004	H/03/00406/059	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 320 MG/12,5 MG FILMSKO	SE/H/0565/004	H/03/00406/051	NOVARTIS PHARMA GMBH	SI

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
OBLOŽENE TABLETE				
CO-DIOVAN 320 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/004	H/03/00406/052	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 320 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/004	H/03/00406/053	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 320 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/004	H/03/00406/054	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 320 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/004	H/03/00406/055	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 320 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/004	H/03/00406/056	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 320 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/004	H/03/00406/057	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 320 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/004	H/03/00406/058	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 320 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/004	H/03/00406/059	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 320 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/004	H/03/00406/051	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 320 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/004	H/03/00406/052	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 320 MG/12,5 MG FILMSKO	SE/H/0565/004	H/03/00406/053	NOVARTIS PHARMA GMBH	SI

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
OBLOŽENE TABLETE				
CO-DIOVAN 320 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/004	H/03/00406/054	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 320 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/004	H/03/00406/055	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 320 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/004	H/03/00406/056	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 320 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/004	H/03/00406/057	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 320 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/004	H/03/00406/058	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 320 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/004	H/03/00406/059	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 320 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/004	H/03/00406/051	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 320 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/004	H/03/00406/052	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 320 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/004	H/03/00406/053	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 320 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/004	H/03/00406/054	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 320 MG/12,5 MG FILMSKO	SE/H/0565/004	H/03/00406/055	NOVARTIS PHARMA GMBH	SI

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
OBLOŽENE TABLETE				
CO-DIOVAN 320 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/004	H/03/00406/056	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 320 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/004	H/03/00406/057	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 320 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/004	H/03/00406/058	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 320 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/004	H/03/00406/059	NOVARTIS PHARMA GMBH	SI
Co-Diovan 320 mg/12.5 mg film-coated tablets	SE/H/0565/004	PA 13/91/4	NOVARTIS PHARMACEUTICALS UK LIMITED	IE
Co-Diovan 320 mg/12.5 mg film-coated tablets	SE/H/0565/004	PA 13/91/4	NOVARTIS PHARMACEUTICALS UK LIMITED	IE
Co-Diovan 320 mg/12.5 mg film-coated tablets	SE/H/0565/004	PA 13/91/4	NOVARTIS PHARMACEUTICALS UK LIMITED	IE
Co-Diovan 320 mg/12.5 mg film-coated tablets	SE/H/0565/004	PA 13/91/4	NOVARTIS PHARMACEUTICALS UK LIMITED	IE
CO-DIOVAN 320 MG/12.5 MG FILM-COATED TABLETS	SE/H/0565/004	088/00404	NOVARTIS PHARMACEUTICALS UK LIMITED	MT
CO-DIOVAN 320 MG/12.5 MG FILM-COATED TABLETS	SE/H/0565/004	088/00404	NOVARTIS PHARMACEUTICALS UK LIMITED	MT
CO-DIOVAN 320 MG/12.5 MG FILM-	SE/H/0565/004	088/00404	NOVARTIS PHARMACEUTICALS UK	MT

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
COATED TABLETS			LIMITED	
CO-DIOVAN 320 MG/12.5 MG FILM-COATED TABLETS	SE/H/0565/004	088/00404	NOVARTIS PHARMACEUTICALS UK LIMITED	MT
Co-Diovan 320 mg/25 mg - Filmtabletten	SE/H/0565/005	1-27298	NOVARTIS PHARMA GMBH	AT
Co-Diovan 320 mg/25 mg - Filmtabletten	SE/H/0565/005	1-27298	NOVARTIS PHARMA GMBH	AT
Co-Diovan 320 mg/25 mg - Filmtabletten	SE/H/0565/005	1-27298	NOVARTIS PHARMA GMBH	AT
Co-Diovan 320 mg/25 mg - Filmtabletten	SE/H/0565/005	1-27298	NOVARTIS PHARMA GMBH	AT
Co-Diovan 320 mg/25 mg film-coated tablets	SE/H/0565/005	PA 13/91/5	NOVARTIS PHARMACEUTICALS UK LIMITED	IE
Co-Diovan 320 mg/25 mg film-coated tablets	SE/H/0565/005	PA 13/91/5	NOVARTIS PHARMACEUTICALS UK LIMITED	IE
Co-Diovan 320 mg/25 mg film-coated tablets	SE/H/0565/005	PA 13/91/5	NOVARTIS PHARMACEUTICALS UK LIMITED	IE
Co-Diovan 320 mg/25 mg film-coated tablets	SE/H/0565/005	PA 13/91/5	NOVARTIS PHARMACEUTICALS UK LIMITED	IE
CO-DIOVAN 320 MG/25 MG FILM-COATED TABLETS	SE/H/0565/005	088/00405	NOVARTIS PHARMACEUTICALS UK LIMITED	MT
CO-DIOVAN 320 MG/25 MG FILM-COATED TABLETS	SE/H/0565/005	088/00405	NOVARTIS PHARMACEUTICALS UK LIMITED	MT
CO-DIOVAN 320 MG/25 MG FILM-COATED TABLETS	SE/H/0565/005	088/00405	NOVARTIS PHARMACEUTICALS UK LIMITED	MT

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
CO-DIOVAN 320 MG/25 MG FILM-COATED TABLETS	SE/H/0565/005	088/00405	NOVARTIS PHARMACEUTICALS UK LIMITED	MT
CO-DIOVAN 320 MG/25 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/005	H/03/00406/060	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 320 MG/25 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/005	H/03/00406/061	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 320 MG/25 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/005	H/03/00406/062	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 320 MG/25 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/005	H/03/00406/063	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 320 MG/25 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/005	H/03/00406/064	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 320 MG/25 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/005	H/03/00406/065	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 320 MG/25 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/005	H/03/00406/066	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 320 MG/25 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/005	H/03/00406/067	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 320 MG/25 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/005	H/03/00406/068	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 320 MG/25 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/005	H/03/00406/060	NOVARTIS PHARMA GMBH	SI

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
CO-DIOVAN 320 MG/25 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/005	H/03/00406/061	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 320 MG/25 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/005	H/03/00406/062	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 320 MG/25 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/005	H/03/00406/063	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 320 MG/25 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/005	H/03/00406/064	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 320 MG/25 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/005	H/03/00406/065	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 320 MG/25 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/005	H/03/00406/066	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 320 MG/25 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/005	H/03/00406/067	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 320 MG/25 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/005	H/03/00406/068	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 320 MG/25 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/005	H/03/00406/060	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 320 MG/25 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/005	H/03/00406/061	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 320 MG/25 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/005	H/03/00406/062	NOVARTIS PHARMA GMBH	SI

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
CO-DIOVAN 320 MG/25 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/005	H/03/00406/063	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 320 MG/25 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/005	H/03/00406/064	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 320 MG/25 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/005	H/03/00406/065	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 320 MG/25 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/005	H/03/00406/066	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 320 MG/25 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/005	H/03/00406/067	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 320 MG/25 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/005	H/03/00406/068	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 320 MG/25 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/005	H/03/00406/060	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 320 MG/25 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/005	H/03/00406/061	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 320 MG/25 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/005	H/03/00406/062	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 320 MG/25 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/005	H/03/00406/063	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 320 MG/25 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/005	H/03/00406/064	NOVARTIS PHARMA GMBH	SI

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
CO-DIOVAN 320 MG/25 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/005	H/03/00406/065	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 320 MG/25 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/005	H/03/00406/066	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 320 MG/25 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/005	H/03/00406/067	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 320 MG/25 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/005	H/03/00406/068	NOVARTIS PHARMA GMBH	SI
Co-Diovan 320/12,5 mg õhukese polümeerikattega tabletid	SE/H/0565/004	541107	NOVARTIS FINLAND OY	EE
Co-Diovan 320/12,5 mg õhukese polümeerikattega tabletid	SE/H/0565/004	541107	NOVARTIS FINLAND OY	EE
Co-Diovan 320/12,5 mg õhukese polümeerikattega tabletid	SE/H/0565/004	541107	NOVARTIS FINLAND OY	EE
Co-Diovan 320/12,5 mg õhukese polümeerikattega tabletid	SE/H/0565/004	541107	NOVARTIS FINLAND OY	EE
Co-Diovan 320/12,5, 320 mg/12,5 mg filmomhulde tabletten	SE/H/0565/004	RVG 100453	NOVARTIS PHARMA B.V.	NL
Co-Diovan 320/12,5, 320 mg/12,5 mg filmomhulde tabletten	SE/H/0565/004	RVG 100453	NOVARTIS PHARMA B.V.	NL

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
Co-Diovan 320/12,5, 320 mg/12,5 mg filmomhulde tabletten	SE/H/0565/004	RVG 100453	NOVARTIS PHARMA B.V.	NL
Co-Diovan 320/12,5, 320 mg/12,5 mg filmomhulde tabletten	SE/H/0565/004	RVG 100453	NOVARTIS PHARMA B.V.	NL
Co-Diovan 320/12.5 mg filmom obložene tablete	SE/H/0565/004	HR-H-174157013-01	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 320/12.5 mg filmom obložene tablete	SE/H/0565/004	HR-H-174157013-02	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 320/12.5 mg filmom obložene tablete	SE/H/0565/004	HR-H-174157013-03	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 320/12.5 mg filmom obložene tablete	SE/H/0565/004	HR-H-174157013-04	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 320/12.5 mg filmom obložene tablete	SE/H/0565/004	HR-H-174157013-05	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 320/12.5 mg filmom obložene tablete	SE/H/0565/004	HR-H-174157013-06	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 320/12.5 mg filmom obložene tablete	SE/H/0565/004	HR-H-174157013-07	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 320/12.5 mg filmom obložene tablete	SE/H/0565/004	HR-H-174157013-08	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 320/12.5 mg filmom obložene tablete	SE/H/0565/004	HR-H-174157013-09	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 320/12.5 mg filmom obložene tablete	SE/H/0565/004	HR-H-174157013-01	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 320/12.5 mg filmom obložene tablete	SE/H/0565/004	HR-H-174157013-02	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 320/12.5 mg filmom obložene tablete	SE/H/0565/004	HR-H-174157013-03	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 320/12.5 mg filmom obložene tablete	SE/H/0565/004	HR-H-174157013-04	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 320/12.5 mg	SE/H/0565/004	HR-H-174157013-05	NOVARTIS HRVATSKA	HR

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
filmom obložene tablete			D.O.O.	
Co-Diovan 320/12.5 mg filmom obložene tablete	SE/H/0565/004	HR-H-174157013-06	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 320/12.5 mg filmom obložene tablete	SE/H/0565/004	HR-H-174157013-07	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 320/12.5 mg filmom obložene tablete	SE/H/0565/004	HR-H-174157013-08	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 320/12.5 mg filmom obložene tablete	SE/H/0565/004	HR-H-174157013-09	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 320/12.5 mg filmom obložene tablete	SE/H/0565/004	HR-H-174157013-01	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 320/12.5 mg filmom obložene tablete	SE/H/0565/004	HR-H-174157013-02	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 320/12.5 mg filmom obložene tablete	SE/H/0565/004	HR-H-174157013-03	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 320/12.5 mg filmom obložene tablete	SE/H/0565/004	HR-H-174157013-04	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 320/12.5 mg filmom obložene tablete	SE/H/0565/004	HR-H-174157013-05	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 320/12.5 mg filmom obložene tablete	SE/H/0565/004	HR-H-174157013-06	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 320/12.5 mg filmom obložene tablete	SE/H/0565/004	HR-H-174157013-07	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 320/12.5 mg filmom obložene tablete	SE/H/0565/004	HR-H-174157013-08	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 320/12.5 mg filmom obložene tablete	SE/H/0565/004	HR-H-174157013-09	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 320/12.5 mg filmom obložene tablete	SE/H/0565/004	HR-H-174157013-01	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 320/12.5 mg filmom obložene tablete	SE/H/0565/004	HR-H-174157013-02	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 320/12.5 mg filmom obložene tablete	SE/H/0565/004	HR-H-174157013-03	NOVARTIS HRVATSKA D.O.O.	HR

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
Co-Diovan 320/12.5 mg filmom obložene tablete	SE/H/0565/004	HR-H-174157013-04	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 320/12.5 mg filmom obložene tablete	SE/H/0565/004	HR-H-174157013-05	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 320/12.5 mg filmom obložene tablete	SE/H/0565/004	HR-H-174157013-06	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 320/12.5 mg filmom obložene tablete	SE/H/0565/004	HR-H-174157013-07	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 320/12.5 mg filmom obložene tablete	SE/H/0565/004	HR-H-174157013-08	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 320/12.5 mg filmom obložene tablete	SE/H/0565/004	HR-H-174157013-09	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 320/25 mg comprimidos revestidos por película	SE/H/0565/005	SE/H/0565/005	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 320/25 mg comprimidos revestidos por película	SE/H/0565/005	SE/H/0565/005	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 320/25 mg comprimidos revestidos por película	SE/H/0565/005	SE/H/0565/005	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 320/25 mg comprimidos revestidos por película	SE/H/0565/005	SE/H/0565/005	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 320/25 mg comprimidos revestidos por película	SE/H/0565/005	5072434	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 320/25 mg comprimidos revestidos por película	SE/H/0565/005	5072442	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT

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
Co-Diovan 320/25 mg comprimidos revestidos por película	SE/H/0565/005	5072459	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 320/25 mg comprimidos revestidos por película	SE/H/0565/005	SE/H/0565/005	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 320/25 mg comprimidos revestidos por película	SE/H/0565/005	SE/H/0565/005	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 320/25 mg comprimidos revestidos por película	SE/H/0565/005	SE/H/0565/005	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 320/25 mg comprimidos revestidos por película	SE/H/0565/005	SE/H/0565/005	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 320/25 mg comprimidos revestidos por película	SE/H/0565/005	SE/H/0565/005	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 320/25 mg comprimidos revestidos por película	SE/H/0565/005	5072434	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 320/25 mg comprimidos revestidos por película	SE/H/0565/005	5072442	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 320/25 mg comprimidos revestidos por película	SE/H/0565/005	5072459	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 320/25 mg comprimidos revestidos por película	SE/H/0565/005	SE/H/0565/005	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 320/25 mg comprimidos revestidos por película	SE/H/0565/005	SE/H/0565/005	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT

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
Co-Diovan 320/25 mg comprimidos revestidos por película	SE/H/0565/005	SE/H/0565/005	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 320/25 mg comprimidos revestidos por película	SE/H/0565/005	SE/H/0565/005	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 320/25 mg comprimidos revestidos por película	SE/H/0565/005	SE/H/0565/005	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 320/25 mg comprimidos revestidos por película	SE/H/0565/005	5072434	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 320/25 mg comprimidos revestidos por película	SE/H/0565/005	5072442	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 320/25 mg comprimidos revestidos por película	SE/H/0565/005	5072459	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 320/25 mg comprimidos revestidos por película	SE/H/0565/005	SE/H/0565/005	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 320/25 mg comprimidos revestidos por película	SE/H/0565/005	SE/H/0565/005	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 320/25 mg comprimidos revestidos por película	SE/H/0565/005	SE/H/0565/005	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 320/25 mg comprimidos revestidos por película	SE/H/0565/005	SE/H/0565/005	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 320/25 mg comprimidos revestidos por película	SE/H/0565/005	SE/H/0565/005	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT

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
Co-Diovan 320/25 mg comprimidos revestidos por película	SE/H/0565/005	5072434	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 320/25 mg comprimidos revestidos por película	SE/H/0565/005	5072442	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 320/25 mg comprimidos revestidos por película	SE/H/0565/005	5072459	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 320/25 mg filmom obložene tablete	SE/H/0565/005	HR-H-063837675-01	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 320/25 mg filmom obložene tablete	SE/H/0565/005	HR-H-063837675-02	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 320/25 mg filmom obložene tablete	SE/H/0565/005	HR-H-063837675-03	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 320/25 mg filmom obložene tablete	SE/H/0565/005	HR-H-063837675-04	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 320/25 mg filmom obložene tablete	SE/H/0565/005	HR-H-063837675-05	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 320/25 mg filmom obložene tablete	SE/H/0565/005	HR-H-063837675-06	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 320/25 mg filmom obložene tablete	SE/H/0565/005	HR-H-063837675-07	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 320/25 mg filmom obložene tablete	SE/H/0565/005	HR-H-063837675-08	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 320/25 mg filmom obložene tablete	SE/H/0565/005	HR-H-063837675-09	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 320/25 mg filmom obložene tablete	SE/H/0565/005	HR-H-063837675-01	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 320/25 mg filmom obložene tablete	SE/H/0565/005	HR-H-063837675-02	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 320/25 mg filmom obložene tablete	SE/H/0565/005	HR-H-063837675-03	NOVARTIS HRVATSKA D.O.O.	HR

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
Co-Diovan 320/25 mg filmom obložene tablete	SE/H/0565/005	HR-H-063837675-04	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 320/25 mg filmom obložene tablete	SE/H/0565/005	HR-H-063837675-05	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 320/25 mg filmom obložene tablete	SE/H/0565/005	HR-H-063837675-06	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 320/25 mg filmom obložene tablete	SE/H/0565/005	HR-H-063837675-07	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 320/25 mg filmom obložene tablete	SE/H/0565/005	HR-H-063837675-08	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 320/25 mg filmom obložene tablete	SE/H/0565/005	HR-H-063837675-09	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 320/25 mg filmom obložene tablete	SE/H/0565/005	HR-H-063837675-01	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 320/25 mg filmom obložene tablete	SE/H/0565/005	HR-H-063837675-02	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 320/25 mg filmom obložene tablete	SE/H/0565/005	HR-H-063837675-03	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 320/25 mg filmom obložene tablete	SE/H/0565/005	HR-H-063837675-04	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 320/25 mg filmom obložene tablete	SE/H/0565/005	HR-H-063837675-05	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 320/25 mg filmom obložene tablete	SE/H/0565/005	HR-H-063837675-06	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 320/25 mg filmom obložene tablete	SE/H/0565/005	HR-H-063837675-07	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 320/25 mg filmom obložene tablete	SE/H/0565/005	HR-H-063837675-08	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 320/25 mg filmom obložene tablete	SE/H/0565/005	HR-H-063837675-09	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 320/25 mg filmom obložene tablete	SE/H/0565/005	HR-H-063837675-01	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 320/25 mg	SE/H/0565/005	HR-H-063837675-02	NOVARTIS HRVATSKA	HR

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
filmom oblozene tablete			D.O.O.	
Co-Diovan 320/25 mg filmom oblozene tablete	SE/H/0565/005	HR-H-063837675-03	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 320/25 mg filmom oblozene tablete	SE/H/0565/005	HR-H-063837675-04	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 320/25 mg filmom oblozene tablete	SE/H/0565/005	HR-H-063837675-05	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 320/25 mg filmom oblozene tablete	SE/H/0565/005	HR-H-063837675-06	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 320/25 mg filmom oblozene tablete	SE/H/0565/005	HR-H-063837675-07	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 320/25 mg filmom oblozene tablete	SE/H/0565/005	HR-H-063837675-08	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 320/25 mg filmom oblozene tablete	SE/H/0565/005	HR-H-063837675-09	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 320/25 mg õhukese polümeerikattega tabletid	SE/H/0565/005	541207	NOVARTIS FINLAND OY	EE
Co-Diovan 320/25 mg õhukese polümeerikattega tabletid	SE/H/0565/005	541207	NOVARTIS FINLAND OY	EE
Co-Diovan 320/25 mg õhukese polümeerikattega tabletid	SE/H/0565/005	541207	NOVARTIS FINLAND OY	EE
Co-Diovan 320/25 mg õhukese polümeerikattega tabletid	SE/H/0565/005	541207	NOVARTIS FINLAND OY	EE
Co-Diovan 320/25, 320 mg/25 mg filmomhulde	SE/H/0565/005	RVG 100454	NOVARTIS PHARMA B.V.	NL

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
tabletten				
Co-Diovan 320/25, 320 mg/25 mg filmomhulde tabletten	SE/H/0565/005	RVG 100454	NOVARTIS PHARMA B.V.	NL
Co-Diovan 320/25, 320 mg/25 mg filmomhulde tabletten	SE/H/0565/005	RVG 100454	NOVARTIS PHARMA B.V.	NL
Co-Diovan 320/25, 320 mg/25 mg filmomhulde tabletten	SE/H/0565/005	RVG 100454	NOVARTIS PHARMA B.V.	NL
Co-Diovan 80 mg /12,5 mg comprimidos revestidos por película	SE/H/0565/001	2644383	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 80 mg /12,5 mg comprimidos revestidos por película	SE/H/0565/001	2644482	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 80 mg /12,5 mg comprimidos revestidos por película	SE/H/0565/001	2644581	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 80 mg /12,5 mg comprimidos revestidos por película	SE/H/0565/001	5022744	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 80 mg /12,5 mg comprimidos revestidos por película	SE/H/0565/001	5022751	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 80 mg /12,5 mg comprimidos revestidos por película	SE/H/0565/001	2644383	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 80 mg /12,5 mg comprimidos revestidos por película	SE/H/0565/001	2644482	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 80 mg /12,5 mg comprimidos	SE/H/0565/001	2644581	NOVARTIS FARMA - PRODUTOS	PT

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
revestidos por película			FARMACÊUTICOS S.A.	
Co-Diovan 80 mg /12,5 mg comprimidos revestidos por película	SE/H/0565/001	5022744	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 80 mg /12,5 mg comprimidos revestidos por película	SE/H/0565/001	5022751	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 80 mg /12,5 mg comprimidos revestidos por película	SE/H/0565/001	2644383	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 80 mg /12,5 mg comprimidos revestidos por película	SE/H/0565/001	2644482	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 80 mg /12,5 mg comprimidos revestidos por película	SE/H/0565/001	2644581	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 80 mg /12,5 mg comprimidos revestidos por película	SE/H/0565/001	5022744	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 80 mg /12,5 mg comprimidos revestidos por película	SE/H/0565/001	5022751	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 80 mg /12,5 mg comprimidos revestidos por película	SE/H/0565/001	2644383	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 80 mg /12,5 mg comprimidos revestidos por película	SE/H/0565/001	2644482	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 80 mg /12,5 mg comprimidos revestidos por película	SE/H/0565/001	2644581	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 80 mg /12,5 mg comprimidos	SE/H/0565/001	5022744	NOVARTIS FARMA - PRODUTOS	PT

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
revestidos por película			FARMACÊUTICOS S.A.	
Co-Diovan 80 mg /12,5 mg comprimidos revestidos por película	SE/H/0565/001	5022751	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Co-Diovan 80 mg/12,5 mg – Filmtabletten	SE/H/0565/001	1 - 22463	NOVARTIS PHARMA GMBH	AT
Co-Diovan 80 mg/12,5 mg – Filmtabletten	SE/H/0565/001	1 - 22463	NOVARTIS PHARMA GMBH	AT
Co-Diovan 80 mg/12,5 mg – Filmtabletten	SE/H/0565/001	1 - 22463	NOVARTIS PHARMA GMBH	AT
Co-Diovan 80 mg/12,5 mg – Filmtabletten	SE/H/0565/001	1 - 22463	NOVARTIS PHARMA GMBH	AT
CO-DIOVAN 80 MG/12,5 MG APVALKOTĀS TABLETES	SE/H/0565/001	98-0310	NOVARTIS FINLAND OY	LV
CO-DIOVAN 80 MG/12,5 MG APVALKOTĀS TABLETES	SE/H/0565/001	98-0310	NOVARTIS FINLAND OY	LV
CO-DIOVAN 80 MG/12,5 MG APVALKOTĀS TABLETES	SE/H/0565/001	98-0310	NOVARTIS FINLAND OY	LV
CO-DIOVAN 80 MG/12,5 MG APVALKOTĀS TABLETES	SE/H/0565/001	98-0310	NOVARTIS FINLAND OY	LV
Co-Diován 80 mg/12,5 mg comprimidos recubiertos con película	SE/H/0565/001	62.206	NOVARTIS FARMACÉUTICA S.A.	ES
Co-Diován 80 mg/12,5 mg comprimidos recubiertos con película	SE/H/0565/001	62.206	NOVARTIS FARMACÉUTICA S.A.	ES
Co-Diován 80 mg/12,5 mg comprimidos recubiertos con película	SE/H/0565/001	62.206	NOVARTIS FARMACÉUTICA S.A.	ES

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
Co-Diován 80 mg/12,5 mg comprimidos recubiertos con película	SE/H/0565/001	62.206	NOVARTIS FARMACÉUTICA S.A.	ES
CO-DIOVAN 80 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/001	H/03/00406/001	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 80 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/001	H/03/00406/002	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 80 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/001	H/03/00406/003	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 80 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/001	H/03/00406/004	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 80 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/001	H/03/00406/005	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 80 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/001	H/03/00406/006	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 80 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/001	H/03/00406/007	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 80 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/001	H/03/00406/008	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 80 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/001	H/03/00406/009	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 80 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/001	H/03/00406/010	NOVARTIS PHARMA GMBH	SI

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
CO-DIOVAN 80 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/001	H/03/00406/011	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 80 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/001	H/03/00406/012	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 80 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/001	H/03/00406/013	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 80 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/001	H/03/00406/014	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 80 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/001	H/03/00406/015	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 80 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/001	H/03/00406/016	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 80 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/001	H/03/00406/017	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 80 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/001	H/03/00406/018	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 80 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/001	H/03/00406/001	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 80 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/001	H/03/00406/002	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 80 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/001	H/03/00406/003	NOVARTIS PHARMA GMBH	SI

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
CO-DIOVAN 80 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/001	H/03/00406/004	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 80 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/001	H/03/00406/005	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 80 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/001	H/03/00406/006	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 80 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/001	H/03/00406/007	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 80 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/001	H/03/00406/008	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 80 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/001	H/03/00406/009	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 80 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/001	H/03/00406/010	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 80 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/001	H/03/00406/011	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 80 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/001	H/03/00406/012	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 80 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/001	H/03/00406/013	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 80 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/001	H/03/00406/014	NOVARTIS PHARMA GMBH	SI

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
CO-DIOVAN 80 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/001	H/03/00406/015	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 80 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/001	H/03/00406/016	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 80 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/001	H/03/00406/017	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 80 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/001	H/03/00406/018	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 80 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/001	H/03/00406/001	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 80 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/001	H/03/00406/002	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 80 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/001	H/03/00406/003	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 80 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/001	H/03/00406/004	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 80 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/001	H/03/00406/005	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 80 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/001	H/03/00406/006	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 80 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/001	H/03/00406/007	NOVARTIS PHARMA GMBH	SI

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
CO-DIOVAN 80 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/001	H/03/00406/008	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 80 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/001	H/03/00406/009	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 80 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/001	H/03/00406/010	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 80 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/001	H/03/00406/011	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 80 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/001	H/03/00406/012	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 80 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/001	H/03/00406/013	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 80 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/001	H/03/00406/014	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 80 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/001	H/03/00406/015	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 80 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/001	H/03/00406/016	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 80 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/001	H/03/00406/017	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 80 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/001	H/03/00406/018	NOVARTIS PHARMA GMBH	SI

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
CO-DIOVAN 80 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/001	H/03/00406/001	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 80 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/001	H/03/00406/002	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 80 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/001	H/03/00406/003	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 80 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/001	H/03/00406/004	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 80 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/001	H/03/00406/005	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 80 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/001	H/03/00406/006	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 80 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/001	H/03/00406/007	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 80 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/001	H/03/00406/008	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 80 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/001	H/03/00406/009	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 80 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/001	H/03/00406/010	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 80 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/001	H/03/00406/011	NOVARTIS PHARMA GMBH	SI

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
CO-DIOVAN 80 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/001	H/03/00406/012	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 80 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/001	H/03/00406/013	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 80 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/001	H/03/00406/014	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 80 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/001	H/03/00406/015	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 80 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/001	H/03/00406/016	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 80 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/001	H/03/00406/017	NOVARTIS PHARMA GMBH	SI
CO-DIOVAN 80 MG/12,5 MG FILMSKO OBLOŽENE TABLETE	SE/H/0565/001	H/03/00406/018	NOVARTIS PHARMA GMBH	SI
Co-Diovan 80 mg/12,5 mg plėvele dengtos tabletės	SE/H/0565/001	LT/1/01/1235/001	NOVARTIS FINLAND OY	LT
Co-Diovan 80 mg/12,5 mg plėvele dengtos tabletės	SE/H/0565/001	LT/1/01/1235/001	NOVARTIS FINLAND OY	LT
Co-Diovan 80 mg/12,5 mg plėvele dengtos tabletės	SE/H/0565/001	LT/1/01/1235/001	NOVARTIS FINLAND OY	LT
Co-Diovan 80 mg/12,5 mg plėvele dengtos tabletės	SE/H/0565/001	LT/1/01/1235/001	NOVARTIS FINLAND OY	LT

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
Co-Diovan 80 mg/12,5 mg επικαλυμμένα με λεπτό υμένιο δισκία	SE/H/0565/001	17851	NOVARTIS PHARMACEUTICALS UK LIMITED	CY
Co-Diovan 80 mg/12,5 mg επικαλυμμένα με λεπτό υμένιο δισκία	SE/H/0565/001	17851	NOVARTIS PHARMACEUTICALS UK LIMITED	CY
Co-Diovan 80 mg/12,5 mg επικαλυμμένα με λεπτό υμένιο δισκία	SE/H/0565/001	17851	NOVARTIS PHARMACEUTICALS UK LIMITED	CY
Co-Diovan 80 mg/12,5 mg επικαλυμμένα με λεπτό υμένιο δισκία	SE/H/0565/001	17851	NOVARTIS PHARMACEUTICALS UK LIMITED	CY
CO-DIOVAN 80 MG/12.5 MG FILM-COATED TABLETS	SE/H/0565/001	PA 13/91/1	NOVARTIS PHARMACEUTICALS UK LIMITED	IE
CO-DIOVAN 80 MG/12.5 MG FILM-COATED TABLETS	SE/H/0565/001	PA 13/91/1	NOVARTIS PHARMACEUTICALS UK LIMITED	IE
CO-DIOVAN 80 MG/12.5 MG FILM-COATED TABLETS	SE/H/0565/001	PA 13/91/1	NOVARTIS PHARMACEUTICALS UK LIMITED	IE
CO-DIOVAN 80 MG/12.5 MG FILM-COATED TABLETS	SE/H/0565/001	PA 13/91/1	NOVARTIS PHARMACEUTICALS UK LIMITED	IE
CO-DIOVAN 80 MG/12.5 MG FILM-COATED TABLETS	SE/H/0565/001	088/00401	NOVARTIS PHARMACEUTICALS UK LIMITED	MT
CO-DIOVAN 80 MG/12.5 MG FILM-COATED TABLETS	SE/H/0565/001	088/00401	NOVARTIS PHARMACEUTICALS UK LIMITED	MT
CO-DIOVAN 80 MG/12.5 MG FILM-COATED TABLETS	SE/H/0565/001	088/00401	NOVARTIS PHARMACEUTICALS UK LIMITED	MT

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
CO-DIOVAN 80 MG/12.5 MG FILM-COATED TABLETS	SE/H/0565/001	088/00401	NOVARTIS PHARMACEUTICALS UK LIMITED	MT
Co-Diovan 80/12,5 mg õhukese polümeerikattega tabletid	SE/H/0565/001	246499	NOVARTIS FINLAND OY	EE
Co-Diovan 80/12,5 mg õhukese polümeerikattega tabletid	SE/H/0565/001	246499	NOVARTIS FINLAND OY	EE
Co-Diovan 80/12,5 mg õhukese polümeerikattega tabletid	SE/H/0565/001	246499	NOVARTIS FINLAND OY	EE
Co-Diovan 80/12,5 mg õhukese polümeerikattega tabletid	SE/H/0565/001	246499	NOVARTIS FINLAND OY	EE
Co-Diovan 80/12,5, 80 mg/12,5 mg filmomhulde tabletten	SE/H/0565/001	RVG 22365	NOVARTIS PHARMA B.V.	NL
Co-Diovan 80/12,5, 80 mg/12,5 mg filmomhulde tabletten	SE/H/0565/001	RVG 22365	NOVARTIS PHARMA B.V.	NL
Co-Diovan 80/12,5, 80 mg/12,5 mg filmomhulde tabletten	SE/H/0565/001	RVG 22365	NOVARTIS PHARMA B.V.	NL
Co-Diovan 80/12,5, 80 mg/12,5 mg filmomhulde tabletten	SE/H/0565/001	RVG 22365	NOVARTIS PHARMA B.V.	NL
Co-Diovan 80/12.5 mg filmom oblozene tablete	SE/H/0565/001	HR-H-035681755-01	NOVARTIS HRVATSKA D.O.O.	HR

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
Co-Diovan 80/12.5 mg filmom obložene tablete	SE/H/0565/001	HR-H-035681755-02	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 80/12.5 mg filmom obložene tablete	SE/H/0565/001	HR-H-035681755-03	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 80/12.5 mg filmom obložene tablete	SE/H/0565/001	HR-H-035681755-04	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 80/12.5 mg filmom obložene tablete	SE/H/0565/001	HR-H-035681755-05	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 80/12.5 mg filmom obložene tablete	SE/H/0565/001	HR-H-035681755-06	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 80/12.5 mg filmom obložene tablete	SE/H/0565/001	HR-H-035681755-07	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 80/12.5 mg filmom obložene tablete	SE/H/0565/001	HR-H-035681755-08	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 80/12.5 mg filmom obložene tablete	SE/H/0565/001	HR-H-035681755-09	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 80/12.5 mg filmom obložene tablete	SE/H/0565/001	HR-H-035681755-01	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 80/12.5 mg filmom obložene tablete	SE/H/0565/001	HR-H-035681755-02	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 80/12.5 mg filmom obložene tablete	SE/H/0565/001	HR-H-035681755-03	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 80/12.5 mg filmom obložene tablete	SE/H/0565/001	HR-H-035681755-04	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 80/12.5 mg filmom obložene tablete	SE/H/0565/001	HR-H-035681755-05	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 80/12.5 mg filmom obložene tablete	SE/H/0565/001	HR-H-035681755-06	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 80/12.5 mg filmom obložene tablete	SE/H/0565/001	HR-H-035681755-07	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 80/12.5 mg filmom obložene tablete	SE/H/0565/001	HR-H-035681755-08	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 80/12.5 mg	SE/H/0565/001	HR-H-035681755-09	NOVARTIS HRVATSKA	HR

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
filmom obložene tablete			D.O.O.	
Co-Diovan 80/12.5 mg filmom obložene tablete	SE/H/0565/001	HR-H-035681755-01	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 80/12.5 mg filmom obložene tablete	SE/H/0565/001	HR-H-035681755-02	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 80/12.5 mg filmom obložene tablete	SE/H/0565/001	HR-H-035681755-03	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 80/12.5 mg filmom obložene tablete	SE/H/0565/001	HR-H-035681755-04	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 80/12.5 mg filmom obložene tablete	SE/H/0565/001	HR-H-035681755-05	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 80/12.5 mg filmom obložene tablete	SE/H/0565/001	HR-H-035681755-06	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 80/12.5 mg filmom obložene tablete	SE/H/0565/001	HR-H-035681755-07	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 80/12.5 mg filmom obložene tablete	SE/H/0565/001	HR-H-035681755-08	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 80/12.5 mg filmom obložene tablete	SE/H/0565/001	HR-H-035681755-09	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 80/12.5 mg filmom obložene tablete	SE/H/0565/001	HR-H-035681755-01	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 80/12.5 mg filmom obložene tablete	SE/H/0565/001	HR-H-035681755-02	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 80/12.5 mg filmom obložene tablete	SE/H/0565/001	HR-H-035681755-03	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 80/12.5 mg filmom obložene tablete	SE/H/0565/001	HR-H-035681755-04	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 80/12.5 mg filmom obložene tablete	SE/H/0565/001	HR-H-035681755-05	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 80/12.5 mg filmom obložene tablete	SE/H/0565/001	HR-H-035681755-06	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 80/12.5 mg filmom obložene tablete	SE/H/0565/001	HR-H-035681755-07	NOVARTIS HRVATSKA D.O.O.	HR

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
Co-Diovan 80/12.5 mg filmom obložene tablete	SE/H/0565/001	HR-H-035681755-08	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan 80/12.5 mg filmom obložene tablete	SE/H/0565/001	HR-H-035681755-09	NOVARTIS HRVATSKA D.O.O.	HR
Co-Diovan forte 160 mg/12,5 mg – Filmtabletten	SE/H/0565/002	1 – 25101	NOVARTIS PHARMA GMBH	AT
Co-Diovan forte 160 mg/12,5 mg – Filmtabletten	SE/H/0565/002	1 – 25101	NOVARTIS PHARMA GMBH	AT
Co-Diovan forte 160 mg/12,5 mg – Filmtabletten	SE/H/0565/002	1 – 25101	NOVARTIS PHARMA GMBH	AT
Co-Diovan forte 160 mg/12,5 mg – Filmtabletten	SE/H/0565/002	1 – 25101	NOVARTIS PHARMA GMBH	AT
Co-Diován Forte 160 mg/25 mg comprimidos recubiertos con película	SE/H/0565/003	66.526	NOVARTIS FARMACÉUTICA S.A.	ES
Co-Diován Forte 160 mg/25 mg comprimidos recubiertos con película	SE/H/0565/003	66.526	NOVARTIS FARMACÉUTICA S.A.	ES
Co-Diován Forte 160 mg/25 mg comprimidos recubiertos con película	SE/H/0565/003	66.526	NOVARTIS FARMACÉUTICA S.A.	ES
Co-Diován Forte 160 mg/25 mg comprimidos recubiertos con película	SE/H/0565/003	66.526	NOVARTIS FARMACÉUTICA S.A.	ES
Co-Diován Forte 320 mg/25 mg comprimidos recubiertos con película	SE/H/0565/005	69.942	NOVARTIS FARMACÉUTICA S.A.	ES
Co-Diován Forte 320 mg/25 mg comprimidos	SE/H/0565/005	69.942	NOVARTIS FARMACÉUTICA S.A.	ES

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
recubiertos con película				
Co-Diován Forte 320 mg/25 mg comprimidos recubiertos con película	SE/H/0565/005	69.942	NOVARTIS FARMACÉUTICA S.A.	ES
Co-Diován Forte 320 mg/25 mg comprimidos recubiertos con película	SE/H/0565/005	69.942	NOVARTIS FARMACÉUTICA S.A.	ES
Co-Diovan fortissimum 160 mg/25 mg - Filmdabletten	SE/H/0565/003	1-25735	NOVARTIS PHARMA GMBH	AT
Co-Diovan fortissimum 160 mg/25 mg - Filmdabletten	SE/H/0565/003	1-25735	NOVARTIS PHARMA GMBH	AT
Co-Diovan fortissimum 160 mg/25 mg - Filmdabletten	SE/H/0565/003	1-25735	NOVARTIS PHARMA GMBH	AT
Co-Diovan fortissimum 160 mg/25 mg - Filmdabletten	SE/H/0565/003	1-25735	NOVARTIS PHARMA GMBH	AT
Co-Diovan, 160/25 mg õhukese polümeerikattega tabletid	SE/H/0565/003	380702	NOVARTIS FINLAND OY	EE
Co-Diovan, 160/25 mg õhukese polümeerikattega tabletid	SE/H/0565/003	380702	NOVARTIS FINLAND OY	EE
Co-Diovan, 160/25 mg õhukese polümeerikattega tabletid	SE/H/0565/003	380702	NOVARTIS FINLAND OY	EE
Co-Diovan, 160/25 mg õhukese	SE/H/0565/003	380702	NOVARTIS FINLAND OY	EE

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
polümeerikattega tabletid				
CoDiovan® 160 mg/12,5 mg Filmtabletten	SE/H/0565/002	52884.00.00	NOVARTIS PHARMA GMBH	DE
CoDiovan® 160 mg/12,5 mg Filmtabletten	SE/H/0565/002	52884.00.00	NOVARTIS PHARMA GMBH	DE
CoDiovan® 160 mg/12,5 mg Filmtabletten	SE/H/0565/002	52884.00.00	NOVARTIS PHARMA GMBH	DE
CoDiovan® 160 mg/12,5 mg Filmtabletten	SE/H/0565/002	52884.00.00	NOVARTIS PHARMA GMBH	DE
Co-Diovan® 160/12.5 mg film-coated tablets	SE/H/0565/002	PL 00101/0650	NOVARTIS PHARMACEUTICALS UK LIMITED	UK
Co-Diovan® 160/12.5 mg film-coated tablets	SE/H/0565/002	PL 00101/0650	NOVARTIS PHARMACEUTICALS UK LIMITED	UK
Co-Diovan® 160/12.5 mg film-coated tablets	SE/H/0565/002	PL 00101/0650	NOVARTIS PHARMACEUTICALS UK LIMITED	UK
Co-Diovan® 160/12.5 mg film-coated tablets	SE/H/0565/002	PL 00101/0650	NOVARTIS PHARMACEUTICALS UK LIMITED	UK
Co-Diovan® 160/25 mg film-coated tablets	SE/H/0565/003	PL 00101/0651	NOVARTIS PHARMACEUTICALS UK LIMITED	UK
Co-Diovan® 160/25 mg film-coated tablets	SE/H/0565/003	PL 00101/0651	NOVARTIS PHARMACEUTICALS UK LIMITED	UK
Co-Diovan® 160/25 mg film-coated tablets	SE/H/0565/003	PL 00101/0651	NOVARTIS PHARMACEUTICALS UK LIMITED	UK
Co-Diovan® 160/25 mg film-coated tablets	SE/H/0565/003	PL 00101/0651	NOVARTIS PHARMACEUTICALS UK	UK

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
			LIMITED	
CoDiovan® 320 mg/12,5 mg Filmdabletten	SE/H/0565/004	69620.00.00	NOVARTIS PHARMA GMBH	DE
CoDiovan® 320 mg/12,5 mg Filmdabletten	SE/H/0565/004	69620.00.00	NOVARTIS PHARMA GMBH	DE
CoDiovan® 320 mg/12,5 mg Filmdabletten	SE/H/0565/004	69620.00.00	NOVARTIS PHARMA GMBH	DE
CoDiovan® 320 mg/12,5 mg Filmdabletten	SE/H/0565/004	69620.00.00	NOVARTIS PHARMA GMBH	DE
CoDiovan® 80 mg/12,5 mg Filmdabletten	SE/H/0565/001	40763.00.00	NOVARTIS PHARMA GMBH	DE
CoDiovan® 80 mg/12,5 mg Filmdabletten	SE/H/0565/001	40763.00.00	NOVARTIS PHARMA GMBH	DE
CoDiovan® 80 mg/12,5 mg Filmdabletten	SE/H/0565/001	40763.00.00	NOVARTIS PHARMA GMBH	DE
CoDiovan® 80 mg/12,5 mg Filmdabletten	SE/H/0565/001	40763.00.00	NOVARTIS PHARMA GMBH	DE
Co-Diovan® 80/12.5 mg film-coated tablets	SE/H/0565/001	PL 00101/0480	NOVARTIS PHARMACEUTICALS UK LIMITED	UK
Co-Diovan® 80/12.5 mg film-coated tablets	SE/H/0565/001	PL 00101/0480	NOVARTIS PHARMACEUTICALS UK LIMITED	UK
Co-Diovan® 80/12.5 mg film-coated tablets	SE/H/0565/001	PL 00101/0480	NOVARTIS PHARMACEUTICALS UK LIMITED	UK
Co-Diovan® 80/12.5 mg film-coated tablets	SE/H/0565/001	PL 00101/0480	NOVARTIS PHARMACEUTICALS UK LIMITED	UK
CoDiovan® forte 160 mg/25 mg Filmdabletten	SE/H/0565/003	52884.01.00	NOVARTIS PHARMA GMBH	DE
CoDiovan® forte 160 mg/25 mg Filmdabletten	SE/H/0565/003	52884.01.00	NOVARTIS 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
CoDiovan® forte 160 mg/25 mg Filmtabletten	SE/H/0565/003	52884.01.00	NOVARTIS PHARMA GMBH	DE
CoDiovan® forte 160 mg/25 mg Filmtabletten	SE/H/0565/003	52884.01.00	NOVARTIS PHARMA GMBH	DE
CoDiovan® forte 320 mg/25 mg Filmtabletten	SE/H/0565/005	69621.00.00	NOVARTIS PHARMA GMBH	DE
CoDiovan® forte 320 mg/25 mg Filmtabletten	SE/H/0565/005	69621.00.00	NOVARTIS PHARMA GMBH	DE
CoDiovan® forte 320 mg/25 mg Filmtabletten	SE/H/0565/005	69621.00.00	NOVARTIS PHARMA GMBH	DE
CoDiovan® forte 320 mg/25 mg Filmtabletten	SE/H/0565/005	69621.00.00	NOVARTIS PHARMA GMBH	DE
Co-Diothane 160 mg /12,5 mg Filmtabletten	SE/H/0565/002	BE258517	NOVARTIS PHARMA N.V.	BE
Co-Diothane 160 mg /12,5 mg Filmtabletten	SE/H/0565/002	BE258517	NOVARTIS PHARMA N.V.	BE
Co-Diothane 160 mg /12,5 mg Filmtabletten	SE/H/0565/002	BE258517	NOVARTIS PHARMA N.V.	BE
Co-Diothane 160 mg /12,5 mg Filmtabletten	SE/H/0565/002	BE258517	NOVARTIS PHARMA N.V.	BE
Co-Diothane 160 mg /12,5 mg Filmtabletten	SE/H/0565/002	2003100011	NOVARTIS PHARMA N.V.	LU
Co-Diothane 160 mg /12,5 mg Filmtabletten	SE/H/0565/002	2003100011	NOVARTIS PHARMA N.V.	LU
Co-Diothane 160 mg /12,5 mg Filmtabletten	SE/H/0565/002	2003100011	NOVARTIS PHARMA N.V.	LU
Co-Diothane 160 mg /12,5 mg Filmtabletten	SE/H/0565/002	2003100011	NOVARTIS PHARMA N.V.	LU
Co-Diothane 160 mg /25 mg Filmtabletten	SE/H/0565/003	BE271747	NOVARTIS PHARMA N.V.	BE
Co-Diothane 160 mg /25 mg Filmtabletten	SE/H/0565/003	BE271747	NOVARTIS PHARMA N.V.	BE
Co-Diothane 160 mg /25 mg Filmtabletten	SE/H/0565/003	BE271747	NOVARTIS PHARMA N.V.	BE

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
mg Filmtabletten				
Co-Diovane 160 mg /25 mg Filmtabletten	SE/H/0565/003	BE271747	NOVARTIS PHARMA N.V.	BE
Co-Diovane 160 mg /25 mg Filmtabletten	SE/H/0565/003	2005010014	NOVARTIS PHARMA N.V.	LU
Co-Diovane 160 mg /25 mg Filmtabletten	SE/H/0565/003	2005010014	NOVARTIS PHARMA N.V.	LU
Co-Diovane 160 mg /25 mg Filmtabletten	SE/H/0565/003	2005010014	NOVARTIS PHARMA N.V.	LU
Co-Diovane 160 mg /25 mg Filmtabletten	SE/H/0565/003	2005010014	NOVARTIS PHARMA N.V.	LU
Co-Diovane 160 mg/12,5 mg filmomhulde tabletten	SE/H/0565/002	BE258517	NOVARTIS PHARMA N.V.	BE
Co-Diovane 160 mg/12,5 mg filmomhulde tabletten	SE/H/0565/002	BE258517	NOVARTIS PHARMA N.V.	BE
Co-Diovane 160 mg/12,5 mg filmomhulde tabletten	SE/H/0565/002	BE258517	NOVARTIS PHARMA N.V.	BE
Co-Diovane 160 mg/12,5 mg filmomhulde tabletten	SE/H/0565/002	BE258517	NOVARTIS PHARMA N.V.	BE
Co-Diovane 160 mg/12,5 mg, comprimés pelliculés	SE/H/0565/002	BE258517	NOVARTIS PHARMA N.V.	BE
Co-Diovane 160 mg/12,5 mg, comprimés pelliculés	SE/H/0565/002	BE258517	NOVARTIS PHARMA N.V.	BE
Co-Diovane 160 mg/12,5 mg, comprimés pelliculés	SE/H/0565/002	BE258517	NOVARTIS PHARMA N.V.	BE
Co-Diovane 160 mg/12,5 mg, comprimés pelliculés	SE/H/0565/002	BE258517	NOVARTIS PHARMA N.V.	BE
CO-DIOVANE 160 MG/12,5 MG,	SE/H/0565/002	2003100011	NOVARTIS PHARMA N.V.	LU

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
COMPRIMÉS PELLICULÉS				
CO-DIOVANE 160 MG/12,5 MG, COMPRIMÉS PELLICULÉS	SE/H/0565/002	2003100011	NOVARTIS PHARMA N.V.	LU
CO-DIOVANE 160 MG/12,5 MG, COMPRIMÉS PELLICULÉS	SE/H/0565/002	2003100011	NOVARTIS PHARMA N.V.	LU
CO-DIOVANE 160 MG/12,5 MG, COMPRIMÉS PELLICULÉS	SE/H/0565/002	2003100011	NOVARTIS PHARMA N.V.	LU
Co-Diovane 160 mg/25 mg filmomhulde tabletten	SE/H/0565/003	BE271747	NOVARTIS PHARMA N.V.	BE
Co-Diovane 160 mg/25 mg filmomhulde tabletten	SE/H/0565/003	BE271747	NOVARTIS PHARMA N.V.	BE
Co-Diovane 160 mg/25 mg filmomhulde tabletten	SE/H/0565/003	BE271747	NOVARTIS PHARMA N.V.	BE
Co-Diovane 160 mg/25 mg filmomhulde tabletten	SE/H/0565/003	BE271747	NOVARTIS PHARMA N.V.	BE
Co-Diovane 160 mg/25 mg, comprimés pelliculés	SE/H/0565/003	BE271747	NOVARTIS PHARMA N.V.	BE
Co-Diovane 160 mg/25 mg, comprimés pelliculés	SE/H/0565/003	BE271747	NOVARTIS PHARMA N.V.	BE
Co-Diovane 160 mg/25 mg, comprimés pelliculés	SE/H/0565/003	BE271747	NOVARTIS PHARMA N.V.	BE
Co-Diovane 160 mg/25 mg, comprimés pelliculés	SE/H/0565/003	BE271747	NOVARTIS PHARMA N.V.	BE
CO-DIOVANE 160 MG/25 MG, COMPRIMÉS PELLICULÉS	SE/H/0565/003	2005010014	NOVARTIS PHARMA N.V.	LU

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
CO-DIOVANE 160 MG/25 MG, COMPRIMÉS PELLICULÉS	SE/H/0565/003	2005010014	NOVARTIS PHARMA N.V.	LU
CO-DIOVANE 160 MG/25 MG, COMPRIMÉS PELLICULÉS	SE/H/0565/003	2005010014	NOVARTIS PHARMA N.V.	LU
CO-DIOVANE 160 MG/25 MG, COMPRIMÉS PELLICULÉS	SE/H/0565/003	2005010014	NOVARTIS PHARMA N.V.	LU
Co-Diovane 320 mg /12,5 mg Filmdabletten	SE/H/0565/004	BE329025	NOVARTIS PHARMA N.V.	BE
Co-Diovane 320 mg /12,5 mg Filmdabletten	SE/H/0565/004	BE329025	NOVARTIS PHARMA N.V.	BE
Co-Diovane 320 mg /12,5 mg Filmdabletten	SE/H/0565/004	BE329025	NOVARTIS PHARMA N.V.	BE
Co-Diovane 320 mg /12,5 mg Filmdabletten	SE/H/0565/004	BE329025	NOVARTIS PHARMA N.V.	BE
Co-Diovane 320 mg /12,5 mg Filmdabletten	SE/H/0565/004	2008010030	NOVARTIS PHARMA N.V.	LU
Co-Diovane 320 mg /12,5 mg Filmdabletten	SE/H/0565/004	2008010030	NOVARTIS PHARMA N.V.	LU
Co-Diovane 320 mg /12,5 mg Filmdabletten	SE/H/0565/004	2008010030	NOVARTIS PHARMA N.V.	LU
Co-Diovane 320 mg /12,5 mg Filmdabletten	SE/H/0565/004	2008010030	NOVARTIS PHARMA N.V.	LU
Co-Diovane 320 mg /25 mg Filmdabletten	SE/H/0565/005	BE329034	NOVARTIS PHARMA N.V.	BE
Co-Diovane 320 mg /25 mg Filmdabletten	SE/H/0565/005	BE329034	NOVARTIS PHARMA N.V.	BE
Co-Diovane 320 mg /25 mg Filmdabletten	SE/H/0565/005	BE329034	NOVARTIS PHARMA N.V.	BE
Co-Diovane 320 mg /25 mg Filmdabletten	SE/H/0565/005	BE329034	NOVARTIS PHARMA N.V.	BE

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
Co-Diovane 320 mg /25 mg Filmtabletten	SE/H/0565/005	2008010031	NOVARTIS PHARMA N.V.	LU
Co-Diovane 320 mg /25 mg Filmtabletten	SE/H/0565/005	2008010031	NOVARTIS PHARMA N.V.	LU
Co-Diovane 320 mg /25 mg Filmtabletten	SE/H/0565/005	2008010031	NOVARTIS PHARMA N.V.	LU
Co-Diovane 320 mg /25 mg Filmtabletten	SE/H/0565/005	2008010031	NOVARTIS PHARMA N.V.	LU
Co-Diovane 320 mg/12,5 mg filmomhulde tabletten	SE/H/0565/004	BE329025	NOVARTIS PHARMA N.V.	BE
Co-Diovane 320 mg/12,5 mg filmomhulde tabletten	SE/H/0565/004	BE329025	NOVARTIS PHARMA N.V.	BE
Co-Diovane 320 mg/12,5 mg filmomhulde tabletten	SE/H/0565/004	BE329025	NOVARTIS PHARMA N.V.	BE
Co-Diovane 320 mg/12,5 mg filmomhulde tabletten	SE/H/0565/004	BE329025	NOVARTIS PHARMA N.V.	BE
Co-Diovane 320 mg/12,5 mg, comprimés pelliculés	SE/H/0565/004	BE329025	NOVARTIS PHARMA N.V.	BE
Co-Diovane 320 mg/12,5 mg, comprimés pelliculés	SE/H/0565/004	BE329025	NOVARTIS PHARMA N.V.	BE
Co-Diovane 320 mg/12,5 mg, comprimés pelliculés	SE/H/0565/004	BE329025	NOVARTIS PHARMA N.V.	BE
Co-Diovane 320 mg/12,5 mg, comprimés pelliculés	SE/H/0565/004	BE329025	NOVARTIS PHARMA N.V.	BE
CO-DIOVANE 320 MG/12,5 MG, COMPRIMÉS PELLICULÉS	SE/H/0565/004	2008010030	NOVARTIS PHARMA N.V.	LU
CO-DIOVANE 320 MG/12,5 MG,	SE/H/0565/004	2008010030	NOVARTIS PHARMA N.V.	LU

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
COMPRIMÉS PELLICULÉS				
CO-DIOVANE 320 MG/12,5 MG, COMPRIMÉS PELLICULÉS	SE/H/0565/004	2008010030	NOVARTIS PHARMA N.V.	LU
CO-DIOVANE 320 MG/12,5 MG, COMPRIMÉS PELLICULÉS	SE/H/0565/004	2008010030	NOVARTIS PHARMA N.V.	LU
Co-Diovine 320 mg/25 mg filmomhulde tabletten	SE/H/0565/005	BE329034	NOVARTIS PHARMA N.V.	BE
Co-Diovine 320 mg/25 mg filmomhulde tabletten	SE/H/0565/005	BE329034	NOVARTIS PHARMA N.V.	BE
Co-Diovine 320 mg/25 mg filmomhulde tabletten	SE/H/0565/005	BE329034	NOVARTIS PHARMA N.V.	BE
Co-Diovine 320 mg/25 mg filmomhulde tabletten	SE/H/0565/005	BE329034	NOVARTIS PHARMA N.V.	BE
Co-Diovine 320 mg/25 mg, comprimés pelliculés	SE/H/0565/005	BE329034	NOVARTIS PHARMA N.V.	BE
Co-Diovine 320 mg/25 mg, comprimés pelliculés	SE/H/0565/005	BE329034	NOVARTIS PHARMA N.V.	BE
Co-Diovine 320 mg/25 mg, comprimés pelliculés	SE/H/0565/005	BE329034	NOVARTIS PHARMA N.V.	BE
Co-Diovine 320 mg/25 mg, comprimés pelliculés	SE/H/0565/005	BE329034	NOVARTIS PHARMA N.V.	BE
CO-DIOVANE 320 MG/25 MG, COMPRIMÉS PELLICULÉS	SE/H/0565/005	2008010031	NOVARTIS PHARMA N.V.	LU
CO-DIOVANE 320 MG/25 MG, COMPRIMÉS PELLICULÉS	SE/H/0565/005	2008010031	NOVARTIS PHARMA N.V.	LU

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
CO-DIOVANE 320 MG/25 MG, COMPRIMÉS PELLICULÉS	SE/H/0565/005	2008010031	NOVARTIS PHARMA N.V.	LU
CO-DIOVANE 320 MG/25 MG, COMPRIMÉS PELLICULÉS	SE/H/0565/005	2008010031	NOVARTIS PHARMA N.V.	LU
Co-Diovane 80 mg /12,5 mg Filmtabletten	SE/H/0565/001	BE192893	NOVARTIS PHARMA N.V.	BE
Co-Diovane 80 mg /12,5 mg Filmtabletten	SE/H/0565/001	BE192893	NOVARTIS PHARMA N.V.	BE
Co-Diovane 80 mg /12,5 mg Filmtabletten	SE/H/0565/001	BE192893	NOVARTIS PHARMA N.V.	BE
Co-Diovane 80 mg /12,5 mg Filmtabletten	SE/H/0565/001	BE192893	NOVARTIS PHARMA N.V.	BE
Co-Diovane 80 mg /12,5 mg Filmtabletten	SE/H/0565/001	2002107201	NOVARTIS PHARMA N.V.	LU
Co-Diovane 80 mg /12,5 mg Filmtabletten	SE/H/0565/001	2002107201	NOVARTIS PHARMA N.V.	LU
Co-Diovane 80 mg /12,5 mg Filmtabletten	SE/H/0565/001	2002107201	NOVARTIS PHARMA N.V.	LU
Co-Diovane 80 mg /12,5 mg Filmtabletten	SE/H/0565/001	2002107201	NOVARTIS PHARMA N.V.	LU
Co-Diovane 80 mg/12,5 mg filmomhulde tabletten	SE/H/0565/001	BE192893	NOVARTIS PHARMA N.V.	BE
Co-Diovane 80 mg/12,5 mg filmomhulde tabletten	SE/H/0565/001	BE192893	NOVARTIS PHARMA N.V.	BE
Co-Diovane 80 mg/12,5 mg filmomhulde tabletten	SE/H/0565/001	BE192893	NOVARTIS PHARMA N.V.	BE
Co-Diovane 80 mg/12,5 mg filmomhulde	SE/H/0565/001	BE192893	NOVARTIS PHARMA N.V.	BE

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
tabletten				
Co-Diovane 80 mg/12,5 mg, comprimés pelliculés	SE/H/0565/001	BE192893	NOVARTIS PHARMA N.V.	BE
Co-Diovane 80 mg/12,5 mg, comprimés pelliculés	SE/H/0565/001	BE192893	NOVARTIS PHARMA N.V.	BE
Co-Diovane 80 mg/12,5 mg, comprimés pelliculés	SE/H/0565/001	BE192893	NOVARTIS PHARMA N.V.	BE
Co-Diovane 80 mg/12,5 mg, comprimés pelliculés	SE/H/0565/001	BE192893	NOVARTIS PHARMA N.V.	BE
CO-DIOVANE 80 MG/12,5 MG, COMPRIMÉS PELLICULÉS	SE/H/0565/001	2002107201	NOVARTIS PHARMA N.V.	LU
CO-DIOVANE 80 MG/12,5 MG, COMPRIMÉS PELLICULÉS	SE/H/0565/001	2002107201	NOVARTIS PHARMA N.V.	LU
CO-DIOVANE 80 MG/12,5 MG, COMPRIMÉS PELLICULÉS	SE/H/0565/001	2002107201	NOVARTIS PHARMA N.V.	LU
CO-DIOVANE 80 MG/12,5 MG, COMPRIMÉS PELLICULÉS	SE/H/0565/001	2002107201	NOVARTIS PHARMA N.V.	LU
COMBISARTAN 160 mg/12,5 mg compresse rivestite con film	not available	034134078	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

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 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/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	034134155	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 160 mg/25 mg compresse rivestite con film	not available	034134142	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

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	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
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

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/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
COMBISARTAN 80 mg/12,5 mg compresse rivestite con film	not available	034134015	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
CORDINATE® 160 MG FILMTABLETTEN	SE/H/0407/002	50724.01.00	NOVARTIS PHARMA GMBH	DE
CORDINATE® 160 MG FILMTABLETTEN	SE/H/0407/002	50724.01.00	NOVARTIS PHARMA GMBH	DE
CORDINATE® 160 MG FILMTABLETTEN	SE/H/0407/002	50724.01.00	NOVARTIS PHARMA GMBH	DE
CORDINATE® 160 MG FILMTABLETTEN	SE/H/0407/002	50724.01.00	NOVARTIS PHARMA GMBH	DE
CORDINATE® 160 MG	SE/H/0407/002	50724.01.00	NOVARTIS 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
FILMTABLETTE				
CORDINATE® 160 MG FILMTABLETTE	SE/H/0407/002	50724.01.00	NOVARTIS PHARMA GMBH	DE
CORDINATE® 320 MG FILMTABLETTE	SE/H/0407/004	67299.00.00	NOVARTIS PHARMA GMBH	DE
CORDINATE® 320 MG FILMTABLETTE	SE/H/0407/004	67299.00.00	NOVARTIS PHARMA GMBH	DE
CORDINATE® 320 MG FILMTABLETTE	SE/H/0407/004	67299.00.00	NOVARTIS PHARMA GMBH	DE
CORDINATE® 320 MG FILMTABLETTE	SE/H/0407/004	67299.00.00	NOVARTIS PHARMA GMBH	DE
CORDINATE® 320 MG FILMTABLETTE	SE/H/0407/004	67299.00.00	NOVARTIS PHARMA GMBH	DE
CORDINATE® 320 MG FILMTABLETTE	SE/H/0407/004	67299.00.00	NOVARTIS PHARMA GMBH	DE
CORDINATE® 40 MG FILMTABLETTE	SE/H/0407/003	50724.02.00	NOVARTIS PHARMA GMBH	DE
CORDINATE® 40 MG FILMTABLETTE	SE/H/0407/003	50724.02.00	NOVARTIS PHARMA GMBH	DE
CORDINATE® 40 MG FILMTABLETTE	SE/H/0407/003	50724.02.00	NOVARTIS PHARMA GMBH	DE
CORDINATE® 40 MG FILMTABLETTE	SE/H/0407/003	50724.02.00	NOVARTIS PHARMA GMBH	DE
CORDINATE® 40 MG FILMTABLETTE	SE/H/0407/003	50724.02.00	NOVARTIS PHARMA GMBH	DE
CORDINATE® 40 MG FILMTABLETTE	SE/H/0407/003	50724.02.00	NOVARTIS PHARMA GMBH	DE
CORDINATE® 80 MG FILMTABLETTE	SE/H/0407/001	50724.00.00	NOVARTIS PHARMA GMBH	DE
CORDINATE® 80 MG FILMTABLETTE	SE/H/0407/001	50724.00.00	NOVARTIS PHARMA GMBH	DE
CORDINATE® 80 MG FILMTABLETTE	SE/H/0407/001	50724.00.00	NOVARTIS 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
CORDINATE® 80 MG FILMTABLETTEN	SE/H/0407/001	50724.00.00	NOVARTIS PHARMA GMBH	DE
CORDINATE® 80 MG FILMTABLETTEN	SE/H/0407/001	50724.00.00	NOVARTIS PHARMA GMBH	DE
CORDINATE® 80 MG FILMTABLETTEN	SE/H/0407/001	50724.00.00	NOVARTIS PHARMA GMBH	DE
Cordinate® plus 160 mg/12,5 mg Filmtabletten	SE/H/0566/002	52886.00.00	NOVARTIS PHARMA GMBH	DE
Cordinate® plus 160 mg/12,5 mg Filmtabletten	SE/H/0566/002	52886.00.00	NOVARTIS PHARMA GMBH	DE
Cordinate® plus 160 mg/12,5 mg Filmtabletten	SE/H/0566/002	52886.00.00	NOVARTIS PHARMA GMBH	DE
Cordinate® plus 320 mg/12,5 mg Filmtabletten	SE/H/0566/004	69622.00.00	NOVARTIS PHARMA GMBH	DE
Cordinate® plus 320 mg/12,5 mg Filmtabletten	SE/H/0566/004	69622.00.00	NOVARTIS PHARMA GMBH	DE
Cordinate® plus 320 mg/12,5 mg Filmtabletten	SE/H/0566/004	69622.00.00	NOVARTIS PHARMA GMBH	DE
Cordinate® plus 80 mg/12,5 mg Filmtabletten	SE/H/0566/001	44773.00.00	NOVARTIS PHARMA GMBH	DE
Cordinate® plus 80 mg/12,5 mg Filmtabletten	SE/H/0566/001	44773.00.00	NOVARTIS PHARMA GMBH	DE
Cordinate® plus 80 mg/12,5 mg Filmtabletten	SE/H/0566/001	44773.00.00	NOVARTIS 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
Cordinate® plus forte 160 mg/25 mg Filmdabletten	SE/H/0566/003	52886.01.00	NOVARTIS PHARMA GMBH	DE
Cordinate® plus forte 160 mg/25 mg Filmdabletten	SE/H/0566/003	52886.01.00	NOVARTIS PHARMA GMBH	DE
Cordinate® plus forte 160 mg/25 mg Filmdabletten	SE/H/0566/003	52886.01.00	NOVARTIS PHARMA GMBH	DE
Cordinate® plus forte 320 mg/25 mg Filmdabletten	SE/H/0566/005	69623.00.00	NOVARTIS PHARMA GMBH	DE
Cordinate® plus forte 320 mg/25 mg Filmdabletten	SE/H/0566/005	69623.00.00	NOVARTIS PHARMA GMBH	DE
Cordinate® plus forte 320 mg/25 mg Filmdabletten	SE/H/0566/005	69623.00.00	NOVARTIS PHARMA GMBH	DE
CORIXIL	IT/H/0244/001	034774012	SANDOZ S.P.A.	IT
CORIXIL	IT/H/0244/001	034774024	SANDOZ S.P.A.	IT
CORIXIL	IT/H/0244/002	034774048	SANDOZ S.P.A.	IT
CORIXIL	IT/H/0244/002	034774036	SANDOZ S.P.A.	IT
CORIXIL	IT/H/0244/002	034774051	SANDOZ S.P.A.	IT
CORIXIL	IT/H/0244/002	034774063	SANDOZ S.P.A.	IT
CORIXIL	IT/H/0244/002	034774075	SANDOZ S.P.A.	IT
CORIXIL	IT/H/0244/002	034774087	SANDOZ S.P.A.	IT
CORIXIL	IT/H/0244/002	034774099	SANDOZ S.P.A.	IT
CORIXIL	IT/H/0244/003	034774101	SANDOZ S.P.A.	IT
CORIXIL	IT/H/0244/003	034774113	SANDOZ S.P.A.	IT
CORIXIL	IT/H/0244/003	034774125	SANDOZ S.P.A.	IT
CORIXIL	IT/H/0244/003	034774137	SANDOZ S.P.A.	IT
CORIXIL	IT/H/0244/003	034774149	SANDOZ S.P.A.	IT
CORIXIL	IT/H/0244/003	034774152	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
CORIXIL	IT/H/0244/003	034774164	SANDOZ S.P.A.	IT
CORIXIL	IT/H/0244/004	034774176	SANDOZ S.P.A.	IT
CORIXIL	IT/H/0244/004	034774188	SANDOZ S.P.A.	IT
CORIXIL	IT/H/0244/004	034774190	SANDOZ S.P.A.	IT
CORIXIL	IT/H/0244/004	034774202	SANDOZ S.P.A.	IT
CORIXIL	IT/H/0244/004	034774214	SANDOZ S.P.A.	IT
CORIXIL	IT/H/0244/004	034774226	SANDOZ S.P.A.	IT
CORIXIL	IT/H/0244/004	034774238	SANDOZ S.P.A.	IT
CORIXIL	IT/H/0244/004	034774240	SANDOZ S.P.A.	IT
CORIXIL	IT/H/0244/004	034774341	SANDOZ S.P.A.	IT
CORIXIL	IT/H/0244/005	034774253	SANDOZ S.P.A.	IT
CORIXIL	IT/H/0244/005	034774265	SANDOZ S.P.A.	IT
CORIXIL	IT/H/0244/005	034774277	SANDOZ S.P.A.	IT
CORIXIL	IT/H/0244/005	034774289	SANDOZ S.P.A.	IT
CORIXIL	IT/H/0244/005	034774303	SANDOZ S.P.A.	IT
CORIXIL	IT/H/0244/005	034774315	SANDOZ S.P.A.	IT
CORIXIL	IT/H/0244/005	034774327	SANDOZ S.P.A.	IT
CORIXIL	IT/H/0244/005	034774339	SANDOZ S.P.A.	IT
CORIXIL	IT/H/0244/005	034774354	SANDOZ S.P.A.	IT
CORIXIL	IT/H/0244/001	034774024	SANDOZ S.P.A.	IT
CORIXIL	IT/H/0244/002	034774036	SANDOZ S.P.A.	IT
CORIXIL	IT/H/0244/002	034774048	SANDOZ S.P.A.	IT
CORIXIL	IT/H/0244/002	034774075	SANDOZ S.P.A.	IT
CORIXIL	IT/H/0244/002	034774099	SANDOZ S.P.A.	IT
CORIXIL	IT/H/0244/002	034774087	SANDOZ S.P.A.	IT
CORIXIL	IT/H/0244/003	034774125	SANDOZ S.P.A.	IT
CORIXIL	IT/H/0244/003	034774137	SANDOZ S.P.A.	IT
CORIXIL	IT/H/0244/003	034774149	SANDOZ S.P.A.	IT
CORIXIL	IT/H/0244/004	034774202	SANDOZ S.P.A.	IT
CORIXIL	IT/H/0244/004	034774214	SANDOZ S.P.A.	IT
CORIXIL	IT/H/0244/004	034774238	SANDOZ S.P.A.	IT
CORIXIL	IT/H/0244/004	034774341	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
CORIXIL	IT/H/0244/001	034774012	SANDOZ S.P.A.	IT
CORIXIL	IT/H/0244/002	034774051	SANDOZ S.P.A.	IT
CORIXIL	IT/H/0244/003	034774101	SANDOZ S.P.A.	IT
CORIXIL	IT/H/0244/002	034774063	SANDOZ S.P.A.	IT
CORIXIL	IT/H/0244/003	034774113	SANDOZ S.P.A.	IT
CORIXIL	IT/H/0244/003	034774152	SANDOZ S.P.A.	IT
CORIXIL	IT/H/0244/003	034774164	SANDOZ S.P.A.	IT
CORIXIL	IT/H/0244/004	034774176	SANDOZ S.P.A.	IT
CORIXIL	IT/H/0244/004	034774190	SANDOZ S.P.A.	IT
CORIXIL	IT/H/0244/004	034774226	SANDOZ S.P.A.	IT
CORIXIL	IT/H/0244/004	034774188	SANDOZ S.P.A.	IT
CORIXIL	IT/H/0244/004	034774240	SANDOZ S.P.A.	IT
CORIXIL	IT/H/0244/005	034774265	SANDOZ S.P.A.	IT
CORIXIL	IT/H/0244/005	034774253	SANDOZ S.P.A.	IT
CORIXIL	IT/H/0244/005	034774277	SANDOZ S.P.A.	IT
CORIXIL	IT/H/0244/005	034774289	SANDOZ S.P.A.	IT
CORIXIL	IT/H/0244/005	034774303	SANDOZ S.P.A.	IT
CORIXIL	IT/H/0244/005	034774315	SANDOZ S.P.A.	IT
CORIXIL	IT/H/0244/005	034774327	SANDOZ S.P.A.	IT
CORIXIL	IT/H/0244/005	034774339	SANDOZ S.P.A.	IT
CORIXIL	IT/H/0244/005	034774354	SANDOZ S.P.A.	IT
Co-Tareg 160 mg /12,5 mg comprimidos revestidos por película	IT/H/0244/002	4850988	LABORATORIO NORMAL-PRODUTOS FARMACEUTICOS, LDA	PT
Co-Tareg 160 mg /12,5 mg comprimidos revestidos por película	IT/H/0244/002	4851184	LABORATORIO NORMAL-PRODUTOS FARMACEUTICOS, LDA	PT
Co-Tareg 160 mg /12,5 mg comprimidos revestidos por película	IT/H/0244/002	4851283	LABORATORIO NORMAL-PRODUTOS FARMACEUTICOS, LDA	PT
Co-Tareg 160 mg /12,5 mg comprimidos revestidos por película	IT/H/0244/002	4851085	LABORATORIO NORMAL-PRODUTOS FARMACEUTICOS, LDA	PT

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
Co-Tareg 160 mg /25 mg comprimidos revestidos por película	IT/H/0244/003	5456090	LABORATORIO NORMAL-PRODUTOS FARMACEUTICOS, LDA	PT
Co-Tareg 160 mg /25 mg comprimidos revestidos por película	IT/H/0244/003	5456199	LABORATORIO NORMAL-PRODUTOS FARMACEUTICOS, LDA	PT
Co-Tareg 160 mg /25 mg comprimidos revestidos por película	IT/H/0244/003	5456298	LABORATORIO NORMAL-PRODUTOS FARMACEUTICOS, LDA	PT
Co-Tareg 160 mg /25 mg comprimidos revestidos por película	IT/H/0244/003	5456397	LABORATORIO NORMAL-PRODUTOS FARMACEUTICOS, LDA	PT
Co-Tareg 160 mg /25 mg comprimidos revestidos por película	IT/H/0244/003	5456496	LABORATORIO NORMAL-PRODUTOS FARMACEUTICOS, LDA	PT
Co-Tareg 160 mg /25 mg comprimidos revestidos por película	IT/H/0244/003	5456595	LABORATORIO NORMAL-PRODUTOS FARMACEUTICOS, LDA	PT
Co-Tareg 160 mg /25 mg comprimidos revestidos por película	IT/H/0244/003	5456694	LABORATORIO NORMAL-PRODUTOS FARMACEUTICOS, LDA	PT
Cotareg 160 mg/12,5 mg compresse rivestite con film	SE/H/0565/002	034114417	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 160 mg/12,5 mg compresse rivestite con film	SE/H/0565/002	034114429	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 160 mg/12,5 mg compresse rivestite con film	SE/H/0565/002	034114506	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 160 mg/12,5 mg compresse rivestite con film	SE/H/0565/002	034114518	NOVARTIS EUROPHARM LIMITED	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
Cotareg 160 mg/12,5 mg compresse rivestite con film	SE/H/0565/002	034114520	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 160 mg/12,5 mg compresse rivestite con film	SE/H/0565/002	034114532	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 160 mg/12,5 mg compresse rivestite con film	SE/H/0565/002	034114544	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 160 mg/12,5 mg compresse rivestite con film	SE/H/0565/002	034114557	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 160 mg/12,5 mg compresse rivestite con film	SE/H/0565/002	034114569	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 160 mg/12,5 mg compresse rivestite con film	SE/H/0565/002	034114571	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 160 mg/12,5 mg compresse rivestite con film	SE/H/0565/002	034114064	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 160 mg/12,5 mg compresse rivestite con film	SE/H/0565/002	034114076	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 160 mg/12,5 mg compresse rivestite con film	SE/H/0565/002	034114088	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 160 mg/12,5 mg compresse rivestite con film	SE/H/0565/002	034114090	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 160 mg/12,5 mg compresse rivestite con film	SE/H/0565/002	034114102	NOVARTIS EUROPHARM LIMITED	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
Cotareg 160 mg/12,5 mg compresse rivestite con film	SE/H/0565/002	034114114	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 160 mg/12,5 mg compresse rivestite con film	SE/H/0565/002	034114126	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 160 mg/12,5 mg compresse rivestite con film	SE/H/0565/002	034114417	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 160 mg/12,5 mg compresse rivestite con film	SE/H/0565/002	034114429	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 160 mg/12,5 mg compresse rivestite con film	SE/H/0565/002	034114506	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 160 mg/12,5 mg compresse rivestite con film	SE/H/0565/002	034114518	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 160 mg/12,5 mg compresse rivestite con film	SE/H/0565/002	034114520	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 160 mg/12,5 mg compresse rivestite con film	SE/H/0565/002	034114532	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 160 mg/12,5 mg compresse rivestite con film	SE/H/0565/002	034114544	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 160 mg/12,5 mg compresse rivestite con film	SE/H/0565/002	034114557	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 160 mg/12,5 mg compresse rivestite con film	SE/H/0565/002	034114569	NOVARTIS EUROPHARM LIMITED	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
Cotareg 160 mg/12,5 mg compresse rivestite con film	SE/H/0565/002	034114571	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 160 mg/12,5 mg compresse rivestite con film	SE/H/0565/002	034114064	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 160 mg/12,5 mg compresse rivestite con film	SE/H/0565/002	034114076	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 160 mg/12,5 mg compresse rivestite con film	SE/H/0565/002	034114088	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 160 mg/12,5 mg compresse rivestite con film	SE/H/0565/002	034114090	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 160 mg/12,5 mg compresse rivestite con film	SE/H/0565/002	034114102	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 160 mg/12,5 mg compresse rivestite con film	SE/H/0565/002	034114114	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 160 mg/12,5 mg compresse rivestite con film	SE/H/0565/002	034114126	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 160 mg/12,5 mg compresse rivestite con film	SE/H/0565/002	034114417	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 160 mg/12,5 mg compresse rivestite con film	SE/H/0565/002	034114429	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 160 mg/12,5 mg compresse rivestite con film	SE/H/0565/002	034114506	NOVARTIS EUROPHARM LIMITED	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
Cotareg 160 mg/12,5 mg compresse rivestite con film	SE/H/0565/002	034114518	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 160 mg/12,5 mg compresse rivestite con film	SE/H/0565/002	034114520	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 160 mg/12,5 mg compresse rivestite con film	SE/H/0565/002	034114532	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 160 mg/12,5 mg compresse rivestite con film	SE/H/0565/002	034114544	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 160 mg/12,5 mg compresse rivestite con film	SE/H/0565/002	034114557	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 160 mg/12,5 mg compresse rivestite con film	SE/H/0565/002	034114569	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 160 mg/12,5 mg compresse rivestite con film	SE/H/0565/002	034114571	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 160 mg/12,5 mg compresse rivestite con film	SE/H/0565/002	034114064	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 160 mg/12,5 mg compresse rivestite con film	SE/H/0565/002	034114076	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 160 mg/12,5 mg compresse rivestite con film	SE/H/0565/002	034114088	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 160 mg/12,5 mg compresse rivestite con film	SE/H/0565/002	034114090	NOVARTIS EUROPHARM LIMITED	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
Cotareg 160 mg/12,5 mg compresse rivestite con film	SE/H/0565/002	034114102	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 160 mg/12,5 mg compresse rivestite con film	SE/H/0565/002	034114114	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 160 mg/12,5 mg compresse rivestite con film	SE/H/0565/002	034114126	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 160 mg/12,5 mg compresse rivestite con film	SE/H/0565/002	034114417	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 160 mg/12,5 mg compresse rivestite con film	SE/H/0565/002	034114429	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 160 mg/12,5 mg compresse rivestite con film	SE/H/0565/002	034114506	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 160 mg/12,5 mg compresse rivestite con film	SE/H/0565/002	034114518	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 160 mg/12,5 mg compresse rivestite con film	SE/H/0565/002	034114520	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 160 mg/12,5 mg compresse rivestite con film	SE/H/0565/002	034114532	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 160 mg/12,5 mg compresse rivestite con film	SE/H/0565/002	034114544	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 160 mg/12,5 mg compresse rivestite con film	SE/H/0565/002	034114557	NOVARTIS EUROPHARM LIMITED	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
Cotareg 160 mg/12,5 mg compresse rivestite con film	SE/H/0565/002	034114569	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 160 mg/12,5 mg compresse rivestite con film	SE/H/0565/002	034114571	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 160 mg/12,5 mg compresse rivestite con film	SE/H/0565/002	034114064	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 160 mg/12,5 mg compresse rivestite con film	SE/H/0565/002	034114076	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 160 mg/12,5 mg compresse rivestite con film	SE/H/0565/002	034114088	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 160 mg/12,5 mg compresse rivestite con film	SE/H/0565/002	034114090	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 160 mg/12,5 mg compresse rivestite con film	SE/H/0565/002	034114102	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 160 mg/12,5 mg compresse rivestite con film	SE/H/0565/002	034114114	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 160 mg/12,5 mg compresse rivestite con film	SE/H/0565/002	034114126	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 160 mg/12,5 mg, comprimé pelliculé	SE/H/0565/002	3400936310728	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/12,5 mg, comprimé pelliculé	SE/H/0565/002	3400937125406	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/12,5 mg, comprimé pelliculé	SE/H/0565/002	3400937201377	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
COTAREG 160 mg/12,5 mg, comprimé pelliculé	SE/H/0565/002	3400937201438	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/12,5 mg, comprimé pelliculé	SE/H/0565/002	3400938156591	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/12,5 mg, comprimé pelliculé	SE/H/0565/002	3400938156652	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/12,5 mg, comprimé pelliculé	SE/H/0565/002	3400938156713	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/12,5 mg, comprimé pelliculé	SE/H/0565/002	3400938156881	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/12,5 mg, comprimé pelliculé	SE/H/0565/002	3400938156942	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/12,5 mg, comprimé pelliculé	SE/H/0565/002	3400938157024	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/12,5 mg, comprimé pelliculé	SE/H/0565/002	3400938157192	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/12,5 mg, comprimé pelliculé	SE/H/0565/002	3400956497683	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/12,5 mg, comprimé pelliculé	SE/H/0565/002	3400956497744	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/12,5 mg, comprimé pelliculé	SE/H/0565/002	3400956497805	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/12,5 mg, comprimé pelliculé	SE/H/0565/002	3400936310728	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/12,5 mg, comprimé pelliculé	SE/H/0565/002	3400937125406	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/12,5 mg, comprimé pelliculé	SE/H/0565/002	3400937201377	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/12,5 mg, comprimé pelliculé	SE/H/0565/002	3400937201438	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/12,5 mg, comprimé pelliculé	SE/H/0565/002	3400938156591	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/12,5 mg, comprimé pelliculé	SE/H/0565/002	3400938156652	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
mg, comprimé pelliculé				
COTAREG 160 mg/12,5 mg, comprimé pelliculé	SE/H/0565/002	3400938156713	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/12,5 mg, comprimé pelliculé	SE/H/0565/002	3400938156881	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/12,5 mg, comprimé pelliculé	SE/H/0565/002	3400938156942	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/12,5 mg, comprimé pelliculé	SE/H/0565/002	3400938157024	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/12,5 mg, comprimé pelliculé	SE/H/0565/002	3400938157192	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/12,5 mg, comprimé pelliculé	SE/H/0565/002	3400956497683	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/12,5 mg, comprimé pelliculé	SE/H/0565/002	3400956497744	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/12,5 mg, comprimé pelliculé	SE/H/0565/002	3400956497805	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/12,5 mg, comprimé pelliculé	SE/H/0565/002	3400936310728	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/12,5 mg, comprimé pelliculé	SE/H/0565/002	3400937125406	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/12,5 mg, comprimé pelliculé	SE/H/0565/002	3400937201377	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/12,5 mg, comprimé pelliculé	SE/H/0565/002	3400937201438	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/12,5 mg, comprimé pelliculé	SE/H/0565/002	3400938156591	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/12,5 mg, comprimé pelliculé	SE/H/0565/002	3400938156652	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/12,5 mg, comprimé pelliculé	SE/H/0565/002	3400938156713	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/12,5 mg, comprimé pelliculé	SE/H/0565/002	3400938156881	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
COTAREG 160 mg/12,5 mg, comprimé pelliculé	SE/H/0565/002	3400938156942	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/12,5 mg, comprimé pelliculé	SE/H/0565/002	3400938157024	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/12,5 mg, comprimé pelliculé	SE/H/0565/002	3400938157192	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/12,5 mg, comprimé pelliculé	SE/H/0565/002	3400956497683	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/12,5 mg, comprimé pelliculé	SE/H/0565/002	3400956497744	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/12,5 mg, comprimé pelliculé	SE/H/0565/002	3400956497805	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/12,5 mg, comprimé pelliculé	SE/H/0565/002	3400936310728	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/12,5 mg, comprimé pelliculé	SE/H/0565/002	3400937125406	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/12,5 mg, comprimé pelliculé	SE/H/0565/002	3400937201377	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/12,5 mg, comprimé pelliculé	SE/H/0565/002	3400937201438	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/12,5 mg, comprimé pelliculé	SE/H/0565/002	3400938156591	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/12,5 mg, comprimé pelliculé	SE/H/0565/002	3400938156652	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/12,5 mg, comprimé pelliculé	SE/H/0565/002	3400938156713	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/12,5 mg, comprimé pelliculé	SE/H/0565/002	3400938156881	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/12,5 mg, comprimé pelliculé	SE/H/0565/002	3400938156942	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/12,5 mg, comprimé pelliculé	SE/H/0565/002	3400938157024	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/12,5 mg, comprimé pelliculé	SE/H/0565/002	3400938157192	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
mg, comprimé pelliculé				
COTAREG 160 mg/12,5 mg, comprimé pelliculé	SE/H/0565/002	3400956497683	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/12,5 mg, comprimé pelliculé	SE/H/0565/002	3400956497744	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/12,5 mg, comprimé pelliculé	SE/H/0565/002	3400956497805	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/25 mg compresse rivestite con film	SE/H/0565/003	034114138	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 160 mg/25 mg compresse rivestite con film	SE/H/0565/003	034114140	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 160 mg/25 mg compresse rivestite con film	SE/H/0565/003	034114153	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 160 mg/25 mg compresse rivestite con film	SE/H/0565/003	034114165	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 160 mg/25 mg compresse rivestite con film	SE/H/0565/003	034114177	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 160 mg/25 mg compresse rivestite con film	SE/H/0565/003	034114189	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 160 mg/25 mg compresse rivestite con film	SE/H/0565/003	034114431	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 160 mg/25 mg compresse rivestite con film	SE/H/0565/003	034114443	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 160 mg/25 mg compresse rivestite con	SE/H/0565/003	034114583	NOVARTIS EUROPHARM LIMITED	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
film				
COTAREG 160 mg/25 mg compresse rivestite con film	SE/H/0565/003	034114595	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 160 mg/25 mg compresse rivestite con film	SE/H/0565/003	034114607	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 160 mg/25 mg compresse rivestite con film	SE/H/0565/003	034114619	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 160 mg/25 mg compresse rivestite con film	SE/H/0565/003	034114621	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 160 mg/25 mg compresse rivestite con film	SE/H/0565/003	034114633	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 160 mg/25 mg compresse rivestite con film	SE/H/0565/003	034114645	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 160 mg/25 mg compresse rivestite con film	SE/H/0565/003	034114658	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 160 mg/25 mg compresse rivestite con film	SE/H/0565/003	034114191	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 160 mg/25 mg compresse rivestite con film	SE/H/0565/003	034114138	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 160 mg/25 mg compresse rivestite con film	SE/H/0565/003	034114140	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 160 mg/25 mg compresse rivestite con	SE/H/0565/003	034114153	NOVARTIS EUROPHARM LIMITED	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
film				
COTAREG 160 mg/25 mg compresse rivestite con film	SE/H/0565/003	034114165	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 160 mg/25 mg compresse rivestite con film	SE/H/0565/003	034114177	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 160 mg/25 mg compresse rivestite con film	SE/H/0565/003	034114189	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 160 mg/25 mg compresse rivestite con film	SE/H/0565/003	034114431	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 160 mg/25 mg compresse rivestite con film	SE/H/0565/003	034114443	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 160 mg/25 mg compresse rivestite con film	SE/H/0565/003	034114583	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 160 mg/25 mg compresse rivestite con film	SE/H/0565/003	034114595	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 160 mg/25 mg compresse rivestite con film	SE/H/0565/003	034114607	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 160 mg/25 mg compresse rivestite con film	SE/H/0565/003	034114619	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 160 mg/25 mg compresse rivestite con film	SE/H/0565/003	034114621	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 160 mg/25 mg compresse rivestite con	SE/H/0565/003	034114633	NOVARTIS EUROPHARM LIMITED	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
film				
COTAREG 160 mg/25 mg compresse rivestite con film	SE/H/0565/003	034114645	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 160 mg/25 mg compresse rivestite con film	SE/H/0565/003	034114658	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 160 mg/25 mg compresse rivestite con film	SE/H/0565/003	034114191	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 160 mg/25 mg compresse rivestite con film	SE/H/0565/003	034114138	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 160 mg/25 mg compresse rivestite con film	SE/H/0565/003	034114140	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 160 mg/25 mg compresse rivestite con film	SE/H/0565/003	034114153	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 160 mg/25 mg compresse rivestite con film	SE/H/0565/003	034114165	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 160 mg/25 mg compresse rivestite con film	SE/H/0565/003	034114177	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 160 mg/25 mg compresse rivestite con film	SE/H/0565/003	034114189	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 160 mg/25 mg compresse rivestite con film	SE/H/0565/003	034114431	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 160 mg/25 mg compresse rivestite con	SE/H/0565/003	034114443	NOVARTIS EUROPHARM LIMITED	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
film				
COTAREG 160 mg/25 mg compresse rivestite con film	SE/H/0565/003	034114583	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 160 mg/25 mg compresse rivestite con film	SE/H/0565/003	034114595	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 160 mg/25 mg compresse rivestite con film	SE/H/0565/003	034114607	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 160 mg/25 mg compresse rivestite con film	SE/H/0565/003	034114619	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 160 mg/25 mg compresse rivestite con film	SE/H/0565/003	034114621	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 160 mg/25 mg compresse rivestite con film	SE/H/0565/003	034114633	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 160 mg/25 mg compresse rivestite con film	SE/H/0565/003	034114645	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 160 mg/25 mg compresse rivestite con film	SE/H/0565/003	034114658	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 160 mg/25 mg compresse rivestite con film	SE/H/0565/003	034114191	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 160 mg/25 mg compresse rivestite con film	SE/H/0565/003	034114138	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 160 mg/25 mg compresse rivestite con	SE/H/0565/003	034114140	NOVARTIS EUROPHARM LIMITED	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
film				
COTAREG 160 mg/25 mg compresse rivestite con film	SE/H/0565/003	034114153	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 160 mg/25 mg compresse rivestite con film	SE/H/0565/003	034114165	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 160 mg/25 mg compresse rivestite con film	SE/H/0565/003	034114177	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 160 mg/25 mg compresse rivestite con film	SE/H/0565/003	034114189	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 160 mg/25 mg compresse rivestite con film	SE/H/0565/003	034114431	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 160 mg/25 mg compresse rivestite con film	SE/H/0565/003	034114443	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 160 mg/25 mg compresse rivestite con film	SE/H/0565/003	034114583	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 160 mg/25 mg compresse rivestite con film	SE/H/0565/003	034114595	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 160 mg/25 mg compresse rivestite con film	SE/H/0565/003	034114607	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 160 mg/25 mg compresse rivestite con film	SE/H/0565/003	034114619	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 160 mg/25 mg compresse rivestite con	SE/H/0565/003	034114621	NOVARTIS EUROPHARM LIMITED	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
film				
COTAREG 160 mg/25 mg compresse rivestite con film	SE/H/0565/003	034114633	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 160 mg/25 mg compresse rivestite con film	SE/H/0565/003	034114645	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 160 mg/25 mg compresse rivestite con film	SE/H/0565/003	034114658	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 160 mg/25 mg compresse rivestite con film	SE/H/0565/003	034114191	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 160 mg/25 mg, comprimé pelliculé	SE/H/0565/003	3400935923202	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/25 mg, comprimé pelliculé	SE/H/0565/003	3400937139472	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/25 mg, comprimé pelliculé	SE/H/0565/003	3400937139533	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/25 mg, comprimé pelliculé	SE/H/0565/003	3400937139762	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/25 mg, comprimé pelliculé	SE/H/0565/003	3400938157253	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/25 mg, comprimé pelliculé	SE/H/0565/003	3400938157314	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/25 mg, comprimé pelliculé	SE/H/0565/003	3400938157482	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/25 mg, comprimé pelliculé	SE/H/0565/003	3400938157543	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/25 mg, comprimé pelliculé	SE/H/0565/003	3400938157604	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/25 mg, comprimé pelliculé	SE/H/0565/003	3400938157772	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
COTAREG 160 mg/25 mg, comprimé pelliculé	SE/H/0565/003	3400938157833	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/25 mg, comprimé pelliculé	SE/H/0565/003	3400956368372	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/25 mg, comprimé pelliculé	SE/H/0565/003	3400956368433	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/25 mg, comprimé pelliculé	SE/H/0565/003	3400956368662	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/25 mg, comprimé pelliculé	SE/H/0565/003	3400935923202	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/25 mg, comprimé pelliculé	SE/H/0565/003	3400937139472	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/25 mg, comprimé pelliculé	SE/H/0565/003	3400937139533	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/25 mg, comprimé pelliculé	SE/H/0565/003	3400937139762	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/25 mg, comprimé pelliculé	SE/H/0565/003	3400938157253	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/25 mg, comprimé pelliculé	SE/H/0565/003	3400938157314	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/25 mg, comprimé pelliculé	SE/H/0565/003	3400938157482	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/25 mg, comprimé pelliculé	SE/H/0565/003	3400938157543	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/25 mg, comprimé pelliculé	SE/H/0565/003	3400938157604	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/25 mg, comprimé pelliculé	SE/H/0565/003	3400938157772	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/25 mg, comprimé pelliculé	SE/H/0565/003	3400938157833	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/25 mg, comprimé pelliculé	SE/H/0565/003	3400956368372	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/25 mg, comprimé pelliculé	SE/H/0565/003	3400956368433	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
mg, comprimé pelliculé				
COTAREG 160 mg/25 mg, comprimé pelliculé	SE/H/0565/003	3400956368662	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/25 mg, comprimé pelliculé	SE/H/0565/003	3400935923202	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/25 mg, comprimé pelliculé	SE/H/0565/003	3400937139472	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/25 mg, comprimé pelliculé	SE/H/0565/003	3400937139533	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/25 mg, comprimé pelliculé	SE/H/0565/003	3400937139762	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/25 mg, comprimé pelliculé	SE/H/0565/003	3400938157253	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/25 mg, comprimé pelliculé	SE/H/0565/003	3400938157314	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/25 mg, comprimé pelliculé	SE/H/0565/003	3400938157482	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/25 mg, comprimé pelliculé	SE/H/0565/003	3400938157543	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/25 mg, comprimé pelliculé	SE/H/0565/003	3400938157604	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/25 mg, comprimé pelliculé	SE/H/0565/003	3400938157772	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/25 mg, comprimé pelliculé	SE/H/0565/003	3400938157833	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/25 mg, comprimé pelliculé	SE/H/0565/003	3400956368372	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/25 mg, comprimé pelliculé	SE/H/0565/003	3400956368433	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/25 mg, comprimé pelliculé	SE/H/0565/003	3400956368662	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/25 mg, comprimé pelliculé	SE/H/0565/003	3400935923202	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
COTAREG 160 mg/25 mg, comprimé pelliculé	SE/H/0565/003	3400937139472	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/25 mg, comprimé pelliculé	SE/H/0565/003	3400937139533	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/25 mg, comprimé pelliculé	SE/H/0565/003	3400937139762	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/25 mg, comprimé pelliculé	SE/H/0565/003	3400938157253	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/25 mg, comprimé pelliculé	SE/H/0565/003	3400938157314	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/25 mg, comprimé pelliculé	SE/H/0565/003	3400938157482	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/25 mg, comprimé pelliculé	SE/H/0565/003	3400938157543	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/25 mg, comprimé pelliculé	SE/H/0565/003	3400938157604	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/25 mg, comprimé pelliculé	SE/H/0565/003	3400938157772	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/25 mg, comprimé pelliculé	SE/H/0565/003	3400938157833	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/25 mg, comprimé pelliculé	SE/H/0565/003	3400956368372	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/25 mg, comprimé pelliculé	SE/H/0565/003	3400956368433	NOVARTIS PHARMA S.A.S.	FR
COTAREG 160 mg/25 mg, comprimé pelliculé	SE/H/0565/003	3400956368662	NOVARTIS PHARMA S.A.S.	FR
COTAREG 320 mg/12,5 mg compresse rivestite con film	SE/H/0565/004	034114203	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 320 mg/12,5 mg compresse rivestite con film	SE/H/0565/004	034114215	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 320 mg/12,5	SE/H/0565/004	034114227	NOVARTIS EUROPHARM	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
mg compresse rivestite con film			LIMITED	
COTAREG 320 mg/12,5 mg compresse rivestite con film	SE/H/0565/004	034114239	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 320 mg/12,5 mg compresse rivestite con film	SE/H/0565/004	034114241	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 320 mg/12,5 mg compresse rivestite con film	SE/H/0565/004	034114254	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 320 mg/12,5 mg compresse rivestite con film	SE/H/0565/004	034114266	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 320 mg/12,5 mg compresse rivestite con film	SE/H/0565/004	034114278	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 320 mg/12,5 mg compresse rivestite con film	SE/H/0565/004	034114367	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 320 mg/12,5 mg compresse rivestite con film	SE/H/0565/004	034114203	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 320 mg/12,5 mg compresse rivestite con film	SE/H/0565/004	034114215	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 320 mg/12,5 mg compresse rivestite con film	SE/H/0565/004	034114227	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 320 mg/12,5 mg compresse rivestite con film	SE/H/0565/004	034114239	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 320 mg/12,5	SE/H/0565/004	034114241	NOVARTIS EUROPHARM	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
mg compresse rivestite con film			LIMITED	
COTAREG 320 mg/12,5 mg compresse rivestite con film	SE/H/0565/004	034114254	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 320 mg/12,5 mg compresse rivestite con film	SE/H/0565/004	034114266	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 320 mg/12,5 mg compresse rivestite con film	SE/H/0565/004	034114278	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 320 mg/12,5 mg compresse rivestite con film	SE/H/0565/004	034114367	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 320 mg/12,5 mg compresse rivestite con film	SE/H/0565/004	034114203	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 320 mg/12,5 mg compresse rivestite con film	SE/H/0565/004	034114215	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 320 mg/12,5 mg compresse rivestite con film	SE/H/0565/004	034114227	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 320 mg/12,5 mg compresse rivestite con film	SE/H/0565/004	034114239	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 320 mg/12,5 mg compresse rivestite con film	SE/H/0565/004	034114241	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 320 mg/12,5 mg compresse rivestite con film	SE/H/0565/004	034114254	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 320 mg/12,5	SE/H/0565/004	034114266	NOVARTIS EUROPHARM	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
mg compresse rivestite con film			LIMITED	
COTAREG 320 mg/12,5 mg compresse rivestite con film	SE/H/0565/004	034114278	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 320 mg/12,5 mg compresse rivestite con film	SE/H/0565/004	034114367	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 320 mg/12,5 mg compresse rivestite con film	SE/H/0565/004	034114203	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 320 mg/12,5 mg compresse rivestite con film	SE/H/0565/004	034114215	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 320 mg/12,5 mg compresse rivestite con film	SE/H/0565/004	034114227	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 320 mg/12,5 mg compresse rivestite con film	SE/H/0565/004	034114239	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 320 mg/12,5 mg compresse rivestite con film	SE/H/0565/004	034114241	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 320 mg/12,5 mg compresse rivestite con film	SE/H/0565/004	034114254	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 320 mg/12,5 mg compresse rivestite con film	SE/H/0565/004	034114266	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 320 mg/12,5 mg compresse rivestite con film	SE/H/0565/004	034114278	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 320 mg/12,5	SE/H/0565/004	034114367	NOVARTIS EUROPHARM	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
mg compresse rivestite con film			LIMITED	
COTAREG 320 mg/25 mg compresse rivestite con film	SE/H/0565/005	034114280	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 320 mg/25 mg compresse rivestite con film	SE/H/0565/005	034114292	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 320 mg/25 mg compresse rivestite con film	SE/H/0565/005	034114304	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 320 mg/25 mg compresse rivestite con film	SE/H/0565/005	034114316	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 320 mg/25 mg compresse rivestite con film	SE/H/0565/005	034114328	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 320 mg/25 mg compresse rivestite con film	SE/H/0565/005	034114330	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 320 mg/25 mg compresse rivestite con film	SE/H/0565/005	034114342	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 320 mg/25 mg compresse rivestite con film	SE/H/0565/005	034114355	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 320 mg/25 mg compresse rivestite con film	SE/H/0565/005	034114379	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 320 mg/25 mg compresse rivestite con film	SE/H/0565/005	034114280	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 320 mg/25 mg	SE/H/0565/005	034114292	NOVARTIS EUROPHARM	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			LIMITED	
COTAREG 320 mg/25 mg compresse rivestite con film	SE/H/0565/005	034114304	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 320 mg/25 mg compresse rivestite con film	SE/H/0565/005	034114316	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 320 mg/25 mg compresse rivestite con film	SE/H/0565/005	034114328	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 320 mg/25 mg compresse rivestite con film	SE/H/0565/005	034114330	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 320 mg/25 mg compresse rivestite con film	SE/H/0565/005	034114342	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 320 mg/25 mg compresse rivestite con film	SE/H/0565/005	034114355	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 320 mg/25 mg compresse rivestite con film	SE/H/0565/005	034114379	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 320 mg/25 mg compresse rivestite con film	SE/H/0565/005	034114280	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 320 mg/25 mg compresse rivestite con film	SE/H/0565/005	034114292	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 320 mg/25 mg compresse rivestite con film	SE/H/0565/005	034114304	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 320 mg/25 mg	SE/H/0565/005	034114316	NOVARTIS EUROPHARM	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			LIMITED	
COTAREG 320 mg/25 mg compresse rivestite con film	SE/H/0565/005	034114328	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 320 mg/25 mg compresse rivestite con film	SE/H/0565/005	034114330	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 320 mg/25 mg compresse rivestite con film	SE/H/0565/005	034114342	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 320 mg/25 mg compresse rivestite con film	SE/H/0565/005	034114355	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 320 mg/25 mg compresse rivestite con film	SE/H/0565/005	034114379	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 320 mg/25 mg compresse rivestite con film	SE/H/0565/005	034114280	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 320 mg/25 mg compresse rivestite con film	SE/H/0565/005	034114292	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 320 mg/25 mg compresse rivestite con film	SE/H/0565/005	034114304	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 320 mg/25 mg compresse rivestite con film	SE/H/0565/005	034114316	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 320 mg/25 mg compresse rivestite con film	SE/H/0565/005	034114328	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 320 mg/25 mg	SE/H/0565/005	034114330	NOVARTIS EUROPHARM	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			LIMITED	
COTAREG 320 mg/25 mg compresse rivestite con film	SE/H/0565/005	034114342	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 320 mg/25 mg compresse rivestite con film	SE/H/0565/005	034114355	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 320 mg/25 mg compresse rivestite con film	SE/H/0565/005	034114379	NOVARTIS EUROPHARM LIMITED	IT
Co-Tareg 80 mg /12,5 mg comprimidos revestidos por película	IT/H/0244/001	2807584	LABORATORIO NORMAL-PRODUTOS FARMACEUTICOS, LDA	PT
Co-Tareg 80 mg /12,5 mg comprimidos revestidos por película	IT/H/0244/001	2807683	LABORATORIO NORMAL-PRODUTOS FARMACEUTICOS, LDA	PT
Co-Tareg 80 mg /12,5 mg comprimidos revestidos por película	IT/H/0244/001	2807782	LABORATORIO NORMAL-PRODUTOS FARMACEUTICOS, LDA	PT
Cotareg 80 mg/12,5 mg compresse rivestite con film	SE/H/0565/001	034114037	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 80 mg/12,5 mg compresse rivestite con film	SE/H/0565/001	034114405	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 80 mg/12,5 mg compresse rivestite con film	SE/H/0565/001	034114393	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 80 mg/12,5 mg compresse rivestite con film	SE/H/0565/001	034114381	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 80 mg/12,5 mg	SE/H/0565/001	034114052	NOVARTIS EUROPHARM	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			LIMITED	
Cotareg 80 mg/12,5 mg compresse rivestite con film	SE/H/0565/001	034114049	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 80 mg/12,5 mg compresse rivestite con film	SE/H/0565/001	034114708	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 80 mg/12,5 mg compresse rivestite con film	SE/H/0565/001	034114672	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 80 mg/12,5 mg compresse rivestite con film	SE/H/0565/001	034114660	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 80 mg/12,5 mg compresse rivestite con film	SE/H/0565/001	034114494	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 80 mg/12,5 mg compresse rivestite con film	SE/H/0565/001	034114696	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 80 mg/12,5 mg compresse rivestite con film	SE/H/0565/001	034114684	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 80 mg/12,5 mg compresse rivestite con film	SE/H/0565/001	034114482	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 80 mg/12,5 mg compresse rivestite con film	SE/H/0565/001	034114470	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 80 mg/12,5 mg compresse rivestite con film	SE/H/0565/001	034114468	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 80 mg/12,5 mg	SE/H/0565/001	034114456	NOVARTIS EUROPHARM	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			LIMITED	
Cotareg 80 mg/12,5 mg compresse rivestite con film	SE/H/0565/001	034114025	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 80 mg/12,5 mg compresse rivestite con film	SE/H/0565/001	034114013	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 80 mg/12,5 mg compresse rivestite con film	SE/H/0565/001	034114037	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 80 mg/12,5 mg compresse rivestite con film	SE/H/0565/001	034114405	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 80 mg/12,5 mg compresse rivestite con film	SE/H/0565/001	034114393	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 80 mg/12,5 mg compresse rivestite con film	SE/H/0565/001	034114381	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 80 mg/12,5 mg compresse rivestite con film	SE/H/0565/001	034114052	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 80 mg/12,5 mg compresse rivestite con film	SE/H/0565/001	034114049	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 80 mg/12,5 mg compresse rivestite con film	SE/H/0565/001	034114708	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 80 mg/12,5 mg compresse rivestite con film	SE/H/0565/001	034114672	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 80 mg/12,5 mg	SE/H/0565/001	034114660	NOVARTIS EUROPHARM	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			LIMITED	
Cotareg 80 mg/12,5 mg compresse rivestite con film	SE/H/0565/001	034114494	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 80 mg/12,5 mg compresse rivestite con film	SE/H/0565/001	034114696	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 80 mg/12,5 mg compresse rivestite con film	SE/H/0565/001	034114684	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 80 mg/12,5 mg compresse rivestite con film	SE/H/0565/001	034114482	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 80 mg/12,5 mg compresse rivestite con film	SE/H/0565/001	034114470	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 80 mg/12,5 mg compresse rivestite con film	SE/H/0565/001	034114468	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 80 mg/12,5 mg compresse rivestite con film	SE/H/0565/001	034114456	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 80 mg/12,5 mg compresse rivestite con film	SE/H/0565/001	034114025	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 80 mg/12,5 mg compresse rivestite con film	SE/H/0565/001	034114013	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 80 mg/12,5 mg compresse rivestite con film	SE/H/0565/001	034114037	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 80 mg/12,5 mg	SE/H/0565/001	034114405	NOVARTIS EUROPHARM	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			LIMITED	
Cotareg 80 mg/12,5 mg compresse rivestite con film	SE/H/0565/001	034114393	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 80 mg/12,5 mg compresse rivestite con film	SE/H/0565/001	034114381	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 80 mg/12,5 mg compresse rivestite con film	SE/H/0565/001	034114052	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 80 mg/12,5 mg compresse rivestite con film	SE/H/0565/001	034114049	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 80 mg/12,5 mg compresse rivestite con film	SE/H/0565/001	034114708	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 80 mg/12,5 mg compresse rivestite con film	SE/H/0565/001	034114672	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 80 mg/12,5 mg compresse rivestite con film	SE/H/0565/001	034114660	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 80 mg/12,5 mg compresse rivestite con film	SE/H/0565/001	034114494	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 80 mg/12,5 mg compresse rivestite con film	SE/H/0565/001	034114696	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 80 mg/12,5 mg compresse rivestite con film	SE/H/0565/001	034114684	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 80 mg/12,5 mg	SE/H/0565/001	034114482	NOVARTIS EUROPHARM	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			LIMITED	
Cotareg 80 mg/12,5 mg compresse rivestite con film	SE/H/0565/001	034114470	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 80 mg/12,5 mg compresse rivestite con film	SE/H/0565/001	034114468	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 80 mg/12,5 mg compresse rivestite con film	SE/H/0565/001	034114456	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 80 mg/12,5 mg compresse rivestite con film	SE/H/0565/001	034114025	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 80 mg/12,5 mg compresse rivestite con film	SE/H/0565/001	034114013	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 80 mg/12,5 mg compresse rivestite con film	SE/H/0565/001	034114037	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 80 mg/12,5 mg compresse rivestite con film	SE/H/0565/001	034114405	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 80 mg/12,5 mg compresse rivestite con film	SE/H/0565/001	034114393	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 80 mg/12,5 mg compresse rivestite con film	SE/H/0565/001	034114381	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 80 mg/12,5 mg compresse rivestite con film	SE/H/0565/001	034114052	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 80 mg/12,5 mg	SE/H/0565/001	034114049	NOVARTIS EUROPHARM	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			LIMITED	
Cotareg 80 mg/12,5 mg compresse rivestite con film	SE/H/0565/001	034114708	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 80 mg/12,5 mg compresse rivestite con film	SE/H/0565/001	034114672	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 80 mg/12,5 mg compresse rivestite con film	SE/H/0565/001	034114660	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 80 mg/12,5 mg compresse rivestite con film	SE/H/0565/001	034114494	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 80 mg/12,5 mg compresse rivestite con film	SE/H/0565/001	034114696	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 80 mg/12,5 mg compresse rivestite con film	SE/H/0565/001	034114684	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 80 mg/12,5 mg compresse rivestite con film	SE/H/0565/001	034114482	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 80 mg/12,5 mg compresse rivestite con film	SE/H/0565/001	034114470	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 80 mg/12,5 mg compresse rivestite con film	SE/H/0565/001	034114468	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 80 mg/12,5 mg compresse rivestite con film	SE/H/0565/001	034114456	NOVARTIS EUROPHARM LIMITED	IT
Cotareg 80 mg/12,5 mg	SE/H/0565/001	034114025	NOVARTIS EUROPHARM	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
comprimé rivestito con film			LIMITED	
Cotareg 80 mg/12,5 mg comprimé rivestito con film	SE/H/0565/001	034114013	NOVARTIS EUROPHARM LIMITED	IT
COTAREG 80 mg/12,5 mg, comprimé pelliculé	SE/H/0565/001	3400934430053	NOVARTIS PHARMA S.A.S.	FR
COTAREG 80 mg/12,5 mg, comprimé pelliculé	SE/H/0565/001	3400934430114	NOVARTIS PHARMA S.A.S.	FR
COTAREG 80 mg/12,5 mg, comprimé pelliculé	SE/H/0565/001	3400934430282	NOVARTIS PHARMA S.A.S.	FR
COTAREG 80 mg/12,5 mg, comprimé pelliculé	SE/H/0565/001	3400934430343	NOVARTIS PHARMA S.A.S.	FR
COTAREG 80 mg/12,5 mg, comprimé pelliculé	SE/H/0565/001	3400937196192	NOVARTIS PHARMA S.A.S.	FR
COTAREG 80 mg/12,5 mg, comprimé pelliculé	SE/H/0565/001	3400937196253	NOVARTIS PHARMA S.A.S.	FR
COTAREG 80 mg/12,5 mg, comprimé pelliculé	SE/H/0565/001	3400937196314	NOVARTIS PHARMA S.A.S.	FR
COTAREG 80 mg/12,5 mg, comprimé pelliculé	SE/H/0565/001	3400938155761	NOVARTIS PHARMA S.A.S.	FR
COTAREG 80 mg/12,5 mg, comprimé pelliculé	SE/H/0565/001	3400938155822	NOVARTIS PHARMA S.A.S.	FR
COTAREG 80 mg/12,5 mg, comprimé pelliculé	SE/H/0565/001	3400938155990	NOVARTIS PHARMA S.A.S.	FR
COTAREG 80 mg/12,5 mg, comprimé pelliculé	SE/H/0565/001	3400938156072	NOVARTIS PHARMA S.A.S.	FR
COTAREG 80 mg/12,5 mg, comprimé pelliculé	SE/H/0565/001	3400938156133	NOVARTIS PHARMA S.A.S.	FR
COTAREG 80 mg/12,5 mg, comprimé pelliculé	SE/H/0565/001	3400938156362	NOVARTIS PHARMA S.A.S.	FR
COTAREG 80 mg/12,5 mg, comprimé pelliculé	SE/H/0565/001	3400938156423	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
COTAREG 80 mg/12,5 mg, comprimé pelliculé	SE/H/0565/001	3400934430053	NOVARTIS PHARMA S.A.S.	FR
COTAREG 80 mg/12,5 mg, comprimé pelliculé	SE/H/0565/001	3400934430114	NOVARTIS PHARMA S.A.S.	FR
COTAREG 80 mg/12,5 mg, comprimé pelliculé	SE/H/0565/001	3400934430282	NOVARTIS PHARMA S.A.S.	FR
COTAREG 80 mg/12,5 mg, comprimé pelliculé	SE/H/0565/001	3400934430343	NOVARTIS PHARMA S.A.S.	FR
COTAREG 80 mg/12,5 mg, comprimé pelliculé	SE/H/0565/001	3400937196192	NOVARTIS PHARMA S.A.S.	FR
COTAREG 80 mg/12,5 mg, comprimé pelliculé	SE/H/0565/001	3400937196253	NOVARTIS PHARMA S.A.S.	FR
COTAREG 80 mg/12,5 mg, comprimé pelliculé	SE/H/0565/001	3400937196314	NOVARTIS PHARMA S.A.S.	FR
COTAREG 80 mg/12,5 mg, comprimé pelliculé	SE/H/0565/001	3400938155761	NOVARTIS PHARMA S.A.S.	FR
COTAREG 80 mg/12,5 mg, comprimé pelliculé	SE/H/0565/001	3400938155822	NOVARTIS PHARMA S.A.S.	FR
COTAREG 80 mg/12,5 mg, comprimé pelliculé	SE/H/0565/001	3400938155990	NOVARTIS PHARMA S.A.S.	FR
COTAREG 80 mg/12,5 mg, comprimé pelliculé	SE/H/0565/001	3400938156072	NOVARTIS PHARMA S.A.S.	FR
COTAREG 80 mg/12,5 mg, comprimé pelliculé	SE/H/0565/001	3400938156133	NOVARTIS PHARMA S.A.S.	FR
COTAREG 80 mg/12,5 mg, comprimé pelliculé	SE/H/0565/001	3400938156362	NOVARTIS PHARMA S.A.S.	FR
COTAREG 80 mg/12,5 mg, comprimé pelliculé	SE/H/0565/001	3400938156423	NOVARTIS PHARMA S.A.S.	FR
COTAREG 80 mg/12,5 mg, comprimé pelliculé	SE/H/0565/001	3400934430053	NOVARTIS PHARMA S.A.S.	FR
COTAREG 80 mg/12,5 mg, comprimé pelliculé	SE/H/0565/001	3400934430114	NOVARTIS PHARMA S.A.S.	FR
COTAREG 80 mg/12,5 mg, comprimé pelliculé	SE/H/0565/001	3400934430282	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
mg, comprimé pelliculé				
COTAREG 80 mg/12,5 mg, comprimé pelliculé	SE/H/0565/001	3400934430343	NOVARTIS PHARMA S.A.S.	FR
COTAREG 80 mg/12,5 mg, comprimé pelliculé	SE/H/0565/001	3400937196192	NOVARTIS PHARMA S.A.S.	FR
COTAREG 80 mg/12,5 mg, comprimé pelliculé	SE/H/0565/001	3400937196253	NOVARTIS PHARMA S.A.S.	FR
COTAREG 80 mg/12,5 mg, comprimé pelliculé	SE/H/0565/001	3400937196314	NOVARTIS PHARMA S.A.S.	FR
COTAREG 80 mg/12,5 mg, comprimé pelliculé	SE/H/0565/001	3400938155761	NOVARTIS PHARMA S.A.S.	FR
COTAREG 80 mg/12,5 mg, comprimé pelliculé	SE/H/0565/001	3400938155822	NOVARTIS PHARMA S.A.S.	FR
COTAREG 80 mg/12,5 mg, comprimé pelliculé	SE/H/0565/001	3400938155990	NOVARTIS PHARMA S.A.S.	FR
COTAREG 80 mg/12,5 mg, comprimé pelliculé	SE/H/0565/001	3400938156072	NOVARTIS PHARMA S.A.S.	FR
COTAREG 80 mg/12,5 mg, comprimé pelliculé	SE/H/0565/001	3400938156133	NOVARTIS PHARMA S.A.S.	FR
COTAREG 80 mg/12,5 mg, comprimé pelliculé	SE/H/0565/001	3400938156362	NOVARTIS PHARMA S.A.S.	FR
COTAREG 80 mg/12,5 mg, comprimé pelliculé	SE/H/0565/001	3400938156423	NOVARTIS PHARMA S.A.S.	FR
COTAREG 80 mg/12,5 mg, comprimé pelliculé	SE/H/0565/001	3400934430053	NOVARTIS PHARMA S.A.S.	FR
COTAREG 80 mg/12,5 mg, comprimé pelliculé	SE/H/0565/001	3400934430114	NOVARTIS PHARMA S.A.S.	FR
COTAREG 80 mg/12,5 mg, comprimé pelliculé	SE/H/0565/001	3400934430282	NOVARTIS PHARMA S.A.S.	FR
COTAREG 80 mg/12,5 mg, comprimé pelliculé	SE/H/0565/001	3400934430343	NOVARTIS PHARMA S.A.S.	FR
COTAREG 80 mg/12,5 mg, comprimé pelliculé	SE/H/0565/001	3400937196192	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
COTAREG 80 mg/12,5 mg, comprimé pelliculé	SE/H/0565/001	3400937196253	NOVARTIS PHARMA S.A.S.	FR
COTAREG 80 mg/12,5 mg, comprimé pelliculé	SE/H/0565/001	3400937196314	NOVARTIS PHARMA S.A.S.	FR
COTAREG 80 mg/12,5 mg, comprimé pelliculé	SE/H/0565/001	3400938155761	NOVARTIS PHARMA S.A.S.	FR
COTAREG 80 mg/12,5 mg, comprimé pelliculé	SE/H/0565/001	3400938155822	NOVARTIS PHARMA S.A.S.	FR
COTAREG 80 mg/12,5 mg, comprimé pelliculé	SE/H/0565/001	3400938155990	NOVARTIS PHARMA S.A.S.	FR
COTAREG 80 mg/12,5 mg, comprimé pelliculé	SE/H/0565/001	3400938156072	NOVARTIS PHARMA S.A.S.	FR
COTAREG 80 mg/12,5 mg, comprimé pelliculé	SE/H/0565/001	3400938156133	NOVARTIS PHARMA S.A.S.	FR
COTAREG 80 mg/12,5 mg, comprimé pelliculé	SE/H/0565/001	3400938156362	NOVARTIS PHARMA S.A.S.	FR
COTAREG 80 mg/12,5 mg, comprimé pelliculé	SE/H/0565/001	3400938156423	NOVARTIS PHARMA S.A.S.	FR
Co-Vals 160 mg/12,5 mg comprimidos recubiertos con película	not available	65.797	LABORATORIOS DR. ESTEVE S.A.	ES
Co-Vals 160 mg/25 mg comprimidos recubiertos con película	not available	66.676	LABORATORIOS DR. ESTEVE S.A.	ES
Co-Vals 320 mg/12,5 mg comprimidos recubiertos con película	not available	70.361	LABORATORIOS DR. ESTEVE S.A.	ES
Co-Vals 320 mg/25 mg comprimidos recubiertos con película	not available	70.363	LABORATORIOS DR. ESTEVE S.A.	ES
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190420	NOVARTIS (HELLAS) S.A.C.I.	GR

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
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190421	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190422	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190423	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190424	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190425	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190426	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190427	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190428	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190429	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190430	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190431	NOVARTIS (HELLAS) S.A.C.I.	GR

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
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190432	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190401	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190402	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190403	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190404	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190405	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190406	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190407	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190408	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190409	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190410	NOVARTIS (HELLAS) S.A.C.I.	GR

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
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190411	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190412	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190413	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190414	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190415	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190416	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190417	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190418	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190419	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190401	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190402	NOVARTIS (HELLAS) S.A.C.I.	GR

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
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190403	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190404	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190405	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190406	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190407	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190408	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190409	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190410	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190411	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190412	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190413	NOVARTIS (HELLAS) S.A.C.I.	GR

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
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190414	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190415	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190416	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190417	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190418	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190419	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190420	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190421	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190422	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190423	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190424	NOVARTIS (HELLAS) S.A.C.I.	GR

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
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190425	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190426	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190427	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190428	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190429	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190430	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190431	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190432	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190401	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190402	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190403	NOVARTIS (HELLAS) S.A.C.I.	GR

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
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190404	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190405	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190406	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190407	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190408	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190409	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190410	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190411	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190412	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190413	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190414	NOVARTIS (HELLAS) S.A.C.I.	GR

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
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190415	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190416	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190417	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190418	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190419	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190420	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190421	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190422	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190423	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190424	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190425	NOVARTIS (HELLAS) S.A.C.I.	GR

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
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190426	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190427	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190428	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190429	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190430	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190431	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190432	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190420	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190421	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190422	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190423	NOVARTIS (HELLAS) S.A.C.I.	GR

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
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190424	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190425	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190426	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190427	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190428	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190429	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190430	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190431	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190432	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190401	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190402	NOVARTIS (HELLAS) S.A.C.I.	GR

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
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190403	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190404	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190405	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190406	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190407	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190408	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190409	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190410	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190411	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190412	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190413	NOVARTIS (HELLAS) S.A.C.I.	GR

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
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190414	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190415	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190416	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190417	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190418	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190419	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190420	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190421	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190422	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190423	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190424	NOVARTIS (HELLAS) S.A.C.I.	GR

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
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190425	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190426	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190427	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190428	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190429	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190430	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190431	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190432	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190401	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190402	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190403	NOVARTIS (HELLAS) S.A.C.I.	GR

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
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190404	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190405	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190406	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190407	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190408	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190409	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190410	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190411	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190412	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190413	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190414	NOVARTIS (HELLAS) S.A.C.I.	GR

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
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190415	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190416	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190417	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190418	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190419	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190420	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190421	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190422	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190423	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190424	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190425	NOVARTIS (HELLAS) S.A.C.I.	GR

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
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190426	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190427	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190428	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190429	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190430	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190431	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190432	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190401	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190402	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190403	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190404	NOVARTIS (HELLAS) S.A.C.I.	GR

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
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190405	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190406	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190407	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190408	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190409	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190410	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190411	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190412	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190413	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190414	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190415	NOVARTIS (HELLAS) S.A.C.I.	GR

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
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190416	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190417	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190418	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0407/002	247190419	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190601	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190602	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190603	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190604	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190605	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190606	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190607	NOVARTIS (HELLAS) S.A.C.I.	GR

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
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190608	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190609	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190610	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190611	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190612	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190613	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190614	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190615	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190616	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190617	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190618	NOVARTIS (HELLAS) S.A.C.I.	GR

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
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190619	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190620	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190621	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190622	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190623	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190624	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190625	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190626	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190627	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190628	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190629	NOVARTIS (HELLAS) S.A.C.I.	GR

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
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190630	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190631	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190632	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190601	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190602	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190603	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190604	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190605	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190606	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190607	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190608	NOVARTIS (HELLAS) S.A.C.I.	GR

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
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190609	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190610	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190611	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190612	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190613	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190614	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190615	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190616	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190617	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190618	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190619	NOVARTIS (HELLAS) S.A.C.I.	GR

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
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190620	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190621	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190622	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190623	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190624	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190625	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190626	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190627	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190628	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190629	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190630	NOVARTIS (HELLAS) S.A.C.I.	GR

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
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190631	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190632	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190601	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190602	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190603	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190604	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190605	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190606	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190607	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190608	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190609	NOVARTIS (HELLAS) S.A.C.I.	GR

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
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190610	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190611	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190612	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190613	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190614	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190615	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190616	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190617	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190618	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190619	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190620	NOVARTIS (HELLAS) S.A.C.I.	GR

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
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190621	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190622	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190623	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190624	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190625	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190626	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190627	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190628	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190629	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190630	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190631	NOVARTIS (HELLAS) S.A.C.I.	GR

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
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190632	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190601	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190602	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190603	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190604	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190605	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190606	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190607	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190608	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190609	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190610	NOVARTIS (HELLAS) S.A.C.I.	GR

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
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190611	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190612	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190613	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190614	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190615	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190616	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190617	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190618	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190619	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190620	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190621	NOVARTIS (HELLAS) S.A.C.I.	GR

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
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190622	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190623	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190624	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190625	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190626	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190627	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190628	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190629	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190630	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190631	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190632	NOVARTIS (HELLAS) S.A.C.I.	GR

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
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190601	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190602	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190603	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190604	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190605	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190606	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190607	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190608	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190609	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190610	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190611	NOVARTIS (HELLAS) S.A.C.I.	GR

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
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190612	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190613	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190614	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190615	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190616	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190617	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190618	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190619	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190620	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190621	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190622	NOVARTIS (HELLAS) S.A.C.I.	GR

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
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190623	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190624	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190625	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190626	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190627	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190628	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190629	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190630	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190631	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190632	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190601	NOVARTIS (HELLAS) S.A.C.I.	GR

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
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190602	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190603	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190604	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190605	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190606	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190607	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190608	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190609	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190610	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190611	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190612	NOVARTIS (HELLAS) S.A.C.I.	GR

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
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190613	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190614	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190615	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190616	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190617	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190618	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190619	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190620	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190621	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190622	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190623	NOVARTIS (HELLAS) S.A.C.I.	GR

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
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190624	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190625	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190626	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190627	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190628	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190629	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190630	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190631	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0407/004	247190632	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190501	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190502	NOVARTIS (HELLAS) S.A.C.I.	GR

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
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190503	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190504	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190505	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190506	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190507	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190508	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190509	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190510	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190511	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190512	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190513	NOVARTIS (HELLAS) S.A.C.I.	GR

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
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190514	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190515	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190516	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190517	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190518	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190519	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190520	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190521	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190522	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190523	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190524	NOVARTIS (HELLAS) S.A.C.I.	GR

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
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190525	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190526	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190527	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190528	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190529	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190530	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190531	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190532	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190501	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190502	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190503	NOVARTIS (HELLAS) S.A.C.I.	GR

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
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190504	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190505	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190506	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190507	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190508	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190509	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190510	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190511	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190512	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190513	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190514	NOVARTIS (HELLAS) S.A.C.I.	GR

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
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190515	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190516	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190517	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190518	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190519	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190520	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190521	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190522	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190523	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190524	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190525	NOVARTIS (HELLAS) S.A.C.I.	GR

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
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190526	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190527	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190528	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190529	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190530	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190531	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190532	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190501	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190502	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190503	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190504	NOVARTIS (HELLAS) S.A.C.I.	GR

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
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190505	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190506	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190507	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190508	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190509	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190510	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190511	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190512	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190513	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190514	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190515	NOVARTIS (HELLAS) S.A.C.I.	GR

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
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190516	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190517	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190518	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190519	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190520	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190521	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190522	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190523	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190524	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190525	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190526	NOVARTIS (HELLAS) S.A.C.I.	GR

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
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190527	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190528	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190529	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190530	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190531	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190532	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190501	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190502	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190503	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190504	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190505	NOVARTIS (HELLAS) S.A.C.I.	GR

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
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190506	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190507	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190508	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190509	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190510	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190511	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190512	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190513	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190514	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190515	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190516	NOVARTIS (HELLAS) S.A.C.I.	GR

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
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190517	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190518	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190519	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190520	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190521	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190522	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190523	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190524	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190525	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190526	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190527	NOVARTIS (HELLAS) S.A.C.I.	GR

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
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190528	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190529	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190530	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190531	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190532	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190501	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190502	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190503	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190504	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190505	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190506	NOVARTIS (HELLAS) S.A.C.I.	GR

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
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190507	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190508	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190509	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190510	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190511	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190512	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190513	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190514	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190515	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190516	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190517	NOVARTIS (HELLAS) S.A.C.I.	GR

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
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190518	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190519	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190520	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190521	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190522	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190523	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190524	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190525	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190526	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190527	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190528	NOVARTIS (HELLAS) S.A.C.I.	GR

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
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190529	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190530	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190531	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190532	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190501	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190502	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190503	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190504	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190505	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190506	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190507	NOVARTIS (HELLAS) S.A.C.I.	GR

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
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190508	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190509	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190510	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190511	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190512	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190513	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190514	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190515	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190516	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190517	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190518	NOVARTIS (HELLAS) S.A.C.I.	GR

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
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190519	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190520	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190521	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190522	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190523	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190524	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190525	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190526	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190527	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190528	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190529	NOVARTIS (HELLAS) S.A.C.I.	GR

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
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190530	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190531	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0407/003	247190532	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190301	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190302	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190303	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190304	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190305	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190306	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190307	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190308	NOVARTIS (HELLAS) S.A.C.I.	GR

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
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190309	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190310	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190311	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190312	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190313	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190314	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190315	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190316	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190317	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190318	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190319	NOVARTIS (HELLAS) S.A.C.I.	GR

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
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190320	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190321	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190322	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190323	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190324	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190325	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190326	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190327	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190328	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190329	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190330	NOVARTIS (HELLAS) S.A.C.I.	GR

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
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190331	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190332	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190301	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190302	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190303	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190304	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190305	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190306	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190307	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190308	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190309	NOVARTIS (HELLAS) S.A.C.I.	GR

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
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190310	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190311	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190312	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190313	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190314	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190315	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190316	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190317	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190318	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190319	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190320	NOVARTIS (HELLAS) S.A.C.I.	GR

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
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190321	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190322	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190323	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190324	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190325	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190326	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190327	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190328	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190329	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190330	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190331	NOVARTIS (HELLAS) S.A.C.I.	GR

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
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190332	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190301	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190302	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190303	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190304	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190305	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190306	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190307	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190308	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190309	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190310	NOVARTIS (HELLAS) S.A.C.I.	GR

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
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190311	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190312	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190313	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190314	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190315	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190316	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190317	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190318	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190319	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190320	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190321	NOVARTIS (HELLAS) S.A.C.I.	GR

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
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190322	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190323	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190324	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190325	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190326	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190327	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190328	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190329	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190330	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190331	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190332	NOVARTIS (HELLAS) S.A.C.I.	GR

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
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190301	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190302	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190303	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190304	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190305	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190306	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190307	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190308	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190309	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190310	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190311	NOVARTIS (HELLAS) S.A.C.I.	GR

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
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190312	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190313	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190314	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190315	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190316	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190317	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190318	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190319	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190320	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190321	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190322	NOVARTIS (HELLAS) S.A.C.I.	GR

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
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190323	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190324	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190325	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190326	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190327	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190328	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190329	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190330	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190331	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190332	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190301	NOVARTIS (HELLAS) S.A.C.I.	GR

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
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190302	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190303	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190304	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190305	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190306	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190307	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190308	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190309	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190310	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190311	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190312	NOVARTIS (HELLAS) S.A.C.I.	GR

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
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190313	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190314	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190315	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190316	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190317	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190318	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190319	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190320	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190321	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190322	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190323	NOVARTIS (HELLAS) S.A.C.I.	GR

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
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190324	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190325	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190326	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190327	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190328	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190329	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190330	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190331	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190332	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190301	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190302	NOVARTIS (HELLAS) S.A.C.I.	GR

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
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190303	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190304	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190305	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190306	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190307	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190308	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190309	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190310	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190311	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190312	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190313	NOVARTIS (HELLAS) S.A.C.I.	GR

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
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190314	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190315	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190316	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190317	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190318	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190319	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190320	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190321	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190322	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190323	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190324	NOVARTIS (HELLAS) S.A.C.I.	GR

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
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190325	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190326	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190327	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190328	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190329	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190330	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190331	NOVARTIS (HELLAS) S.A.C.I.	GR
DALZAD ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0407/001	247190332	NOVARTIS (HELLAS) S.A.C.I.	GR
Dalzac, πόσιμο διάλυμα 3mg/ml	SE/H/0407/005	247190701	NOVARTIS (HELLAS) S.A.C.I.	GR
Dalzac, πόσιμο διάλυμα 3mg/ml	SE/H/0407/005	247190701	NOVARTIS (HELLAS) S.A.C.I.	GR
Dalzac, πόσιμο διάλυμα 3mg/ml	SE/H/0407/005	247190701	NOVARTIS (HELLAS) S.A.C.I.	GR
Dalzac, πόσιμο διάλυμα 3mg/ml	SE/H/0407/005	247190701	NOVARTIS (HELLAS) S.A.C.I.	GR
Dalzac, πόσιμο διάλυμα	SE/H/0407/005	247190701	NOVARTIS (HELLAS)	GR

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
3mg/ml			S.A.C.I.	
Dalzad, πόσιμο διάλυμα 3mg/ml	SE/H/0407/005	247190701	NOVARTIS (HELLAS) S.A.C.I.	GR
Diovan	SE/H/0406/003	32751	NOVARTIS HEALTHCARE A/S	DK
Diovan	SE/H/0406/004	32752	NOVARTIS HEALTHCARE A/S	DK
Diovan	SE/H/0406/005	37642	NOVARTIS HEALTHCARE A/S	DK
Diovan	SE/H/0406/006	39956	NOVARTIS HEALTHCARE A/S	DK
Diovan 160 mg apvalkotās tabletes	SE/H/0406/004	02-0392	NOVARTIS FINLAND OY	LV
DIOVAN 160 MG COMPRIMATE FILMATE	SE/H/0406/004	3947/2011/01	NOVARTIS PHARMA GMBH	RO
DIOVAN 160 MG COMPRIMATE FILMATE	SE/H/0406/004	3947/2011/02	NOVARTIS PHARMA GMBH	RO
DIOVAN 160 MG COMPRIMATE FILMATE	SE/H/0406/004	3947/2011/03	NOVARTIS PHARMA GMBH	RO
DIOVAN 160 MG COMPRIMATE FILMATE	SE/H/0406/004	3947/2011/04	NOVARTIS PHARMA GMBH	RO
DIOVAN 160 MG COMPRIMATE FILMATE	SE/H/0406/004	3947/2011/05	NOVARTIS PHARMA GMBH	RO
DIOVAN 160 MG COMPRIMATE FILMATE	SE/H/0406/004	3947/2011/06	NOVARTIS PHARMA GMBH	RO
DIOVAN 160 MG COMPRIMATE FILMATE	SE/H/0406/004	3947/2011/07	NOVARTIS PHARMA GMBH	RO
DIOVAN 160 MG COMPRIMATE FILMATE	SE/H/0406/004	3947/2011/08	NOVARTIS PHARMA GMBH	RO
DIOVAN 160 MG COMPRIMATE FILMATE	SE/H/0406/004	3947/2011/09	NOVARTIS PHARMA GMBH	RO
DIOVAN 160 MG COMPRIMATE FILMATE	SE/H/0406/004	3947/2011/10	NOVARTIS PHARMA GMBH	RO
DIOVAN 160 MG COMPRIMATE FILMATE	SE/H/0406/004	3947/2011/11	NOVARTIS PHARMA GMBH	RO
Diovan 160 mg comprimate filmate	SE/H/0406/004	3947/2011/12	NOVARTIS PHARMA GMBH	RO

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 160 mg comprimate filmate	SE/H/0406/004	3947/2011/14	NOVARTIS PHARMA GMBH	RO
Diovan 160 mg comprimate filmate	SE/H/0406/004	3947/2011/13	NOVARTIS PHARMA GMBH	RO
Diovan 160 mg comprimate filmate	SE/H/0406/004	3947/2011/21	NOVARTIS PHARMA GMBH	RO
Diovan 160 mg comprimate filmate	SE/H/0406/004	3947/2011/20	NOVARTIS PHARMA GMBH	RO
Diovan 160 mg comprimate filmate	SE/H/0406/004	3947/2011/19	NOVARTIS PHARMA GMBH	RO
Diovan 160 mg comprimate filmate	SE/H/0406/004	3947/2011/18	NOVARTIS PHARMA GMBH	RO
Diovan 160 mg comprimate filmate	SE/H/0406/004	3947/2011/17	NOVARTIS PHARMA GMBH	RO
Diovan 160 mg comprimate filmate	SE/H/0406/004	3947/2011/16	NOVARTIS PHARMA GMBH	RO
Diovan 160 mg comprimate filmate	SE/H/0406/004	3947/2011/15	NOVARTIS PHARMA GMBH	RO
Diovan 160 mg comprimate filmate	SE/H/0406/004	3947/2011/22	NOVARTIS PHARMA GMBH	RO
Diovan 160 mg comprimate filmate	SE/H/0406/004	3947/2011/24	NOVARTIS PHARMA GMBH	RO
Diovan 160 mg comprimate filmate	SE/H/0406/004	3947/2011/23	NOVARTIS PHARMA GMBH	RO
Diovan 160 mg comprimate filmate	SE/H/0406/004	3947/2011/30	NOVARTIS PHARMA GMBH	RO
Diovan 160 mg comprimate filmate	SE/H/0406/004	3947/2011/29	NOVARTIS PHARMA GMBH	RO
Diovan 160 mg comprimate filmate	SE/H/0406/004	3947/2011/28	NOVARTIS PHARMA GMBH	RO
Diovan 160 mg comprimate filmate	SE/H/0406/004	3947/2011/27	NOVARTIS PHARMA GMBH	RO
Diovan 160 mg	SE/H/0406/004	3947/2011/26	NOVARTIS PHARMA GMBH	RO

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
comprimato filmate				
Diovan 160 mg comprimato filmate	SE/H/0406/004	3947/2011/25	NOVARTIS PHARMA GMBH	RO
Diován 160 mg comprimidos recubiertos con película	SE/H/0406/004	64.473	NOVARTIS FARMACÉUTICA S.A.	ES
Diovan 160 mg comprimidos revestidos por película	SE/H/0406/004	3787785	NOVARTIS FARMA - PRODUTOS FARMACÉUTICOS S.A.	PT
Diovan 160 mg comprimidos revestidos por película	SE/H/0406/004	3787884	NOVARTIS FARMA - PRODUTOS FARMACÉUTICOS S.A.	PT
Diovan 160 mg comprimidos revestidos por película	SE/H/0406/004	3787983	NOVARTIS FARMA - PRODUTOS FARMACÉUTICOS S.A.	PT
Diovan 160 mg comprimidos revestidos por película	SE/H/0406/004	3804184	NOVARTIS FARMA - PRODUTOS FARMACÉUTICOS S.A.	PT
Diovan 160 mg comprimidos revestidos por película	SE/H/0406/004	3804283	NOVARTIS FARMA - PRODUTOS FARMACÉUTICOS S.A.	PT
Diovan 160 mg comprimidos revestidos por película	SE/H/0406/004	5068333	NOVARTIS FARMA - PRODUTOS FARMACÉUTICOS S.A.	PT
DIOVAN 160 MG FILM-COATED TABLETS	SE/H/0406/004	PA 13/79/4	NOVARTIS PHARMACEUTICALS UK LIMITED	IE
DIOVAN 160 MG FILM-COATED TABLETS	SE/H/0406/004	088/00602	NOVARTIS PHARMACEUTICALS UK LIMITED	MT
Diovan 160 mg filmdragerade tabletter	SE/H/0406/004	16709	NOVARTIS FINLAND OY	FI
Diovan 160 mg	SE/H/0406/004	17495	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
film dragee tabletter				
Diovan 160 mg filmom obložene tablete	SE/H/0406/005	HR-H-178507347-01	NOVARTIS HRVATSKA D.O.O.	HR
Diovan 160 mg filmom obložene tablete	SE/H/0406/005	HR-H-178507347-02	NOVARTIS HRVATSKA D.O.O.	HR
Diovan 160 mg filmom obložene tablete	SE/H/0406/005	HR-H-178507347-03	NOVARTIS HRVATSKA D.O.O.	HR
Diovan 160 mg filmom obložene tablete	SE/H/0406/005	HR-H-178507347-04	NOVARTIS HRVATSKA D.O.O.	HR
Diovan 160 mg filmom obložene tablete	SE/H/0406/005	HR-H-178507347-05	NOVARTIS HRVATSKA D.O.O.	HR
Diovan 160 mg filmom obložene tablete	SE/H/0406/005	HR-H-178507347-06	NOVARTIS HRVATSKA D.O.O.	HR
Diovan 160 mg filmom obložene tablete	SE/H/0406/005	HR-H-178507347-07	NOVARTIS HRVATSKA D.O.O.	HR
Diovan 160 mg filmom obložene tablete	SE/H/0406/005	HR-H-178507347-08	NOVARTIS HRVATSKA D.O.O.	HR
Diovan 160 mg filmom obložene tablete	SE/H/0406/005	HR-H-178507347-09	NOVARTIS HRVATSKA D.O.O.	HR
Diovan 160 mg filmom obložene tablete	SE/H/0406/005	HR-H-178507347-10	NOVARTIS HRVATSKA D.O.O.	HR
Diovan 160 mg filmom obložene tablete	SE/H/0406/005	HR-H-178507347-11	NOVARTIS HRVATSKA D.O.O.	HR
Diovan 160 mg filmom obložene tablete	SE/H/0406/005	HR-H-178507347-12	NOVARTIS HRVATSKA D.O.O.	HR
Diovan 160 mg filmom obložene tablete	SE/H/0406/005	HR-H-178507347-13	NOVARTIS HRVATSKA D.O.O.	HR
Diovan 160 mg filmom obložene tablete	SE/H/0406/005	HR-H-178507347-14	NOVARTIS HRVATSKA D.O.O.	HR
Diovan 160 mg filmom obložene tablete	SE/H/0406/005	HR-H-178507347-15	NOVARTIS HRVATSKA D.O.O.	HR
Diovan 160 mg filmsko obložene tablete	SE/H/0406/004	H/03/00476/088	NOVARTIS PHARMA GMBH	SI

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 160 mg filmsko obložene tablete	SE/H/0406/004	H/03/00476/089	NOVARTIS PHARMA GMBH	SI
Diovan 160 mg filmsko obložene tablete	SE/H/0406/004	H/03/00476/090	NOVARTIS PHARMA GMBH	SI
Diovan 160 mg filmsko obložene tablete	SE/H/0406/004	H/03/00476/061	NOVARTIS PHARMA GMBH	SI
Diovan 160 mg filmsko obložene tablete	SE/H/0406/004	H/03/00476/062	NOVARTIS PHARMA GMBH	SI
Diovan 160 mg filmsko obložene tablete	SE/H/0406/004	H/03/00476/063	NOVARTIS PHARMA GMBH	SI
Diovan 160 mg filmsko obložene tablete	SE/H/0406/004	H/03/00476/064	NOVARTIS PHARMA GMBH	SI
Diovan 160 mg filmsko obložene tablete	SE/H/0406/004	H/03/00476/065	NOVARTIS PHARMA GMBH	SI
Diovan 160 mg filmsko obložene tablete	SE/H/0406/004	H/03/00476/066	NOVARTIS PHARMA GMBH	SI
Diovan 160 mg filmsko obložene tablete	SE/H/0406/004	H/03/00476/067	NOVARTIS PHARMA GMBH	SI
Diovan 160 mg filmsko obložene tablete	SE/H/0406/004	H/03/00476/068	NOVARTIS PHARMA GMBH	SI
Diovan 160 mg filmsko obložene tablete	SE/H/0406/004	H/03/00476/069	NOVARTIS PHARMA GMBH	SI
Diovan 160 mg filmsko obložene tablete	SE/H/0406/004	H/03/00476/070	NOVARTIS PHARMA GMBH	SI
Diovan 160 mg filmsko obložene tablete	SE/H/0406/004	H/03/00476/071	NOVARTIS PHARMA GMBH	SI
Diovan 160 mg filmsko obložene tablete	SE/H/0406/004	H/03/00476/072	NOVARTIS PHARMA GMBH	SI
Diovan 160 mg filmsko obložene tablete	SE/H/0406/004	H/03/00476/073	NOVARTIS PHARMA GMBH	SI
Diovan 160 mg filmsko obložene tablete	SE/H/0406/004	H/03/00476/074	NOVARTIS PHARMA GMBH	SI
Diovan 160 mg filmsko obložene tablete	SE/H/0406/004	H/03/00476/075	NOVARTIS PHARMA GMBH	SI

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
oblozene tablete				
Diovan 160 mg filmsko oblozene tablete	SE/H/0406/004	H/03/00476/076	NOVARTIS PHARMA GMBH	SI
Diovan 160 mg filmsko oblozene tablete	SE/H/0406/004	H/03/00476/077	NOVARTIS PHARMA GMBH	SI
Diovan 160 mg filmsko oblozene tablete	SE/H/0406/004	H/03/00476/078	NOVARTIS PHARMA GMBH	SI
Diovan 160 mg filmsko oblozene tablete	SE/H/0406/004	H/03/00476/079	NOVARTIS PHARMA GMBH	SI
Diovan 160 mg filmsko oblozene tablete	SE/H/0406/004	H/03/00476/080	NOVARTIS PHARMA GMBH	SI
Diovan 160 mg filmsko oblozene tablete	SE/H/0406/004	H/03/00476/081	NOVARTIS PHARMA GMBH	SI
Diovan 160 mg filmsko oblozene tablete	SE/H/0406/004	H/03/00476/082	NOVARTIS PHARMA GMBH	SI
Diovan 160 mg filmsko oblozene tablete	SE/H/0406/004	H/03/00476/083	NOVARTIS PHARMA GMBH	SI
Diovan 160 mg filmsko oblozene tablete	SE/H/0406/004	H/03/00476/084	NOVARTIS PHARMA GMBH	SI
Diovan 160 mg filmsko oblozene tablete	SE/H/0406/004	H/03/00476/085	NOVARTIS PHARMA GMBH	SI
Diovan 160 mg filmsko oblozene tablete	SE/H/0406/004	H/03/00476/086	NOVARTIS PHARMA GMBH	SI
Diovan 160 mg filmsko oblozene tablete	SE/H/0406/004	H/03/00476/087	NOVARTIS PHARMA GMBH	SI
Diovan 160 mg filmtabletta	SE/H/0406/004	OGYI-T-8484/01	NOVARTIS HUNGÁRIA KFT. PHARMA	HU
Diovan 160 mg filmtabletta	SE/H/0406/004	OGYI-T-8484/02	NOVARTIS HUNGÁRIA KFT. PHARMA	HU
Diovan 160 mg filmtabletta	SE/H/0406/004	OGYI-T-8484/13	NOVARTIS HUNGÁRIA KFT. PHARMA	HU
Diovan 160 mg filmtabletta	SE/H/0406/004	OGYI-T-8484/14	NOVARTIS HUNGÁRIA KFT. PHARMA	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
DIOVAN 160 MG FILMTABLETTEN	SE/H/0406/004	1-24276	NOVARTIS PHARMA GMBH	AT
Diovan 160 mg filmhúðaðar töflur	SE/H/0406/004	IS/1/01/042/02	NOVARTIS HEALTHCARE A/S	IS
Diovan 160 mg kalvopäällysteinen tabletti	SE/H/0406/004	16709	NOVARTIS FINLAND OY	FI
Diovan 160 mg plêvele dengtos tabletès	SE/H/0406/004	LT/1/01/1236/011	NOVARTIS FINLAND OY	LT
Diovan 160 mg plêvele dengtos tabletès	SE/H/0406/004	LT/1/01/1236/042	NOVARTIS FINLAND OY	LT
Diovan 160 mg plêvele dengtos tabletès	SE/H/0406/004	LT/1/01/1236/043	NOVARTIS FINLAND OY	LT
Diovan 160 mg plêvele dengtos tabletès	SE/H/0406/004	LT/1/01/1236/044	NOVARTIS FINLAND OY	LT
Diovan 160 mg plêvele dengtos tabletès	SE/H/0406/004	LT/1/01/1236/045	NOVARTIS FINLAND OY	LT
Diovan 160 mg plêvele dengtos tabletès	SE/H/0406/004	LT/1/01/1236/046	NOVARTIS FINLAND OY	LT
Diovan 160 mg plêvele dengtos tabletès	SE/H/0406/004	LT/1/01/1236/047	NOVARTIS FINLAND OY	LT
Diovan 160 mg plêvele dengtos tabletès	SE/H/0406/004	LT/1/01/1236/048	NOVARTIS FINLAND OY	LT
Diovan 160 mg plêvele dengtos tabletès	SE/H/0406/004	LT/1/01/1236/049	NOVARTIS FINLAND OY	LT
Diovan 160 mg plêvele dengtos tabletès	SE/H/0406/004	LT/1/01/1236/050	NOVARTIS FINLAND OY	LT
Diovan 160 mg plêvele dengtos tabletès	SE/H/0406/004	LT/1/01/1236/051	NOVARTIS FINLAND OY	LT
Diovan 160 mg plêvele dengtos tabletès	SE/H/0406/004	LT/1/01/1236/052	NOVARTIS FINLAND OY	LT
Diovan 160 mg plêvele dengtos tabletès	SE/H/0406/004	LT/1/01/1236/053	NOVARTIS FINLAND OY	LT

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 160 mg plėvele dengtos tabletės	SE/H/0406/004	LT/1/01/1236/054	NOVARTIS FINLAND OY	LT
Diovan 160 mg plėvele dengtos tabletės	SE/H/0406/004	LT/1/01/1236/055	NOVARTIS FINLAND OY	LT
Diovan 160 mg tabletki powlekane	SE/H/0406/004	9292	NOVARTIS POLAND SP. Z O. O.	PL
DIOVAN 160, 160 MG FILMOMHULDE TABLETTEN	SE/H/0406/004	RVG 26940	NOVARTIS PHARMA B.V.	NL
Diovan 3 mg/ml belsőleges oldat	SE/H/0406/007	OGYI-T-8484/15	NOVARTIS HUNGÁRIA KFT. PHARMA	HU
DIOVAN 3 MG/ML DRANK	SE/H/0406/007	RVG 107481	NOVARTIS PHARMA B.V.	NL
Diovan 3 mg/ml geriamasis tirpalas	SE/H/0406/007	LT/1/01/1236/014	NOVARTIS FINLAND OY	LT
DIOVAN 3 MG/ML LÖSUNG ZUM EINNEHMEN	SE/H/0406/007	1-29223	NOVARTIS PHARMA GMBH	AT
Diovan 3 mg/ml mikstur, oppløsning	SE/H/0406/007	10-7589	NOVARTIS NORGE AS	NO
Diovan 3 mg/ml mixtúra, lausn	SE/H/0406/007	IS/1/10/078/01	NOVARTIS HEALTHCARE A/S	IS
Diovan 3 mg/ml oraaliliuos	SE/H/0406/007	27966	NOVARTIS FINLAND OY	FI
Diovan 3 mg/ml oral lösning	SE/H/0406/007	43158	NOVARTIS SVERIGE AB	SE
DIOVAN 3 MG/ML ORAL SOLUTION	SE/H/0406/007	PA 13/79/7	NOVARTIS PHARMACEUTICALS UK LIMITED	IE
DIOVAN 3 MG/ML ORAL SOLUTION	SE/H/0406/007	088/00605	NOVARTIS PHARMACEUTICALS UK LIMITED	MT
DIOVAN 3 MG/ML ORAL SOLUTION	SE/H/0406/007	PL 00101/0956	NOVARTIS PHARMACEUTICALS UK	UK

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
			LIMITED	
Diovan 3 mg/ml peroralna raztopina	SE/H/0406/007	H/03/00476/121	NOVARTIS PHARMA GMBH	SI
Diovan 3 mg/ml perorálny roztok	SE/H/0406/007	58/0287/10-S	NOVARTIS, S.R.O.	SK
Diovan 3 mg/ml roztwór doustny	SE/H/0406/007	17043	NOVARTIS POLAND SP. Z O. O.	PL
Diovan 3 mg/ml šķīdums iekšķīgai lietošanai	SE/H/0406/007	10-0320	NOVARTIS FINLAND OY	LV
Diovan 3 mg/ml solução oral	SE/H/0406/007	5303441	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
DIOVAN 3 mg/ml solución oral	SE/H/0406/007	72.385	NOVARTIS FARMACÉUTICA S.A.	ES
Diovan 3 mg/ml soluție orală	SE/H/0406/007	9051/2016/01	NOVARTIS PHARMA GMBH	RO
Diovan 3 mg/ml suukaudne lahus	SE/H/0406/007	687310	NOVARTIS FINLAND OY	EE
Diovan 3 mg/ml, perorální roztok	SE/H/0406/007	58/550/10-C	NOVARTIS, S.R.O.	CZ
Diovan 3 mg/ml oral lösning	SE/H/0406/007	27966	NOVARTIS FINLAND OY	FI
Diován 320 mg comprimidos recubiertos con película	SE/H/0406/006	68.817	NOVARTIS FARMACÉUTICA S.A.	ES
DIOVAN 320 MG FILM-COATED TABLETS	SE/H/0406/006	PA 13/79/6	NOVARTIS PHARMACEUTICALS UK LIMITED	IE
DIOVAN 320 MG FILM-COATED TABLETS	SE/H/0406/006	088/00603	NOVARTIS PHARMACEUTICALS UK LIMITED	MT
Diovan 320 mg filmdragerade tabletter	SE/H/0406/006	22668	NOVARTIS FINLAND OY	FI

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 320 mg filmdragerade tabletter	SE/H/0406/006	22840	NOVARTIS SVERIGE AB	SE
Diovan 320 mg filmom obložene tablete	SE/H/0406/006	HR-H-512719548-01	NOVARTIS HRVATSKA D.O.O.	HR
Diovan 320 mg filmom obložene tablete	SE/H/0406/006	HR-H-512719548-02	NOVARTIS HRVATSKA D.O.O.	HR
Diovan 320 mg filmom obložene tablete	SE/H/0406/006	HR-H-512719548-03	NOVARTIS HRVATSKA D.O.O.	HR
Diovan 320 mg filmom obložene tablete	SE/H/0406/006	HR-H-512719548-04	NOVARTIS HRVATSKA D.O.O.	HR
Diovan 320 mg filmom obložene tablete	SE/H/0406/006	HR-H-512719548-05	NOVARTIS HRVATSKA D.O.O.	HR
Diovan 320 mg filmom obložene tablete	SE/H/0406/006	HR-H-512719548-06	NOVARTIS HRVATSKA D.O.O.	HR
Diovan 320 mg filmom obložene tablete	SE/H/0406/006	HR-H-512719548-07	NOVARTIS HRVATSKA D.O.O.	HR
Diovan 320 mg filmom obložene tablete	SE/H/0406/006	HR-H-512719548-08	NOVARTIS HRVATSKA D.O.O.	HR
Diovan 320 mg filmom obložene tablete	SE/H/0406/006	HR-H-512719548-09	NOVARTIS HRVATSKA D.O.O.	HR
Diovan 320 mg filmom obložene tablete	SE/H/0406/006	HR-H-512719548-10	NOVARTIS HRVATSKA D.O.O.	HR
Diovan 320 mg filmom obložene tablete	SE/H/0406/006	HR-H-512719548-11	NOVARTIS HRVATSKA D.O.O.	HR
Diovan 320 mg filmom obložene tablete	SE/H/0406/006	HR-H-512719548-12	NOVARTIS HRVATSKA D.O.O.	HR
Diovan 320 mg filmom obložene tablete	SE/H/0406/006	HR-H-512719548-13	NOVARTIS HRVATSKA D.O.O.	HR
Diovan 320 mg filmom obložene tablete	SE/H/0406/006	HR-H-512719548-14	NOVARTIS HRVATSKA D.O.O.	HR
Diovan 320 mg filmom obložene tablete	SE/H/0406/006	HR-H-512719548-15	NOVARTIS HRVATSKA D.O.O.	HR
Diovan 320 mg filmsko	SE/H/0406/006	H/03/00476/091	NOVARTIS PHARMA GMBH	SI

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
oblozene tablete				
Diovan 320 mg filmsko oblozene tablete	SE/H/0406/006	H/03/00476/092	NOVARTIS PHARMA GMBH	SI
Diovan 320 mg filmsko oblozene tablete	SE/H/0406/006	H/03/00476/093	NOVARTIS PHARMA GMBH	SI
Diovan 320 mg filmsko oblozene tablete	SE/H/0406/006	H/03/00476/094	NOVARTIS PHARMA GMBH	SI
Diovan 320 mg filmsko oblozene tablete	SE/H/0406/006	H/03/00476/095	NOVARTIS PHARMA GMBH	SI
Diovan 320 mg filmsko oblozene tablete	SE/H/0406/006	H/03/00476/096	NOVARTIS PHARMA GMBH	SI
Diovan 320 mg filmsko oblozene tablete	SE/H/0406/006	H/03/00476/097	NOVARTIS PHARMA GMBH	SI
Diovan 320 mg filmsko oblozene tablete	SE/H/0406/006	H/03/00476/098	NOVARTIS PHARMA GMBH	SI
Diovan 320 mg filmsko oblozene tablete	SE/H/0406/006	H/03/00476/099	NOVARTIS PHARMA GMBH	SI
Diovan 320 mg filmsko oblozene tablete	SE/H/0406/006	H/03/00476/100	NOVARTIS PHARMA GMBH	SI
Diovan 320 mg filmsko oblozene tablete	SE/H/0406/006	H/03/00476/101	NOVARTIS PHARMA GMBH	SI
Diovan 320 mg filmsko oblozene tablete	SE/H/0406/006	H/03/00476/102	NOVARTIS PHARMA GMBH	SI
Diovan 320 mg filmsko oblozene tablete	SE/H/0406/006	H/03/00476/103	NOVARTIS PHARMA GMBH	SI
Diovan 320 mg filmsko oblozene tablete	SE/H/0406/006	H/03/00476/104	NOVARTIS PHARMA GMBH	SI
Diovan 320 mg filmsko oblozene tablete	SE/H/0406/006	H/03/00476/105	NOVARTIS PHARMA GMBH	SI
Diovan 320 mg filmsko oblozene tablete	SE/H/0406/006	H/03/00476/106	NOVARTIS PHARMA GMBH	SI
Diovan 320 mg filmsko oblozene tablete	SE/H/0406/006	H/03/00476/107	NOVARTIS PHARMA GMBH	SI

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 320 mg filmsko obložene tablete	SE/H/0406/006	H/03/00476/108	NOVARTIS PHARMA GMBH	SI
Diovan 320 mg filmsko obložene tablete	SE/H/0406/006	H/03/00476/109	NOVARTIS PHARMA GMBH	SI
Diovan 320 mg filmsko obložene tablete	SE/H/0406/006	H/03/00476/110	NOVARTIS PHARMA GMBH	SI
Diovan 320 mg filmsko obložene tablete	SE/H/0406/006	H/03/00476/111	NOVARTIS PHARMA GMBH	SI
Diovan 320 mg filmsko obložene tablete	SE/H/0406/006	H/03/00476/112	NOVARTIS PHARMA GMBH	SI
Diovan 320 mg filmsko obložene tablete	SE/H/0406/006	H/03/00476/113	NOVARTIS PHARMA GMBH	SI
Diovan 320 mg filmsko obložene tablete	SE/H/0406/006	H/03/00476/114	NOVARTIS PHARMA GMBH	SI
Diovan 320 mg filmsko obložene tablete	SE/H/0406/006	H/03/00476/115	NOVARTIS PHARMA GMBH	SI
Diovan 320 mg filmsko obložene tablete	SE/H/0406/006	H/03/00476/116	NOVARTIS PHARMA GMBH	SI
Diovan 320 mg filmsko obložene tablete	SE/H/0406/006	H/03/00476/117	NOVARTIS PHARMA GMBH	SI
Diovan 320 mg filmsko obložene tablete	SE/H/0406/006	H/03/00476/118	NOVARTIS PHARMA GMBH	SI
Diovan 320 mg filmsko obložene tablete	SE/H/0406/006	H/03/00476/119	NOVARTIS PHARMA GMBH	SI
Diovan 320 mg filmsko obložene tablete	SE/H/0406/006	H/03/00476/120	NOVARTIS PHARMA GMBH	SI
DIOVAN 320 MG FILMTABLETTEN	SE/H/0406/006	1-26881	NOVARTIS PHARMA GMBH	AT
Diovan 320 mg filmuhúðaðar töflur	SE/H/0406/006	IS/1/06/055/01	NOVARTIS HEALTHCARE A/S	IS
Diovan 320 mg kalvopäällysteinen tabletti	SE/H/0406/006	22668	NOVARTIS FINLAND OY	FI

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 320 mg plêvele dengtos tabletès	SE/H/0406/006	LT/1/01/1236/001	NOVARTIS FINLAND OY	LT
Diovan 320 mg plêvele dengtos tabletès	SE/H/0406/006	LT/1/01/1236/002	NOVARTIS FINLAND OY	LT
Diovan 320 mg plêvele dengtos tabletès	SE/H/0406/006	LT/1/01/1236/003	NOVARTIS FINLAND OY	LT
Diovan 320 mg plêvele dengtos tabletès	SE/H/0406/006	LT/1/01/1236/004	NOVARTIS FINLAND OY	LT
Diovan 320 mg plêvele dengtos tabletès	SE/H/0406/006	LT/1/01/1236/005	NOVARTIS FINLAND OY	LT
Diovan 320 mg plêvele dengtos tabletès	SE/H/0406/006	LT/1/01/1236/007	NOVARTIS FINLAND OY	LT
Diovan 320 mg plêvele dengtos tabletès	SE/H/0406/006	LT/1/01/1236/008	NOVARTIS FINLAND OY	LT
Diovan 320 mg plêvele dengtos tabletès	SE/H/0406/006	LT/1/01/1236/009	NOVARTIS FINLAND OY	LT
Diovan 320 mg plêvele dengtos tabletès	SE/H/0406/006	LT/1/01/1236/056	NOVARTIS FINLAND OY	LT
Diovan 320 mg plêvele dengtos tabletès	SE/H/0406/006	LT/1/01/1236/057	NOVARTIS FINLAND OY	LT
Diovan 320 mg plêvele dengtos tabletès	SE/H/0406/006	LT/1/01/1236/058	NOVARTIS FINLAND OY	LT
Diovan 320 mg plêvele dengtos tabletès	SE/H/0406/006	LT/1/01/1236/059	NOVARTIS FINLAND OY	LT
Diovan 320 mg plêvele dengtos tabletès	SE/H/0406/006	LT/1/01/1236/060	NOVARTIS FINLAND OY	LT
Diovan 320 mg plêvele dengtos tabletès	SE/H/0406/006	LT/1/01/1236/061	NOVARTIS FINLAND OY	LT
Diovan 320 mg plêvele dengtos tabletès	SE/H/0406/006	LT/1/01/1236/062	NOVARTIS FINLAND OY	LT
DIOVAN 320 mg tabletter, filmdrasjerte	SE/H/0406/006	05-3646	NOVARTIS NORGE AS	NO
DIOVAN 320, 320 MG	SE/H/0406/006	RVG 34472	NOVARTIS PHARMA B.V.	NL

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
FILMOMHULDE TABLETTEN				
Diovan 40 mg apvalkotās tabletes	SE/H/0406/005	02-0390	NOVARTIS FINLAND OY	LV
Diovan 40 mg comprimate filmate	SE/H/0406/005	3945/2011/01	NOVARTIS PHARMA GMBH	RO
Diovan 40 mg comprimate filmate	SE/H/0406/005	3945/2011/02	NOVARTIS PHARMA GMBH	RO
Diovan 40 mg comprimate filmate	SE/H/0406/005	3945/2011/03	NOVARTIS PHARMA GMBH	RO
Diovan 40 mg comprimate filmate	SE/H/0406/005	3945/2011/04	NOVARTIS PHARMA GMBH	RO
Diovan 40 mg comprimate filmate	SE/H/0406/005	3945/2011/05	NOVARTIS PHARMA GMBH	RO
Diovan 40 mg comprimate filmate	SE/H/0406/005	3945/2011/06	NOVARTIS PHARMA GMBH	RO
Diovan 40 mg comprimate filmate	SE/H/0406/005	3945/2011/07	NOVARTIS PHARMA GMBH	RO
Diovan 40 mg comprimate filmate	SE/H/0406/005	3945/2011/08	NOVARTIS PHARMA GMBH	RO
Diovan 40 mg comprimate filmate	SE/H/0406/005	3945/2011/09	NOVARTIS PHARMA GMBH	RO
Diovan 40 mg comprimate filmate	SE/H/0406/005	3945/2011/10	NOVARTIS PHARMA GMBH	RO
Diovan 40 mg comprimate filmate	SE/H/0406/005	3945/2011/11	NOVARTIS PHARMA GMBH	RO
Diovan 40 mg comprimate filmate	SE/H/0406/005	3945/2011/12	NOVARTIS PHARMA GMBH	RO
Diovan 40 mg comprimate filmate	SE/H/0406/005	3945/2011/13	NOVARTIS PHARMA GMBH	RO
Diovan 40 mg comprimate filmate	SE/H/0406/005	3945/2011/14	NOVARTIS PHARMA GMBH	RO
Diovan 40 mg	SE/H/0406/005	3945/2011/15	NOVARTIS PHARMA GMBH	RO

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
comprimate filmate				
Diovan 40 mg comprimate filmate	SE/H/0406/005	3945/2011/16	NOVARTIS PHARMA GMBH	RO
Diovan 40 mg comprimate filmate	SE/H/0406/005	3945/2011/17	NOVARTIS PHARMA GMBH	RO
Diovan 40 mg comprimate filmate	SE/H/0406/005	3945/2011/18	NOVARTIS PHARMA GMBH	RO
Diovan 40 mg comprimate filmate	SE/H/0406/005	3945/2011/19	NOVARTIS PHARMA GMBH	RO
Diovan 40 mg comprimate filmate	SE/H/0406/005	3945/2011/20	NOVARTIS PHARMA GMBH	RO
Diovan 40 mg comprimate filmate	SE/H/0406/005	3945/2011/21	NOVARTIS PHARMA GMBH	RO
Diovan 40 mg comprimate filmate	SE/H/0406/005	3945/2011/22	NOVARTIS PHARMA GMBH	RO
Diovan 40 mg comprimate filmate	SE/H/0406/005	3945/2011/23	NOVARTIS PHARMA GMBH	RO
Diovan 40 mg comprimate filmate	SE/H/0406/005	3945/2011/24	NOVARTIS PHARMA GMBH	RO
Diovan 40 mg comprimate filmate	SE/H/0406/005	3945/2011/25	NOVARTIS PHARMA GMBH	RO
Diovan 40 mg comprimate filmate	SE/H/0406/005	3945/2011/26	NOVARTIS PHARMA GMBH	RO
Diovan 40 mg comprimate filmate	SE/H/0406/005	3945/2011/27	NOVARTIS PHARMA GMBH	RO
Diovan 40 mg comprimate filmate	SE/H/0406/005	3945/2011/28	NOVARTIS PHARMA GMBH	RO
Diovan 40 mg comprimate filmate	SE/H/0406/005	3945/2011/29	NOVARTIS PHARMA GMBH	RO
Diovan 40 mg comprimate filmate	SE/H/0406/005	3945/2011/30	NOVARTIS PHARMA GMBH	RO
Diovan 40 mg comprimidos revestidos	SE/H/0406/005	5068317	NOVARTIS FARMA - PRODUTOS	PT

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
por película			FARMACÊUTICOS S.A.	
Diovan 40 mg comprimidos revestidos por película	SE/H/0406/005	5448881	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Diovan 40 mg comprimidos revestidos por película	SE/H/0406/005	5448980	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Diovan 40 mg comprimidos revestidos por película	SE/H/0406/005	5449087	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Diovan 40 mg comprimidos revestidos por película	SE/H/0406/005	5449186	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
DIOVAN 40 MG FILM-COATED TABLETS	SE/H/0406/005	PA 13/79/5	NOVARTIS PHARMACEUTICALS UK LIMITED	IE
DIOVAN 40 MG FILM-COATED TABLETS	SE/H/0406/005	088/00604	NOVARTIS PHARMACEUTICALS UK LIMITED	MT
Diovan 40 mg filmdragerade tabletter	SE/H/0406/005	20510	NOVARTIS FINLAND OY	FI
Diovan 40 mg filmdragerade tabletter	SE/H/0406/005	20464	NOVARTIS SVERIGE AB	SE
Diovan 40 mg filmom oblozene tablete	SE/H/0406/003	HR-H-590752475-01	NOVARTIS HRVATSKA D.O.O.	HR
Diovan 40 mg filmom oblozene tablete	SE/H/0406/003	HR-H-590752475-02	NOVARTIS HRVATSKA D.O.O.	HR
Diovan 40 mg filmom oblozene tablete	SE/H/0406/003	HR-H-590752475-03	NOVARTIS HRVATSKA D.O.O.	HR
Diovan 40 mg filmom oblozene tablete	SE/H/0406/003	HR-H-590752475-04	NOVARTIS HRVATSKA D.O.O.	HR
Diovan 40 mg filmom oblozene tablete	SE/H/0406/003	HR-H-590752475-05	NOVARTIS HRVATSKA D.O.O.	HR

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 40 mg filmom obložene tablete	SE/H/0406/003	HR-H-590752475-06	NOVARTIS HRVATSKA D.O.O.	HR
Diovan 40 mg filmom obložene tablete	SE/H/0406/003	HR-H-590752475-07	NOVARTIS HRVATSKA D.O.O.	HR
Diovan 40 mg filmom obložene tablete	SE/H/0406/003	HR-H-590752475-08	NOVARTIS HRVATSKA D.O.O.	HR
Diovan 40 mg filmom obložene tablete	SE/H/0406/003	HR-H-590752475-09	NOVARTIS HRVATSKA D.O.O.	HR
Diovan 40 mg filmom obložene tablete	SE/H/0406/003	HR-H-590752475-10	NOVARTIS HRVATSKA D.O.O.	HR
Diovan 40 mg filmom obložene tablete	SE/H/0406/003	HR-H-590752475-11	NOVARTIS HRVATSKA D.O.O.	HR
Diovan 40 mg filmom obložene tablete	SE/H/0406/003	HR-H-590752475-12	NOVARTIS HRVATSKA D.O.O.	HR
Diovan 40 mg filmom obložene tablete	SE/H/0406/003	HR-H-590752475-13	NOVARTIS HRVATSKA D.O.O.	HR
Diovan 40 mg filmom obložene tablete	SE/H/0406/003	HR-H-590752475-14	NOVARTIS HRVATSKA D.O.O.	HR
Diovan 40 mg filmom obložene tablete	SE/H/0406/003	HR-H-590752475-15	NOVARTIS HRVATSKA D.O.O.	HR
Diovan 40 mg filmsko obložene tablete	SE/H/0406/005	H/03/00476/024	NOVARTIS PHARMA GMBH	SI
Diovan 40 mg filmsko obložene tablete	SE/H/0406/005	H/03/00476/025	NOVARTIS PHARMA GMBH	SI
Diovan 40 mg filmsko obložene tablete	SE/H/0406/005	H/03/00476/026	NOVARTIS PHARMA GMBH	SI
Diovan 40 mg filmsko obložene tablete	SE/H/0406/005	H/03/00476/027	NOVARTIS PHARMA GMBH	SI
Diovan 40 mg filmsko obložene tablete	SE/H/0406/005	H/03/00476/028	NOVARTIS PHARMA GMBH	SI
Diovan 40 mg filmsko obložene tablete	SE/H/0406/005	H/03/00476/029	NOVARTIS PHARMA GMBH	SI
Diovan 40 mg filmsko obložene tablete	SE/H/0406/005	H/03/00476/030	NOVARTIS PHARMA GMBH	SI

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
oblozene tablete				
Diovan 40 mg filmsko oblozene tablete	SE/H/0406/005	H/03/00476/001	NOVARTIS PHARMA GMBH	SI
Diovan 40 mg filmsko oblozene tablete	SE/H/0406/005	H/03/00476/002	NOVARTIS PHARMA GMBH	SI
Diovan 40 mg filmsko oblozene tablete	SE/H/0406/005	H/03/00476/003	NOVARTIS PHARMA GMBH	SI
Diovan 40 mg filmsko oblozene tablete	SE/H/0406/005	H/03/00476/004	NOVARTIS PHARMA GMBH	SI
Diovan 40 mg filmsko oblozene tablete	SE/H/0406/005	H/03/00476/005	NOVARTIS PHARMA GMBH	SI
Diovan 40 mg filmsko oblozene tablete	SE/H/0406/005	H/03/00476/006	NOVARTIS PHARMA GMBH	SI
Diovan 40 mg filmsko oblozene tablete	SE/H/0406/005	H/03/00476/007	NOVARTIS PHARMA GMBH	SI
Diovan 40 mg filmsko oblozene tablete	SE/H/0406/005	H/03/00476/008	NOVARTIS PHARMA GMBH	SI
Diovan 40 mg filmsko oblozene tablete	SE/H/0406/005	H/03/00476/009	NOVARTIS PHARMA GMBH	SI
Diovan 40 mg filmsko oblozene tablete	SE/H/0406/005	H/03/00476/010	NOVARTIS PHARMA GMBH	SI
Diovan 40 mg filmsko oblozene tablete	SE/H/0406/005	H/03/00476/011	NOVARTIS PHARMA GMBH	SI
Diovan 40 mg filmsko oblozene tablete	SE/H/0406/005	H/03/00476/012	NOVARTIS PHARMA GMBH	SI
Diovan 40 mg filmsko oblozene tablete	SE/H/0406/005	H/03/00476/013	NOVARTIS PHARMA GMBH	SI
Diovan 40 mg filmsko oblozene tablete	SE/H/0406/005	H/03/00476/014	NOVARTIS PHARMA GMBH	SI
Diovan 40 mg filmsko oblozene tablete	SE/H/0406/005	H/03/00476/015	NOVARTIS PHARMA GMBH	SI
Diovan 40 mg filmsko oblozene tablete	SE/H/0406/005	H/03/00476/016	NOVARTIS PHARMA GMBH	SI

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 40 mg filmsko obložene tablete	SE/H/0406/005	H/03/00476/017	NOVARTIS PHARMA GMBH	SI
Diovan 40 mg filmsko obložene tablete	SE/H/0406/005	H/03/00476/018	NOVARTIS PHARMA GMBH	SI
Diovan 40 mg filmsko obložene tablete	SE/H/0406/005	H/03/00476/019	NOVARTIS PHARMA GMBH	SI
Diovan 40 mg filmsko obložene tablete	SE/H/0406/005	H/03/00476/020	NOVARTIS PHARMA GMBH	SI
Diovan 40 mg filmsko obložene tablete	SE/H/0406/005	H/03/00476/021	NOVARTIS PHARMA GMBH	SI
Diovan 40 mg filmsko obložene tablete	SE/H/0406/005	H/03/00476/022	NOVARTIS PHARMA GMBH	SI
Diovan 40 mg filmsko obložene tablete	SE/H/0406/005	H/03/00476/023	NOVARTIS PHARMA GMBH	SI
Diovan 40 mg filmtabletta	SE/H/0406/005	OGYI-T-8484/05	NOVARTIS HUNGÁRIA KFT. PHARMA	HU
Diovan 40 mg filmtabletta	SE/H/0406/005	OGYI-T-8484/06	NOVARTIS HUNGÁRIA KFT. PHARMA	HU
Diovan 40 mg filmtabletta	SE/H/0406/005	OGYI-T-8484/07	NOVARTIS HUNGÁRIA KFT. PHARMA	HU
Diovan 40 mg filmtabletta	SE/H/0406/005	OGYI-T-8484/08	NOVARTIS HUNGÁRIA KFT. PHARMA	HU
DIOVAN 40 MG FILMTABLETTEN	SE/H/0406/005	1-25945	NOVARTIS PHARMA GMBH	AT
Diovan 40 mg filmhúðaðar töflur	SE/H/0406/005	IS/1/05/010/01	NOVARTIS HEALTHCARE A/S	IS
Diovan 40 mg kalvopäällysteinen tabletti	SE/H/0406/005	20510	NOVARTIS FINLAND OY	FI
DIOVAN 40 mg tabletter, filmdrasjerte	SE/H/0406/005	04-2470	NOVARTIS NORGE AS	NO
DIOVAN 40, 40 MG FILMOMHULDE	SE/H/0406/005	RVG 32137	NOVARTIS PHARMA B.V.	NL

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
TABLETTEN				
Diovan 80 mg apvalkotās tabletes	SE/H/0406/003	02-0391	NOVARTIS FINLAND OY	LV
Diovan 80 mg comprimate filmate	SE/H/0406/003	3946/2011/01	NOVARTIS PHARMA GMBH	RO
Diovan 80 mg comprimate filmate	SE/H/0406/003	3946/2011/02	NOVARTIS PHARMA GMBH	RO
Diovan 80 mg comprimate filmate	SE/H/0406/003	3946/2011/03	NOVARTIS PHARMA GMBH	RO
Diovan 80 mg comprimate filmate	SE/H/0406/003	3946/2011/04	NOVARTIS PHARMA GMBH	RO
Diovan 80 mg comprimate filmate	SE/H/0406/003	3946/2011/05	NOVARTIS PHARMA GMBH	RO
Diovan 80 mg comprimate filmate	SE/H/0406/003	3946/2011/06	NOVARTIS PHARMA GMBH	RO
Diovan 80 mg comprimate filmate	SE/H/0406/003	3946/2011/07	NOVARTIS PHARMA GMBH	RO
Diovan 80 mg comprimate filmate	SE/H/0406/003	3946/2011/08	NOVARTIS PHARMA GMBH	RO
Diovan 80 mg comprimate filmate	SE/H/0406/003	3946/2011/09	NOVARTIS PHARMA GMBH	RO
Diovan 80 mg comprimate filmate	SE/H/0406/003	3946/2011/10	NOVARTIS PHARMA GMBH	RO
Diovan 80 mg comprimate filmate	SE/H/0406/003	3946/2011/11	NOVARTIS PHARMA GMBH	RO
Diovan 80 mg comprimate filmate	SE/H/0406/003	3946/2011/12	NOVARTIS PHARMA GMBH	RO
Diovan 80 mg comprimate filmate	SE/H/0406/003	3946/2011/13	NOVARTIS PHARMA GMBH	RO
Diovan 80 mg comprimate filmate	SE/H/0406/003	3946/2011/14	NOVARTIS PHARMA GMBH	RO
Diovan 80 mg comprimate filmate	SE/H/0406/003	3946/2011/15	NOVARTIS PHARMA GMBH	RO

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 80 mg comprimate filmate	SE/H/0406/003	3946/2011/16	NOVARTIS PHARMA GMBH	RO
Diovan 80 mg comprimate filmate	SE/H/0406/003	3946/2011/17	NOVARTIS PHARMA GMBH	RO
Diovan 80 mg comprimate filmate	SE/H/0406/003	3946/2011/18	NOVARTIS PHARMA GMBH	RO
Diovan 80 mg comprimate filmate	SE/H/0406/003	3946/2011/19	NOVARTIS PHARMA GMBH	RO
Diovan 80 mg comprimate filmate	SE/H/0406/003	3946/2011/20	NOVARTIS PHARMA GMBH	RO
Diovan 80 mg comprimate filmate	SE/H/0406/003	3946/2011/21	NOVARTIS PHARMA GMBH	RO
Diovan 80 mg comprimate filmate	SE/H/0406/003	3946/2011/22	NOVARTIS PHARMA GMBH	RO
Diovan 80 mg comprimate filmate	SE/H/0406/003	3946/2011/23	NOVARTIS PHARMA GMBH	RO
Diovan 80 mg comprimate filmate	SE/H/0406/003	3946/2011/24	NOVARTIS PHARMA GMBH	RO
Diovan 80 mg comprimate filmate	SE/H/0406/003	3946/2011/25	NOVARTIS PHARMA GMBH	RO
Diovan 80 mg comprimate filmate	SE/H/0406/003	3946/2011/26	NOVARTIS PHARMA GMBH	RO
Diovan 80 mg comprimate filmate	SE/H/0406/003	3946/2011/27	NOVARTIS PHARMA GMBH	RO
Diovan 80 mg comprimate filmate	SE/H/0406/003	3946/2011/28	NOVARTIS PHARMA GMBH	RO
Diovan 80 mg comprimate filmate	SE/H/0406/003	3946/2011/29	NOVARTIS PHARMA GMBH	RO
Diovan 80 mg comprimate filmate	SE/H/0406/003	3946/2011/30	NOVARTIS PHARMA GMBH	RO
Diován 80 mg comprimidos recubiertos con película	SE/H/0406/003	64.454	NOVARTIS FARMACÉUTICA S.A.	ES

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 80 mg comprimidos revestidos por película	SE/H/0406/003	3787488	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Diovan 80 mg comprimidos revestidos por película	SE/H/0406/003	3787587	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Diovan 80 mg comprimidos revestidos por película	SE/H/0406/003	3787686	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Diovan 80 mg comprimidos revestidos por película	SE/H/0406/003	3803988	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Diovan 80 mg comprimidos revestidos por película	SE/H/0406/003	3804085	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Diovan 80 mg comprimidos revestidos por película	SE/H/0406/003	5068325	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
DIOVAN 80 MG FILM-COATED TABLETS	SE/H/0406/003	PA 13/79/3	NOVARTIS PHARMACEUTICALS UK LIMITED	IE
DIOVAN 80 MG FILM-COATED TABLETS	SE/H/0406/003	088/00601	NOVARTIS PHARMACEUTICALS UK LIMITED	MT
Diovan 80 mg filmdragerade tabletter	SE/H/0406/003	16708	NOVARTIS FINLAND OY	FI
Diovan 80 mg filmdragerade tabletter	SE/H/0406/003	17494	NOVARTIS SVERIGE AB	SE
Diovan 80 mg filmom obložene tablete	SE/H/0406/004	HR-H-807143753-01	NOVARTIS HRVATSKA D.O.O.	HR
Diovan 80 mg filmom obložene tablete	SE/H/0406/004	HR-H-807143753-02	NOVARTIS HRVATSKA D.O.O.	HR
Diovan 80 mg filmom	SE/H/0406/004	HR-H-807143753-03	NOVARTIS HRVATSKA	HR

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
oblozene tablete			D.O.O.	
Diovan 80 mg filmom oblozene tablete	SE/H/0406/004	HR-H-807143753-04	NOVARTIS HRVATSKA D.O.O.	HR
Diovan 80 mg filmom oblozene tablete	SE/H/0406/004	HR-H-807143753-05	NOVARTIS HRVATSKA D.O.O.	HR
Diovan 80 mg filmom oblozene tablete	SE/H/0406/004	HR-H-807143753-06	NOVARTIS HRVATSKA D.O.O.	HR
Diovan 80 mg filmom oblozene tablete	SE/H/0406/004	HR-H-807143753-07	NOVARTIS HRVATSKA D.O.O.	HR
Diovan 80 mg filmom oblozene tablete	SE/H/0406/004	HR-H-807143753-08	NOVARTIS HRVATSKA D.O.O.	HR
Diovan 80 mg filmom oblozene tablete	SE/H/0406/004	HR-H-807143753-09	NOVARTIS HRVATSKA D.O.O.	HR
Diovan 80 mg filmom oblozene tablete	SE/H/0406/004	HR-H-807143753-10	NOVARTIS HRVATSKA D.O.O.	HR
Diovan 80 mg filmom oblozene tablete	SE/H/0406/004	HR-H-807143753-11	NOVARTIS HRVATSKA D.O.O.	HR
Diovan 80 mg filmom oblozene tablete	SE/H/0406/004	HR-H-807143753-12	NOVARTIS HRVATSKA D.O.O.	HR
Diovan 80 mg filmom oblozene tablete	SE/H/0406/004	HR-H-807143753-13	NOVARTIS HRVATSKA D.O.O.	HR
Diovan 80 mg filmom oblozene tablete	SE/H/0406/004	HR-H-807143753-14	NOVARTIS HRVATSKA D.O.O.	HR
Diovan 80 mg filmom oblozene tablete	SE/H/0406/004	HR-H-807143753-15	NOVARTIS HRVATSKA D.O.O.	HR
Diovan 80 mg filmsko oblozene tablete	SE/H/0406/003	H/03/00476/031	NOVARTIS PHARMA GMBH	SI
Diovan 80 mg filmsko oblozene tablete	SE/H/0406/003	H/03/00476/032	NOVARTIS PHARMA GMBH	SI
Diovan 80 mg filmsko oblozene tablete	SE/H/0406/003	H/03/00476/033	NOVARTIS PHARMA GMBH	SI
Diovan 80 mg filmsko oblozene tablete	SE/H/0406/003	H/03/00476/034	NOVARTIS PHARMA GMBH	SI

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 80 mg filmsko obložene tablete	SE/H/0406/003	H/03/00476/035	NOVARTIS PHARMA GMBH	SI
Diovan 80 mg filmsko obložene tablete	SE/H/0406/003	H/03/00476/036	NOVARTIS PHARMA GMBH	SI
Diovan 80 mg filmsko obložene tablete	SE/H/0406/003	H/03/00476/037	NOVARTIS PHARMA GMBH	SI
Diovan 80 mg filmsko obložene tablete	SE/H/0406/003	H/03/00476/038	NOVARTIS PHARMA GMBH	SI
Diovan 80 mg filmsko obložene tablete	SE/H/0406/003	H/03/00476/039	NOVARTIS PHARMA GMBH	SI
Diovan 80 mg filmsko obložene tablete	SE/H/0406/003	H/03/00476/040	NOVARTIS PHARMA GMBH	SI
Diovan 80 mg filmsko obložene tablete	SE/H/0406/003	H/03/00476/041	NOVARTIS PHARMA GMBH	SI
Diovan 80 mg filmsko obložene tablete	SE/H/0406/003	H/03/00476/042	NOVARTIS PHARMA GMBH	SI
Diovan 80 mg filmsko obložene tablete	SE/H/0406/003	H/03/00476/043	NOVARTIS PHARMA GMBH	SI
Diovan 80 mg filmsko obložene tablete	SE/H/0406/003	H/03/00476/044	NOVARTIS PHARMA GMBH	SI
Diovan 80 mg filmsko obložene tablete	SE/H/0406/003	H/03/00476/045	NOVARTIS PHARMA GMBH	SI
Diovan 80 mg filmsko obložene tablete	SE/H/0406/003	H/03/00476/046	NOVARTIS PHARMA GMBH	SI
Diovan 80 mg filmsko obložene tablete	SE/H/0406/003	H/03/00476/047	NOVARTIS PHARMA GMBH	SI
Diovan 80 mg filmsko obložene tablete	SE/H/0406/003	H/03/00476/048	NOVARTIS PHARMA GMBH	SI
Diovan 80 mg filmsko obložene tablete	SE/H/0406/003	H/03/00476/049	NOVARTIS PHARMA GMBH	SI
Diovan 80 mg filmsko obložene tablete	SE/H/0406/003	H/03/00476/050	NOVARTIS PHARMA GMBH	SI
Diovan 80 mg filmsko obložene tablete	SE/H/0406/003	H/03/00476/051	NOVARTIS PHARMA GMBH	SI

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
oblozene tablete				
Diovan 80 mg filmsko oblozene tablete	SE/H/0406/003	H/03/00476/052	NOVARTIS PHARMA GMBH	SI
Diovan 80 mg filmsko oblozene tablete	SE/H/0406/003	H/03/00476/053	NOVARTIS PHARMA GMBH	SI
Diovan 80 mg filmsko oblozene tablete	SE/H/0406/003	H/03/00476/054	NOVARTIS PHARMA GMBH	SI
Diovan 80 mg filmsko oblozene tablete	SE/H/0406/003	H/03/00476/055	NOVARTIS PHARMA GMBH	SI
Diovan 80 mg filmsko oblozene tablete	SE/H/0406/003	H/03/00476/056	NOVARTIS PHARMA GMBH	SI
Diovan 80 mg filmsko oblozene tablete	SE/H/0406/003	H/03/00476/057	NOVARTIS PHARMA GMBH	SI
Diovan 80 mg filmsko oblozene tablete	SE/H/0406/003	H/03/00476/058	NOVARTIS PHARMA GMBH	SI
Diovan 80 mg filmsko oblozene tablete	SE/H/0406/003	H/03/00476/059	NOVARTIS PHARMA GMBH	SI
Diovan 80 mg filmsko oblozene tablete	SE/H/0406/003	H/03/00476/060	NOVARTIS PHARMA GMBH	SI
Diovan 80 mg filmtabletta	SE/H/0406/003	OGYI-T-8484/09	NOVARTIS HUNGÁRIA KFT. PHARMA	HU
Diovan 80 mg filmtabletta	SE/H/0406/003	OGYI-T-8484/10	NOVARTIS HUNGÁRIA KFT. PHARMA	HU
Diovan 80 mg filmtabletta	SE/H/0406/003	OGYI-T-8484/11	NOVARTIS HUNGÁRIA KFT. PHARMA	HU
Diovan 80 mg filmtabletta	SE/H/0406/003	OGYI-T-8484/12	NOVARTIS HUNGÁRIA KFT. PHARMA	HU
DIOVAN 80 MG FILMTABLETTEN	SE/H/0406/003	1-24275	NOVARTIS PHARMA GMBH	AT
Diovan 80 mg filmuhúðaðar töflur	SE/H/0406/003	IS/1/01/042/01	NOVARTIS HEALTHCARE A/S	IS
Diovan 80 mg kalvopäällysteinen	SE/H/0406/003	16708	NOVARTIS FINLAND OY	FI

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
tabletti				
Diovan 80 mg plėvele dengtos tabletės	SE/H/0406/003	LT/1/01/1236/010	NOVARTIS FINLAND OY	LT
Diovan 80 mg plėvele dengtos tabletės	SE/H/0406/003	LT/1/01/1236/028	NOVARTIS FINLAND OY	LT
Diovan 80 mg plėvele dengtos tabletės	SE/H/0406/003	LT/1/01/1236/029	NOVARTIS FINLAND OY	LT
Diovan 80 mg plėvele dengtos tabletės	SE/H/0406/003	LT/1/01/1236/030	NOVARTIS FINLAND OY	LT
Diovan 80 mg plėvele dengtos tabletės	SE/H/0406/003	LT/1/01/1236/031	NOVARTIS FINLAND OY	LT
Diovan 80 mg plėvele dengtos tabletės	SE/H/0406/003	LT/1/01/1236/032	NOVARTIS FINLAND OY	LT
Diovan 80 mg plėvele dengtos tabletės	SE/H/0406/003	LT/1/01/1236/033	NOVARTIS FINLAND OY	LT
Diovan 80 mg plėvele dengtos tabletės	SE/H/0406/003	LT/1/01/1236/034	NOVARTIS FINLAND OY	LT
Diovan 80 mg plėvele dengtos tabletės	SE/H/0406/003	LT/1/01/1236/035	NOVARTIS FINLAND OY	LT
Diovan 80 mg plėvele dengtos tabletės	SE/H/0406/003	LT/1/01/1236/036	NOVARTIS FINLAND OY	LT
Diovan 80 mg plėvele dengtos tabletės	SE/H/0406/003	LT/1/01/1236/037	NOVARTIS FINLAND OY	LT
Diovan 80 mg plėvele dengtos tabletės	SE/H/0406/003	LT/1/01/1236/038	NOVARTIS FINLAND OY	LT
Diovan 80 mg plėvele dengtos tabletės	SE/H/0406/003	LT/1/01/1236/039	NOVARTIS FINLAND OY	LT
Diovan 80 mg plėvele dengtos tabletės	SE/H/0406/003	LT/1/01/1236/040	NOVARTIS FINLAND OY	LT
Diovan 80 mg plėvele dengtos tabletės	SE/H/0406/003	LT/1/01/1236/041	NOVARTIS FINLAND OY	LT
Diovan 80 mg tabletki powlekane	SE/H/0406/003	9291	NOVARTIS POLAND SP. Z O. O.	PL

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 80 mg tabletter, filmdrasjerte	SE/H/0406/003	00-6229	NOVARTIS NORGE AS	NO
DIOVAN 80, 80 MG FILMOMHULDE TABLETTEN	SE/H/0406/003	RVG 26939	NOVARTIS PHARMA B.V.	NL
Diován Cardio 40 mg comprimidos recubiertos con película	SE/H/0406/005	66.910	NOVARTIS FARMACÉUTICA S.A.	ES
Diovan Comp 160 mg/12,5 mg	SE/H/0565/002	35127	NOVARTIS HEALTHCARE A/S	DK
Diovan Comp 160 mg/12,5 mg	SE/H/0565/002	35127	NOVARTIS HEALTHCARE A/S	DK
Diovan Comp 160 mg/12,5 mg	SE/H/0565/002	35127	NOVARTIS HEALTHCARE A/S	DK
Diovan Comp 160 mg/12,5 mg	SE/H/0565/002	35127	NOVARTIS HEALTHCARE A/S	DK
Diovan Comp 160 mg/12,5 mg filmdragerade tabletter	SE/H/0565/002	19525	NOVARTIS SVERIGE AB	SE
Diovan Comp 160 mg/12,5 mg filmdragerade tabletter	SE/H/0565/002	19525	NOVARTIS SVERIGE AB	SE
Diovan Comp 160 mg/12,5 mg filmdragerade tabletter	SE/H/0565/002	19525	NOVARTIS SVERIGE AB	SE
Diovan Comp 160 mg/12,5 mg filmdragerade tabletter	SE/H/0565/002	19525	NOVARTIS SVERIGE AB	SE
Diovan Comp 160 mg/12,5 mg filmhúðaðar töflur	SE/H/0565/002	IS/1/03/029/01	NOVARTIS HEALTHCARE A/S	IS
Diovan Comp 160 mg/12,5 mg	SE/H/0565/002	IS/1/03/029/01	NOVARTIS HEALTHCARE A/S	IS

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
filmuhúðaðar töflur				
Diovan Comp 160 mg/12,5 mg filmuhúðaðar töflur	SE/H/0565/002	IS/1/03/029/01	NOVARTIS HEALTHCARE A/S	IS
Diovan Comp 160 mg/12,5 mg filmuhúðaðar töflur	SE/H/0565/002	IS/1/03/029/01	NOVARTIS HEALTHCARE A/S	IS
Diovan Comp 160 mg/12,5 mg kalvopäälysteinen tabletti	SE/H/0565/002	18231	NOVARTIS FINLAND OY	FI
Diovan Comp 160 mg/12,5 mg kalvopäälysteinen tabletti	SE/H/0565/002	18231	NOVARTIS FINLAND OY	FI
Diovan Comp 160 mg/12,5 mg kalvopäälysteinen tabletti	SE/H/0565/002	18231	NOVARTIS FINLAND OY	FI
Diovan Comp 160 mg/12,5 mg kalvopäälysteinen tabletti	SE/H/0565/002	18231	NOVARTIS FINLAND OY	FI
Diovan Comp 160 mg/25 mg	SE/H/0565/003	36787	NOVARTIS HEALTHCARE A/S	DK
Diovan Comp 160 mg/25 mg	SE/H/0565/003	36787	NOVARTIS HEALTHCARE A/S	DK
Diovan Comp 160 mg/25 mg	SE/H/0565/003	36787	NOVARTIS HEALTHCARE A/S	DK
Diovan Comp 160 mg/25 mg	SE/H/0565/003	36787	NOVARTIS HEALTHCARE A/S	DK
Diovan Comp 160 mg/25 mg filmdragerade	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
tabletter				
Diovan Comp 160 mg/25 mg filmdragerade tabletter	SE/H/0565/003	21204	NOVARTIS SVERIGE AB	SE
Diovan Comp 160 mg/25 mg filmdragerade tabletter	SE/H/0565/003	21204	NOVARTIS SVERIGE AB	SE
Diovan Comp 160 mg/25 mg filmdragerade tabletter	SE/H/0565/003	21204	NOVARTIS SVERIGE AB	SE
Diovan Comp 160 mg/25 mg filmhúðaðar töflur	SE/H/0565/003	IS/1/04/043/01	NOVARTIS HEALTHCARE A/S	IS
Diovan Comp 160 mg/25 mg filmhúðaðar töflur	SE/H/0565/003	IS/1/04/043/01	NOVARTIS HEALTHCARE A/S	IS
Diovan Comp 160 mg/25 mg filmhúðaðar töflur	SE/H/0565/003	IS/1/04/043/01	NOVARTIS HEALTHCARE A/S	IS
Diovan Comp 160 mg/25 mg filmhúðaðar töflur	SE/H/0565/003	IS/1/04/043/01	NOVARTIS HEALTHCARE A/S	IS
Diovan Comp 160 mg/25 mg kalvopäällysteinen tabletti	SE/H/0565/003	19618	NOVARTIS FINLAND OY	FI
Diovan Comp 160 mg/25 mg kalvopäällysteinen tabletti	SE/H/0565/003	19618	NOVARTIS FINLAND OY	FI
Diovan Comp 160 mg/25 mg kalvopäällysteinen tabletti	SE/H/0565/003	19618	NOVARTIS FINLAND OY	FI
Diovan Comp 160 mg/25 mg kalvopäällysteinen tabletti	SE/H/0565/003	19618	NOVARTIS FINLAND OY	FI
Diovan Comp 160 mg/25 mg tabletter, filmdrasjerte	SE/H/0565/003	01-10080	NOVARTIS NORGE AS	NO

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 160 mg/25 mg tabletter, filmdrasjerte	SE/H/0565/003	01-10080	NOVARTIS NORGE AS	NO
Diovan Comp 160 mg/25 mg tabletter, filmdrasjerte	SE/H/0565/003	01-10080	NOVARTIS NORGE AS	NO
Diovan Comp 160 mg/25 mg tabletter, filmdrasjerte	SE/H/0565/003	01-10080	NOVARTIS NORGE AS	NO
Diovan Comp 160 mg/12,5 mg filmdragerade tabletter	SE/H/0565/002	18231	NOVARTIS FINLAND OY	FI
Diovan Comp 160 mg/12,5 mg filmdragerade tabletter	SE/H/0565/002	18231	NOVARTIS FINLAND OY	FI
Diovan Comp 160 mg/12,5 mg filmdragerade tabletter	SE/H/0565/002	18231	NOVARTIS FINLAND OY	FI
Diovan Comp 160 mg/12,5 mg filmdragerade tabletter	SE/H/0565/002	18231	NOVARTIS FINLAND OY	FI
Diovan Comp 160 mg/25 mg filmdragerade tabletter	SE/H/0565/003	19618	NOVARTIS FINLAND OY	FI
Diovan Comp 160 mg/25 mg filmdragerade tabletter	SE/H/0565/003	19618	NOVARTIS FINLAND OY	FI
Diovan Comp 160 mg/25 mg filmdragerade tabletter	SE/H/0565/003	19618	NOVARTIS FINLAND OY	FI
Diovan Comp 160 mg/25 mg filmdragerade tabletter	SE/H/0565/003	19618	NOVARTIS FINLAND OY	FI

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	SE/H/0565/004	41588	NOVARTIS HEALTHCARE A/S	DK
Diovan Comp 320 mg/12,5 mg	SE/H/0565/004	41588	NOVARTIS HEALTHCARE A/S	DK
Diovan Comp 320 mg/12,5 mg	SE/H/0565/004	41588	NOVARTIS HEALTHCARE A/S	DK
Diovan Comp 320 mg/12,5 mg	SE/H/0565/004	41588	NOVARTIS HEALTHCARE A/S	DK
Diovan Comp 320 mg/12,5 mg filmdragerade tabletter	SE/H/0565/004	23641	NOVARTIS SVERIGE AB	SE
Diovan Comp 320 mg/12,5 mg filmdragerade tabletter	SE/H/0565/004	23641	NOVARTIS SVERIGE AB	SE
Diovan Comp 320 mg/12,5 mg filmdragerade tabletter	SE/H/0565/004	23641	NOVARTIS SVERIGE AB	SE
Diovan Comp 320 mg/12,5 mg filmdragerade tabletter	SE/H/0565/004	23641	NOVARTIS SVERIGE AB	SE
Diovan Comp 320 mg/12,5 mg filmuhúðaðar töflur	SE/H/0565/004	IS/1/07/051/01	NOVARTIS HEALTHCARE A/S	IS
Diovan Comp 320 mg/12,5 mg filmuhúðaðar töflur	SE/H/0565/004	IS/1/07/051/01	NOVARTIS HEALTHCARE A/S	IS
Diovan Comp 320 mg/12,5 mg filmuhúðaðar töflur	SE/H/0565/004	IS/1/07/051/01	NOVARTIS HEALTHCARE A/S	IS
Diovan Comp 320 mg/12,5 mg filmuhúðaðar töflur	SE/H/0565/004	IS/1/07/051/01	NOVARTIS HEALTHCARE A/S	IS
Diovan Comp 320	SE/H/0565/004	06-4074	NOVARTIS NORGE AS	NO

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
mg/12,5 mg tabletter, filmdrasjerte				
Diovan Comp 320 mg/12,5 mg tabletter, filmdrasjerte	SE/H/0565/004	06-4074	NOVARTIS NORGE AS	NO
Diovan Comp 320 mg/12,5 mg tabletter, filmdrasjerte	SE/H/0565/004	06-4074	NOVARTIS NORGE AS	NO
Diovan Comp 320 mg/12,5 mg tabletter, filmdrasjerte	SE/H/0565/004	06-4074	NOVARTIS NORGE AS	NO
Diovan Comp 320 mg/25 mg	SE/H/0565/005	41589	NOVARTIS HEALTHCARE A/S	DK
Diovan Comp 320 mg/25 mg	SE/H/0565/005	41589	NOVARTIS HEALTHCARE A/S	DK
Diovan Comp 320 mg/25 mg	SE/H/0565/005	41589	NOVARTIS HEALTHCARE A/S	DK
Diovan Comp 320 mg/25 mg	SE/H/0565/005	41589	NOVARTIS HEALTHCARE A/S	DK
Diovan Comp 320 mg/25 mg filmdragerade tabletter	SE/H/0565/005	23642	NOVARTIS SVERIGE AB	SE
Diovan Comp 320 mg/25 mg filmdragerade tabletter	SE/H/0565/005	23642	NOVARTIS SVERIGE AB	SE
Diovan Comp 320 mg/25 mg filmdragerade tabletter	SE/H/0565/005	23642	NOVARTIS SVERIGE AB	SE
Diovan Comp 320 mg/25 mg filmdragerade tabletter	SE/H/0565/005	23642	NOVARTIS SVERIGE AB	SE
Diovan Comp 320 mg/25 mg filmuhúðaðar töflur	SE/H/0565/005	IS/1/07/051/02	NOVARTIS HEALTHCARE A/S	IS

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/25 mg filmuhúðaðar töflur	SE/H/0565/005	IS/1/07/051/02	NOVARTIS HEALTHCARE A/S	IS
Diovan Comp 320 mg/25 mg filmuhúðaðar töflur	SE/H/0565/005	IS/1/07/051/02	NOVARTIS HEALTHCARE A/S	IS
Diovan Comp 320 mg/25 mg filmuhúðaðar töflur	SE/H/0565/005	IS/1/07/051/02	NOVARTIS HEALTHCARE A/S	IS
Diovan Comp 320 mg/25 mg kalvopäällysteinen tabletti	SE/H/0565/005	23724	NOVARTIS FINLAND OY	FI
Diovan Comp 320 mg/25 mg kalvopäällysteinen tabletti	SE/H/0565/005	23724	NOVARTIS FINLAND OY	FI
Diovan Comp 320 mg/25 mg kalvopäällysteinen tabletti	SE/H/0565/005	23724	NOVARTIS FINLAND OY	FI
Diovan Comp 320 mg/25 mg kalvopäällysteinen tabletti	SE/H/0565/005	23724	NOVARTIS FINLAND OY	FI
Diovan Comp 320 mg/25 mg tabletter, filmdrasjerte	SE/H/0565/005	06-4075	NOVARTIS NORGE AS	NO
Diovan Comp 320 mg/25 mg tabletter, filmdrasjerte	SE/H/0565/005	06-4075	NOVARTIS NORGE AS	NO
Diovan Comp 320 mg/25 mg tabletter, filmdrasjerte	SE/H/0565/005	06-4075	NOVARTIS NORGE AS	NO
Diovan Comp 320 mg/25 mg tabletter, filmdrasjerte	SE/H/0565/005	06-4075	NOVARTIS NORGE AS	NO
Diovan Comp 320 mg/12,5 mg filmdragerade tabletter	SE/H/0565/004	23723	NOVARTIS FINLAND OY	FI

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	23723	NOVARTIS FINLAND OY	FI
Diovan Comp 320 mg/12,5 mg filmdragerade tabletter	SE/H/0565/004	23723	NOVARTIS FINLAND OY	FI
Diovan Comp 320 mg/12,5 mg filmdragerade tabletter	SE/H/0565/004	23723	NOVARTIS FINLAND OY	FI
Diovan Comp 320 mg/25 mg filmdragerade tabletter	SE/H/0565/005	23724	NOVARTIS FINLAND OY	FI
Diovan Comp 320 mg/25 mg filmdragerade tabletter	SE/H/0565/005	23724	NOVARTIS FINLAND OY	FI
Diovan Comp 320 mg/25 mg filmdragerade tabletter	SE/H/0565/005	23724	NOVARTIS FINLAND OY	FI
Diovan Comp 320 mg/25 mg filmdragerade tabletter	SE/H/0565/005	23724	NOVARTIS FINLAND OY	FI
Diovan Comp 80 mg/12,5 mg	SE/H/0565/001	19338	NOVARTIS HEALTHCARE A/S	DK
Diovan Comp 80 mg/12,5 mg	SE/H/0565/001	19338	NOVARTIS HEALTHCARE A/S	DK
Diovan Comp 80 mg/12,5 mg	SE/H/0565/001	19338	NOVARTIS HEALTHCARE A/S	DK
Diovan Comp 80 mg/12,5 mg	SE/H/0565/001	19338	NOVARTIS HEALTHCARE A/S	DK
Diovan Comp 80 mg/12,5 mg filmdragerade tabletter	SE/H/0565/001	14098	NOVARTIS SVERIGE AB	SE
Diovan Comp 80	SE/H/0565/001	14098	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
mg/12,5 mg filmdragerade tabletter				
Diovan Comp 80 mg/12,5 mg filmdragerade tabletter	SE/H/0565/001	14098	NOVARTIS SVERIGE AB	SE
Diovan Comp 80 mg/12,5 mg filmdragerade tabletter	SE/H/0565/001	14098	NOVARTIS SVERIGE AB	SE
Diovan Comp 80 mg/12,5 mg filmhúðaðar töflur	SE/H/0565/001	980160	NOVARTIS HEALTHCARE A/S	IS
Diovan Comp 80 mg/12,5 mg filmhúðaðar töflur	SE/H/0565/001	980160	NOVARTIS HEALTHCARE A/S	IS
Diovan Comp 80 mg/12,5 mg filmhúðaðar töflur	SE/H/0565/001	980160	NOVARTIS HEALTHCARE A/S	IS
Diovan Comp 80 mg/12,5 mg filmhúðaðar töflur	SE/H/0565/001	980160	NOVARTIS HEALTHCARE A/S	IS
Diovan Comp 80 mg/12,5 mg kalvopäällysteinen tabletti	SE/H/0565/001	13181	NOVARTIS FINLAND OY	FI
Diovan Comp 80 mg/12,5 mg kalvopäällysteinen tabletti	SE/H/0565/001	13181	NOVARTIS FINLAND OY	FI
Diovan Comp 80 mg/12,5 mg kalvopäällysteinen tabletti	SE/H/0565/001	13181	NOVARTIS FINLAND OY	FI
Diovan Comp 80	SE/H/0565/001	13181	NOVARTIS FINLAND OY	FI

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
mg/12,5 mg kalvopäällysteinen tabletti				
Diovan Comp 80 mg/12,5 mg filmdragerade tabletter	SE/H/0565/001	13181	NOVARTIS FINLAND OY	FI
Diovan Comp 80 mg/12,5 mg filmdragerade tabletter	SE/H/0565/001	13181	NOVARTIS FINLAND OY	FI
Diovan Comp 80 mg/12,5 mg filmdragerade tabletter	SE/H/0565/001	13181	NOVARTIS FINLAND OY	FI
Diovan Comp 80 mg/12,5 mg filmdragerade tabletter	SE/H/0565/001	13181	NOVARTIS FINLAND OY	FI
Diovan HCT 160/12,5 mg filmdragerade tabletter	SE/H/0565/002	OGYI-T-6552/05	NOVARTIS HUNGÁRIA KFT. PHARMA	HU
Diovan HCT 160/12,5 mg filmdragerade tabletter	SE/H/0565/002	OGYI-T-6552/07	NOVARTIS HUNGÁRIA KFT. PHARMA	HU
Diovan HCT 160/12,5 mg filmdragerade tabletter	SE/H/0565/002	OGYI-T-6552/08	NOVARTIS HUNGÁRIA KFT. PHARMA	HU
Diovan HCT 160/12,5 mg filmdragerade tabletter	SE/H/0565/002	OGYI-T-6552/06	NOVARTIS HUNGÁRIA KFT. PHARMA	HU
Diovan HCT 160/12,5 mg filmdragerade tabletter	SE/H/0565/002	OGYI-T-6552/05	NOVARTIS HUNGÁRIA KFT. PHARMA	HU
Diovan HCT 160/12,5 mg filmdragerade tabletter	SE/H/0565/002	OGYI-T-6552/07	NOVARTIS HUNGÁRIA KFT. PHARMA	HU
Diovan HCT 160/12,5 mg filmdragerade tabletter	SE/H/0565/002	OGYI-T-6552/08	NOVARTIS HUNGÁRIA KFT. PHARMA	HU
Diovan HCT 160/12,5 mg filmdragerade tabletter	SE/H/0565/002	OGYI-T-6552/06	NOVARTIS HUNGÁRIA KFT. PHARMA	HU
Diovan HCT 160/12,5 mg filmdragerade tabletter	SE/H/0565/002	OGYI-T-6552/05	NOVARTIS HUNGÁRIA KFT. PHARMA	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
Diovan HCT 160/12,5 mg filmdabletta	SE/H/0565/002	OGYI-T-6552/07	NOVARTIS HUNGÁRIA KFT. PHARMA	HU
Diovan HCT 160/12,5 mg filmdabletta	SE/H/0565/002	OGYI-T-6552/08	NOVARTIS HUNGÁRIA KFT. PHARMA	HU
Diovan HCT 160/12,5 mg filmdabletta	SE/H/0565/002	OGYI-T-6552/06	NOVARTIS HUNGÁRIA KFT. PHARMA	HU
Diovan HCT 160/12,5 mg filmdabletta	SE/H/0565/002	OGYI-T-6552/05	NOVARTIS HUNGÁRIA KFT. PHARMA	HU
Diovan HCT 160/12,5 mg filmdabletta	SE/H/0565/002	OGYI-T-6552/07	NOVARTIS HUNGÁRIA KFT. PHARMA	HU
Diovan HCT 160/12,5 mg filmdabletta	SE/H/0565/002	OGYI-T-6552/08	NOVARTIS HUNGÁRIA KFT. PHARMA	HU
Diovan HCT 160/12,5 mg filmdabletta	SE/H/0565/002	OGYI-T-6552/06	NOVARTIS HUNGÁRIA KFT. PHARMA	HU
Diovan HCT 160/25 mg filmdabletta	SE/H/0565/003	OGYI-T-6552/09	NOVARTIS HUNGÁRIA KFT. PHARMA	HU
Diovan HCT 160/25 mg filmdabletta	SE/H/0565/003	OGYI-T-6552/10	NOVARTIS HUNGÁRIA KFT. PHARMA	HU
Diovan HCT 160/25 mg filmdabletta	SE/H/0565/003	OGYI-T-6552/09	NOVARTIS HUNGÁRIA KFT. PHARMA	HU
Diovan HCT 160/25 mg filmdabletta	SE/H/0565/003	OGYI-T-6552/10	NOVARTIS HUNGÁRIA KFT. PHARMA	HU
Diovan HCT 160/25 mg filmdabletta	SE/H/0565/003	OGYI-T-6552/09	NOVARTIS HUNGÁRIA KFT. PHARMA	HU
Diovan HCT 160/25 mg filmdabletta	SE/H/0565/003	OGYI-T-6552/10	NOVARTIS HUNGÁRIA KFT. PHARMA	HU
Diovan HCT 160/25 mg filmdabletta	SE/H/0565/003	OGYI-T-6552/09	NOVARTIS HUNGÁRIA KFT. PHARMA	HU
Diovan HCT 160/25 mg filmdabletta	SE/H/0565/003	OGYI-T-6552/10	NOVARTIS HUNGÁRIA KFT. PHARMA	HU
Diovan HCT 160/25mg filmdabletta	SE/H/0565/003	OGYI-T-6552/11	NOVARTIS HUNGÁRIA KFT. PHARMA	HU
Diovan HCT 160/25mg	SE/H/0565/003	OGYI-T-6552/12	NOVARTIS HUNGÁRIA 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
filmdobletta			PHARMA	
Diovan HCT 160/25mg filmdobletta	SE/H/0565/003	OGYI-T-6552/11	NOVARTIS HUNGÁRIA KFT. PHARMA	HU
Diovan HCT 160/25mg filmdobletta	SE/H/0565/003	OGYI-T-6552/12	NOVARTIS HUNGÁRIA KFT. PHARMA	HU
Diovan HCT 160/25mg filmdobletta	SE/H/0565/003	OGYI-T-6552/11	NOVARTIS HUNGÁRIA KFT. PHARMA	HU
Diovan HCT 160/25mg filmdobletta	SE/H/0565/003	OGYI-T-6552/12	NOVARTIS HUNGÁRIA KFT. PHARMA	HU
Diovan HCT 160/25mg filmdobletta	SE/H/0565/003	OGYI-T-6552/11	NOVARTIS HUNGÁRIA KFT. PHARMA	HU
Diovan HCT 160/25mg filmdobletta	SE/H/0565/003	OGYI-T-6552/12	NOVARTIS HUNGÁRIA KFT. PHARMA	HU
Diovan HCT 80/12,5 mg filmdobletta	SE/H/0565/001	OGYI-T-6552/01	NOVARTIS HUNGÁRIA KFT. PHARMA	HU
Diovan HCT 80/12,5 mg filmdobletta	SE/H/0565/001	OGYI-T-6552/02	NOVARTIS HUNGÁRIA KFT. PHARMA	HU
Diovan HCT 80/12,5 mg filmdobletta	SE/H/0565/001	OGYI-T-6552/03	NOVARTIS HUNGÁRIA KFT. PHARMA	HU
Diovan HCT 80/12,5 mg filmdobletta	SE/H/0565/001	OGYI-T-6552/04	NOVARTIS HUNGÁRIA KFT. PHARMA	HU
Diovan HCT 80/12,5 mg filmdobletta	SE/H/0565/001	OGYI-T-6552/01	NOVARTIS HUNGÁRIA KFT. PHARMA	HU
Diovan HCT 80/12,5 mg filmdobletta	SE/H/0565/001	OGYI-T-6552/02	NOVARTIS HUNGÁRIA KFT. PHARMA	HU
Diovan HCT 80/12,5 mg filmdobletta	SE/H/0565/001	OGYI-T-6552/03	NOVARTIS HUNGÁRIA KFT. PHARMA	HU
Diovan HCT 80/12,5 mg filmdobletta	SE/H/0565/001	OGYI-T-6552/04	NOVARTIS HUNGÁRIA KFT. PHARMA	HU
Diovan HCT 80/12,5 mg filmdobletta	SE/H/0565/001	OGYI-T-6552/01	NOVARTIS HUNGÁRIA KFT. PHARMA	HU
Diovan HCT 80/12,5 mg filmdobletta	SE/H/0565/001	OGYI-T-6552/02	NOVARTIS HUNGÁRIA KFT. PHARMA	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
Diovan HCT 80/12,5 mg filmdabletta	SE/H/0565/001	OGYI-T-6552/03	NOVARTIS HUNGÁRIA KFT. PHARMA	HU
Diovan HCT 80/12,5 mg filmdabletta	SE/H/0565/001	OGYI-T-6552/04	NOVARTIS HUNGÁRIA KFT. PHARMA	HU
Diovan HCT 80/12,5 mg filmdabletta	SE/H/0565/001	OGYI-T-6552/01	NOVARTIS HUNGÁRIA KFT. PHARMA	HU
Diovan HCT 80/12,5 mg filmdabletta	SE/H/0565/001	OGYI-T-6552/02	NOVARTIS HUNGÁRIA KFT. PHARMA	HU
Diovan HCT 80/12,5 mg filmdabletta	SE/H/0565/001	OGYI-T-6552/03	NOVARTIS HUNGÁRIA KFT. PHARMA	HU
Diovan HCT 80/12,5 mg filmdabletta	SE/H/0565/001	OGYI-T-6552/04	NOVARTIS HUNGÁRIA KFT. PHARMA	HU
DIOVAN ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0406/004	19385	NOVARTIS PHARMACEUTICALS UK LIMITED	CY
DIOVAN ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0406/004	232870401	NOVARTIS (HELLAS) S.A.C.I.	GR
DIOVAN ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0406/004	232870402	NOVARTIS (HELLAS) S.A.C.I.	GR
DIOVAN ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0406/004	232870403	NOVARTIS (HELLAS) S.A.C.I.	GR
DIOVAN ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0406/004	232870404	NOVARTIS (HELLAS) S.A.C.I.	GR
DIOVAN ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0406/004	232870405	NOVARTIS (HELLAS) S.A.C.I.	GR
DIOVAN ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0406/004	232870406	NOVARTIS (HELLAS) S.A.C.I.	GR

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 ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0406/004	232870407	NOVARTIS (HELLAS) S.A.C.I.	GR
DIOVAN ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0406/004	232870408	NOVARTIS (HELLAS) S.A.C.I.	GR
DIOVAN ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0406/004	232870409	NOVARTIS (HELLAS) S.A.C.I.	GR
DIOVAN ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0406/004	232870410	NOVARTIS (HELLAS) S.A.C.I.	GR
DIOVAN ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0406/004	232870411	NOVARTIS (HELLAS) S.A.C.I.	GR
DIOVAN ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0406/004	232870412	NOVARTIS (HELLAS) S.A.C.I.	GR
DIOVAN ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0406/004	232870413	NOVARTIS (HELLAS) S.A.C.I.	GR
DIOVAN ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0406/004	232870414	NOVARTIS (HELLAS) S.A.C.I.	GR
DIOVAN ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0406/004	232870415	NOVARTIS (HELLAS) S.A.C.I.	GR
DIOVAN ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0406/004	232870416	NOVARTIS (HELLAS) S.A.C.I.	GR
DIOVAN ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0406/004	232870417	NOVARTIS (HELLAS) S.A.C.I.	GR

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 ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0406/004	232870418	NOVARTIS (HELLAS) S.A.C.I.	GR
DIOVAN ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0406/004	232870419	NOVARTIS (HELLAS) S.A.C.I.	GR
DIOVAN ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0406/004	232870420	NOVARTIS (HELLAS) S.A.C.I.	GR
DIOVAN ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0406/004	232870421	NOVARTIS (HELLAS) S.A.C.I.	GR
DIOVAN ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0406/004	232870422	NOVARTIS (HELLAS) S.A.C.I.	GR
DIOVAN ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0406/004	232870423	NOVARTIS (HELLAS) S.A.C.I.	GR
DIOVAN ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0406/004	232870424	NOVARTIS (HELLAS) S.A.C.I.	GR
DIOVAN ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0406/004	232870425	NOVARTIS (HELLAS) S.A.C.I.	GR
DIOVAN ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0406/004	232870426	NOVARTIS (HELLAS) S.A.C.I.	GR
DIOVAN ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0406/004	232870427	NOVARTIS (HELLAS) S.A.C.I.	GR
DIOVAN ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0406/004	232870428	NOVARTIS (HELLAS) S.A.C.I.	GR

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
ΔΙΟΒΑΝ ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0406/004	232870429	NOVARTIS (HELLAS) S.A.C.I.	GR
ΔΙΟΒΑΝ ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0406/004	232870430	NOVARTIS (HELLAS) S.A.C.I.	GR
ΔΙΟΒΑΝ ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0406/004	232870431	NOVARTIS (HELLAS) S.A.C.I.	GR
ΔΙΟΒΑΝ ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 160 MG	SE/H/0406/004	232870432	NOVARTIS (HELLAS) S.A.C.I.	GR
ΔΙΟΒΑΝ ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0406/006	20375	NOVARTIS PHARMACEUTICALS UK LIMITED	CY
ΔΙΟΒΑΝ ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0406/006	232870601	NOVARTIS (HELLAS) S.A.C.I.	GR
ΔΙΟΒΑΝ ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0406/006	232870602	NOVARTIS (HELLAS) S.A.C.I.	GR
ΔΙΟΒΑΝ ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0406/006	232870603	NOVARTIS (HELLAS) S.A.C.I.	GR
ΔΙΟΒΑΝ ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0406/006	232870604	NOVARTIS (HELLAS) S.A.C.I.	GR
ΔΙΟΒΑΝ ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0406/006	232870605	NOVARTIS (HELLAS) S.A.C.I.	GR
ΔΙΟΒΑΝ ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0406/006	232870606	NOVARTIS (HELLAS) S.A.C.I.	GR

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 ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0406/006	232870607	NOVARTIS (HELLAS) S.A.C.I.	GR
DIOVAN ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0406/006	232870608	NOVARTIS (HELLAS) S.A.C.I.	GR
DIOVAN ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0406/006	232870609	NOVARTIS (HELLAS) S.A.C.I.	GR
DIOVAN ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0406/006	232870610	NOVARTIS (HELLAS) S.A.C.I.	GR
DIOVAN ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0406/006	232870611	NOVARTIS (HELLAS) S.A.C.I.	GR
DIOVAN ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0406/006	232870612	NOVARTIS (HELLAS) S.A.C.I.	GR
DIOVAN ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0406/006	232870613	NOVARTIS (HELLAS) S.A.C.I.	GR
DIOVAN ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0406/006	232870614	NOVARTIS (HELLAS) S.A.C.I.	GR
DIOVAN ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0406/006	232870615	NOVARTIS (HELLAS) S.A.C.I.	GR
DIOVAN ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0406/006	232870616	NOVARTIS (HELLAS) S.A.C.I.	GR
DIOVAN ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0406/006	232870617	NOVARTIS (HELLAS) S.A.C.I.	GR

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 ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0406/006	232870618	NOVARTIS (HELLAS) S.A.C.I.	GR
DIOVAN ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0406/006	232870619	NOVARTIS (HELLAS) S.A.C.I.	GR
DIOVAN ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0406/006	232870620	NOVARTIS (HELLAS) S.A.C.I.	GR
DIOVAN ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0406/006	232870621	NOVARTIS (HELLAS) S.A.C.I.	GR
DIOVAN ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0406/006	232870622	NOVARTIS (HELLAS) S.A.C.I.	GR
DIOVAN ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0406/006	232870623	NOVARTIS (HELLAS) S.A.C.I.	GR
DIOVAN ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0406/006	232870624	NOVARTIS (HELLAS) S.A.C.I.	GR
DIOVAN ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0406/006	232870625	NOVARTIS (HELLAS) S.A.C.I.	GR
DIOVAN ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0406/006	232870626	NOVARTIS (HELLAS) S.A.C.I.	GR
DIOVAN ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0406/006	232870627	NOVARTIS (HELLAS) S.A.C.I.	GR
DIOVAN ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0406/006	232870628	NOVARTIS (HELLAS) S.A.C.I.	GR

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
ΔΙΟΒΑΝ ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0406/006	232870629	NOVARTIS (HELLAS) S.A.C.I.	GR
ΔΙΟΒΑΝ ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0406/006	232870630	NOVARTIS (HELLAS) S.A.C.I.	GR
ΔΙΟΒΑΝ ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0406/006	232870631	NOVARTIS (HELLAS) S.A.C.I.	GR
ΔΙΟΒΑΝ ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 320 MG	SE/H/0406/006	232870632	NOVARTIS (HELLAS) S.A.C.I.	GR
ΔΙΟΒΑΝ ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0406/005	19635	NOVARTIS PHARMACEUTICALS UK LIMITED	CY
ΔΙΟΒΑΝ ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0406/005	232870501	NOVARTIS (HELLAS) S.A.C.I.	GR
ΔΙΟΒΑΝ ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0406/005	232870502	NOVARTIS (HELLAS) S.A.C.I.	GR
ΔΙΟΒΑΝ ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0406/005	232870503	NOVARTIS (HELLAS) S.A.C.I.	GR
ΔΙΟΒΑΝ ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0406/005	232870504	NOVARTIS (HELLAS) S.A.C.I.	GR
ΔΙΟΒΑΝ ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0406/005	232870505	NOVARTIS (HELLAS) S.A.C.I.	GR
ΔΙΟΒΑΝ ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0406/005	232870506	NOVARTIS (HELLAS) S.A.C.I.	GR

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
ΔΙΟΒΑΝ ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0406/005	232870507	NOVARTIS (HELLAS) S.A.C.I.	GR
ΔΙΟΒΑΝ ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0406/005	232870508	NOVARTIS (HELLAS) S.A.C.I.	GR
ΔΙΟΒΑΝ ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0406/005	232870509	NOVARTIS (HELLAS) S.A.C.I.	GR
ΔΙΟΒΑΝ ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0406/005	232870510	NOVARTIS (HELLAS) S.A.C.I.	GR
ΔΙΟΒΑΝ ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0406/005	232870511	NOVARTIS (HELLAS) S.A.C.I.	GR
ΔΙΟΒΑΝ ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0406/005	232870512	NOVARTIS (HELLAS) S.A.C.I.	GR
ΔΙΟΒΑΝ ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0406/005	232870513	NOVARTIS (HELLAS) S.A.C.I.	GR
ΔΙΟΒΑΝ ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0406/005	232870514	NOVARTIS (HELLAS) S.A.C.I.	GR
ΔΙΟΒΑΝ ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0406/005	232870515	NOVARTIS (HELLAS) S.A.C.I.	GR
ΔΙΟΒΑΝ ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0406/005	232870516	NOVARTIS (HELLAS) S.A.C.I.	GR
ΔΙΟΒΑΝ ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0406/005	232870517	NOVARTIS (HELLAS) S.A.C.I.	GR

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
ΔΙΟΒΑΝ ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0406/005	232870518	NOVARTIS (HELLAS) S.A.C.I.	GR
ΔΙΟΒΑΝ ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0406/005	232870519	NOVARTIS (HELLAS) S.A.C.I.	GR
ΔΙΟΒΑΝ ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0406/005	232870520	NOVARTIS (HELLAS) S.A.C.I.	GR
ΔΙΟΒΑΝ ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0406/005	232870521	NOVARTIS (HELLAS) S.A.C.I.	GR
ΔΙΟΒΑΝ ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0406/005	232870522	NOVARTIS (HELLAS) S.A.C.I.	GR
ΔΙΟΒΑΝ ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0406/005	232870523	NOVARTIS (HELLAS) S.A.C.I.	GR
ΔΙΟΒΑΝ ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0406/005	232870524	NOVARTIS (HELLAS) S.A.C.I.	GR
ΔΙΟΒΑΝ ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0406/005	232870525	NOVARTIS (HELLAS) S.A.C.I.	GR
ΔΙΟΒΑΝ ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0406/005	232870526	NOVARTIS (HELLAS) S.A.C.I.	GR
ΔΙΟΒΑΝ ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0406/005	232870527	NOVARTIS (HELLAS) S.A.C.I.	GR
ΔΙΟΒΑΝ ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0406/005	232870528	NOVARTIS (HELLAS) S.A.C.I.	GR

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
ΔΙΟΒΑΝ ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0406/005	232870529	NOVARTIS (HELLAS) S.A.C.I.	GR
ΔΙΟΒΑΝ ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0406/005	232870530	NOVARTIS (HELLAS) S.A.C.I.	GR
ΔΙΟΒΑΝ ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0406/005	232870531	NOVARTIS (HELLAS) S.A.C.I.	GR
ΔΙΟΒΑΝ ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 40 MG	SE/H/0406/005	232870532	NOVARTIS (HELLAS) S.A.C.I.	GR
ΔΙΟΒΑΝ ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0406/003	19384	NOVARTIS PHARMACEUTICALS UK LIMITED	CY
ΔΙΟΒΑΝ ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0406/003	232870315	NOVARTIS (HELLAS) S.A.C.I.	GR
ΔΙΟΒΑΝ ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0406/003	232870316	NOVARTIS (HELLAS) S.A.C.I.	GR
ΔΙΟΒΑΝ ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0406/003	232870317	NOVARTIS (HELLAS) S.A.C.I.	GR
ΔΙΟΒΑΝ ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0406/003	232870318	NOVARTIS (HELLAS) S.A.C.I.	GR
ΔΙΟΒΑΝ ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0406/003	232870319	NOVARTIS (HELLAS) S.A.C.I.	GR
ΔΙΟΒΑΝ ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0406/003	232870320	NOVARTIS (HELLAS) S.A.C.I.	GR

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
ΔΙΟΒΑΝ ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0406/003	232870321	NOVARTIS (HELLAS) S.A.C.I.	GR
ΔΙΟΒΑΝ ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0406/003	232870322	NOVARTIS (HELLAS) S.A.C.I.	GR
ΔΙΟΒΑΝ ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0406/003	232870323	NOVARTIS (HELLAS) S.A.C.I.	GR
ΔΙΟΒΑΝ ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0406/003	232870324	NOVARTIS (HELLAS) S.A.C.I.	GR
ΔΙΟΒΑΝ ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0406/003	232870325	NOVARTIS (HELLAS) S.A.C.I.	GR
ΔΙΟΒΑΝ ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0406/003	232870326	NOVARTIS (HELLAS) S.A.C.I.	GR
ΔΙΟΒΑΝ ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0406/003	232870327	NOVARTIS (HELLAS) S.A.C.I.	GR
ΔΙΟΒΑΝ ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0406/003	232870328	NOVARTIS (HELLAS) S.A.C.I.	GR
ΔΙΟΒΑΝ ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0406/003	232870329	NOVARTIS (HELLAS) S.A.C.I.	GR
ΔΙΟΒΑΝ ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0406/003	232870330	NOVARTIS (HELLAS) S.A.C.I.	GR
ΔΙΟΒΑΝ ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0406/003	232870331	NOVARTIS (HELLAS) S.A.C.I.	GR

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
ΔΙΟΒΑΝ ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0406/003	232870332	NOVARTIS (HELLAS) S.A.C.I.	GR
ΔΙΟΒΑΝ ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0406/003	232870301	NOVARTIS (HELLAS) S.A.C.I.	GR
ΔΙΟΒΑΝ ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0406/003	232870302	NOVARTIS (HELLAS) S.A.C.I.	GR
ΔΙΟΒΑΝ ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0406/003	232870303	NOVARTIS (HELLAS) S.A.C.I.	GR
ΔΙΟΒΑΝ ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0406/003	232870304	NOVARTIS (HELLAS) S.A.C.I.	GR
ΔΙΟΒΑΝ ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0406/003	232870305	NOVARTIS (HELLAS) S.A.C.I.	GR
ΔΙΟΒΑΝ ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0406/003	232870306	NOVARTIS (HELLAS) S.A.C.I.	GR
ΔΙΟΒΑΝ ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0406/003	232870307	NOVARTIS (HELLAS) S.A.C.I.	GR
ΔΙΟΒΑΝ ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0406/003	232870308	NOVARTIS (HELLAS) S.A.C.I.	GR
ΔΙΟΒΑΝ ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0406/003	232870309	NOVARTIS (HELLAS) S.A.C.I.	GR
ΔΙΟΒΑΝ ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0406/003	232870310	NOVARTIS (HELLAS) S.A.C.I.	GR

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 ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0406/003	232870311	NOVARTIS (HELLAS) S.A.C.I.	GR
DIOVAN ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0406/003	232870312	NOVARTIS (HELLAS) S.A.C.I.	GR
DIOVAN ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0406/003	232870313	NOVARTIS (HELLAS) S.A.C.I.	GR
DIOVAN ΕΠΙΚΑΛΥΜΜΕΝΑ ΜΕ ΛΕΠΤΟ ΥΜΕΝΙΟ ΔΙΣΚΙΑ 80 MG	SE/H/0406/003	232870314	NOVARTIS (HELLAS) S.A.C.I.	GR
DIOVAN ΠΟΣΙΜΟ ΔΙΑΛΥΜΑ 3 MG/ML	SE/H/0406/007	20694	NOVARTIS PHARMACEUTICALS UK LIMITED	CY
DIOVAN ΠΟΣΙΜΟ ΔΙΑΛΥΜΑ 3 MG/ML	SE/H/0406/007	232870701	NOVARTIS (HELLAS) S.A.C.I.	GR
DIOVAN COMP 320 MG/12,5 MG KALVOPÄÄLLYSTEINEN TABLETTI	SE/H/0565/004	23723	NOVARTIS FINLAND OY	FI
DIOVAN COMP 320 MG/12,5 MG KALVOPÄÄLLYSTEINEN TABLETTI	SE/H/0565/004	23723	NOVARTIS FINLAND OY	FI
DIOVAN COMP 320 MG/12,5 MG KALVOPÄÄLLYSTEINEN TABLETTI	SE/H/0565/004	23723	NOVARTIS FINLAND OY	FI
DIOVAN COMP 320 MG/12,5 MG KALVOPÄÄLLYSTEINEN TABLETTI	SE/H/0565/004	23723	NOVARTIS FINLAND OY	FI

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, 160 mg õhukese polümeerikattega tabletid	SE/H/0406/004	375902	NOVARTIS FINLAND OY	EE
Diovan, 320 mg õhukese polümeerikattega tabletid	SE/H/0406/006	514806	NOVARTIS FINLAND OY	EE
Diovan, 40 mg õhukese polümeerikattega tabletid	SE/H/0406/005	452504	NOVARTIS FINLAND OY	EE
Diovan, 80 mg õhukese polümeerikattega tabletid	SE/H/0406/003	376002	NOVARTIS FINLAND OY	EE
Diovan, oral opløsning	SE/H/0406/007	46831	NOVARTIS HEALTHCARE A/S	DK
DIOVAN® 160 mg protect Filmdrasetten	SE/H/0406/004	50477.01.00	NOVARTIS PHARMA GMBH	DE
DIOVAN® 160 mg tabletter, filmdrasjerte	SE/H/0406/004	00-6230	NOVARTIS NORGE AS	NO
DIOVAN® 3 mg/ml Lösung zum Einnehmen	SE/H/0406/007	82267.00.00	NOVARTIS PHARMA GMBH	DE
DIOVAN® 320 mg forte Filmdrasetten	SE/H/0406/006	67298.00.00	NOVARTIS PHARMA GMBH	DE
DIOVAN® 40 mg Filmdrasetten	SE/H/0406/005	50477.02.00	NOVARTIS PHARMA GMBH	DE
DIOVAN® 80 mg Filmdrasetten	SE/H/0406/003	50477.00.00	NOVARTIS PHARMA GMBH	DE
Diovan® Comp 160 mg/12,5 mg tabletter, filmdrasjerte	SE/H/0565/002	01-10079	NOVARTIS NORGE AS	NO
Diovan® Comp 160 mg/12,5 mg tabletter, filmdrasjerte	SE/H/0565/002	01-10079	NOVARTIS NORGE AS	NO
Diovan® Comp 160 mg/12,5 mg tabletter,	SE/H/0565/002	01-10079	NOVARTIS NORGE AS	NO

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
filmdrasjerte				
Diovan® Comp 160 mg/12,5 mg tabletter, filmdrasjerte	SE/H/0565/002	01-10079	NOVARTIS NORGE AS	NO
Diovan® Comp 80 mg/12,5 mg tabletter, filmdrasjerte	SE/H/0565/001	97-2716	NOVARTIS NORGE AS	NO
Diovan® Comp 80 mg/12,5 mg tabletter, filmdrasjerte	SE/H/0565/001	97-2716	NOVARTIS NORGE AS	NO
Diovan® Comp 80 mg/12,5 mg tabletter, filmdrasjerte	SE/H/0565/001	97-2716	NOVARTIS NORGE AS	NO
Diovan® Comp 80 mg/12,5 mg tabletter, filmdrasjerte	SE/H/0565/001	97-2716	NOVARTIS NORGE AS	NO
DIOVAN320 MG COMPRIMIDOS REVESTIDOS POR PELÍCULA	SE/H/0406/006	5018874	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
DIOVAN320 MG COMPRIMIDOS REVESTIDOS POR PELÍCULA	SE/H/0406/006	5018908	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
DIOVAN320 MG COMPRIMIDOS REVESTIDOS POR PELÍCULA	SE/H/0406/006	5018916	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
DIOVAN320 MG COMPRIMIDOS REVESTIDOS POR PELÍCULA	SE/H/0406/006	5018924	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
DIOVAN320 MG	SE/H/0406/006	5018932	NOVARTIS FARMA -	PT

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
COMPRIMIDOS REVESTIDOS POR PELÍCULA			PRODUTOS FARMACÊUTICOS S.A.	
DIOVAN320 MG COMPRIMIDOS REVESTIDOS POR PELÍCULA	SE/H/0406/006	5018940	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
DIOVAN320 MG COMPRIMIDOS REVESTIDOS POR PELÍCULA	SE/H/0406/006	5018957	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
DIOVAN320 MG COMPRIMIDOS REVESTIDOS POR PELÍCULA	SE/H/0406/006	5018965	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
DIOVAN320 MG COMPRIMIDOS REVESTIDOS POR PELÍCULA	SE/H/0406/006	5018973	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Diovane 160 mg filmomhulde tabletten	SE/H/0406/004	BE296423	NOVARTIS PHARMA N.V.	BE
Diovane 160 mg Filmtabletten	SE/H/0406/004	BE296423	NOVARTIS PHARMA N.V.	BE
Diovane 160 mg Filmtabletten	SE/H/0406/004	2001120026	NOVARTIS PHARMA N.V.	LU
Diovane 160 mg, comprimés pelliculés	SE/H/0406/004	BE296423	NOVARTIS PHARMA N.V.	BE
DIOVANE 160 MG, COMPRIMÉS PELLICULÉS	SE/H/0406/004	2001120026	NOVARTIS PHARMA N.V.	LU
Diovane 3 mg/ml drank	SE/H/0406/007	BE371007	NOVARTIS PHARMA N.V.	BE
Diovane 3 mg/ml Lösung zum Einnehmen	SE/H/0406/007	BE371007	NOVARTIS PHARMA N.V.	BE
Diovane 3 mg/ml Lösung	SE/H/0406/007	2010060100	NOVARTIS PHARMA N.V.	LU

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
zum Einnehmen				
Diovane 3 mg/ml solution buvable	SE/H/0406/007	BE371007	NOVARTIS PHARMA N.V.	BE
Diovane 3 mg/ml solution buvable	SE/H/0406/007	2010060100	NOVARTIS PHARMA N.V.	LU
Diovane 320 mg filmomhulde tabletten	SE/H/0406/006	BE297604	NOVARTIS PHARMA N.V.	BE
Diovane 320 mg Filmtabletten	SE/H/0406/006	BE297604	NOVARTIS PHARMA N.V.	BE
Diovane 320 mg Filmtabletten	SE/H/0406/006	2007040038	NOVARTIS PHARMA N.V.	LU
Diovane 320 mg, comprimés pelliculés	SE/H/0406/006	BE297604	NOVARTIS PHARMA N.V.	BE
DIOVANE 320 MG, COMPRIMÉS PELLICULÉS	SE/H/0406/006	2007040038	NOVARTIS PHARMA N.V.	LU
Diovane 40 mg filmomhulde tabletten	SE/H/0406/005	BE296405	NOVARTIS PHARMA N.V.	BE
Diovane 40 mg Filmtabletten	SE/H/0406/005	BE296405	NOVARTIS PHARMA N.V.	BE
Diovane 40 mg Filmtabletten	SE/H/0406/005	2005080009	NOVARTIS PHARMA N.V.	LU
Diovane 40 mg, comprimés pelliculés	SE/H/0406/005	BE296405	NOVARTIS PHARMA N.V.	BE
DIOVANE 40 MG, COMPRIMÉS PELLICULÉS	SE/H/0406/005	2005080009	NOVARTIS PHARMA N.V.	LU
Diovane 80 mg filmomhulde tabletten	SE/H/0406/003	BE296414	NOVARTIS PHARMA N.V.	BE
Diovane 80 mg Filmtabletten	SE/H/0406/003	BE296414	NOVARTIS PHARMA N.V.	BE
Diovane 80 mg Filmtabletten	SE/H/0406/003	2001120025	NOVARTIS PHARMA N.V.	LU
Diovane 80 mg, comprimés pelliculés	SE/H/0406/003	BE296414	NOVARTIS PHARMA N.V.	BE

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
DIOVANE 80 MG, COMPRIMÉS PELLICULÉS	SE/H/0406/003	2001120025	NOVARTIS PHARMA N.V.	LU
Kalpress 160 mg comprimidos recubiertos con película	SE/H/0407/002	64.452	NOVARTIS FARMACÉUTICA S.A.	ES
Kalpress 160 mg comprimidos recubiertos con película	SE/H/0407/002	64.452	NOVARTIS FARMACÉUTICA S.A.	ES
Kalpress 160 mg comprimidos recubiertos con película	SE/H/0407/002	64.452	NOVARTIS FARMACÉUTICA S.A.	ES
Kalpress 160 mg comprimidos recubiertos con película	SE/H/0407/002	64.452	NOVARTIS FARMACÉUTICA S.A.	ES
Kalpress 160 mg comprimidos recubiertos con película	SE/H/0407/002	64.452	NOVARTIS FARMACÉUTICA S.A.	ES
Kalpress 160 mg comprimidos recubiertos con película	SE/H/0407/002	64.452	NOVARTIS FARMACÉUTICA S.A.	ES
Kalpress 320 mg comprimidos recubiertos con película	SE/H/0407/004	69.517	NOVARTIS FARMACÉUTICA S.A.	ES
Kalpress 320 mg comprimidos recubiertos con película	SE/H/0407/004	69.517	NOVARTIS FARMACÉUTICA S.A.	ES
Kalpress 320 mg comprimidos recubiertos con película	SE/H/0407/004	69.517	NOVARTIS FARMACÉUTICA S.A.	ES
Kalpress 320 mg comprimidos recubiertos con película	SE/H/0407/004	69.517	NOVARTIS FARMACÉUTICA S.A.	ES
Kalpress 320 mg	SE/H/0407/004	69.517	NOVARTIS FARMACÉUTICA	ES

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
comprimidos recubiertos con película			S.A.	
Kalpress 320 mg comprimidos recubiertos con película	SE/H/0407/004	69.517	NOVARTIS FARMACÉUTICA S.A.	ES
Kalpress 80 mg comprimidos recubiertos con película	SE/H/0407/001	64.453	NOVARTIS FARMACÉUTICA S.A.	ES
Kalpress 80 mg comprimidos recubiertos con película	SE/H/0407/001	64.453	NOVARTIS FARMACÉUTICA S.A.	ES
Kalpress 80 mg comprimidos recubiertos con película	SE/H/0407/001	64.453	NOVARTIS FARMACÉUTICA S.A.	ES
Kalpress 80 mg comprimidos recubiertos con película	SE/H/0407/001	64.453	NOVARTIS FARMACÉUTICA S.A.	ES
Kalpress 80 mg comprimidos recubiertos con película	SE/H/0407/001	64.453	NOVARTIS FARMACÉUTICA S.A.	ES
Kalpress 80 mg comprimidos recubiertos con película	SE/H/0407/001	64.453	NOVARTIS FARMACÉUTICA S.A.	ES
Kalpress Cardio 40 mg comprimidos recubiertos con película	SE/H/0407/003	67.304	NOVARTIS FARMACÉUTICA S.A.	ES
Kalpress Cardio 40 mg comprimidos recubiertos con película	SE/H/0407/003	67.304	NOVARTIS FARMACÉUTICA S.A.	ES
Kalpress Cardio 40 mg comprimidos recubiertos con película	SE/H/0407/003	67.304	NOVARTIS FARMACÉUTICA S.A.	ES
Kalpress Cardio 40 mg	SE/H/0407/003	67.304	NOVARTIS FARMACÉUTICA	ES

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
comprimidos recubiertos con película			S.A.	
Kalpress Cardio 40 mg comprimidos recubiertos con película	SE/H/0407/003	67.304	NOVARTIS FARMACÉUTICA S.A.	ES
Kalpress Cardio 40 mg comprimidos recubiertos con película	SE/H/0407/003	67.304	NOVARTIS FARMACÉUTICA S.A.	ES
KALPRESS PLUS 160 MG/12,5 MG COMPRIMIDOS RECUBIERTOS CON PELÍCULA	SE/H/0566/002	65.898	NOVARTIS FARMACÉUTICA S.A.	ES
KALPRESS PLUS 160 MG/12,5 MG COMPRIMIDOS RECUBIERTOS CON PELÍCULA	SE/H/0566/002	65.898	NOVARTIS FARMACÉUTICA S.A.	ES
KALPRESS PLUS 160 MG/12,5 MG COMPRIMIDOS RECUBIERTOS CON PELÍCULA	SE/H/0566/002	65.898	NOVARTIS FARMACÉUTICA S.A.	ES
Kalpress Plus 320 mg/12,5 mg comprimidos recubiertos con película	SE/H/0566/004	70.058	NOVARTIS FARMACÉUTICA S.A.	ES
Kalpress Plus 320 mg/12,5 mg comprimidos recubiertos con película	SE/H/0566/004	70.058	NOVARTIS FARMACÉUTICA S.A.	ES
Kalpress Plus 320 mg/12,5 mg comprimidos recubiertos	SE/H/0566/004	70.058	NOVARTIS FARMACÉUTICA S.A.	ES

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 película				
KALPRESS PLUS 80MG/12,5 MG COMPRIMIDOS RECUBIERTOS CON PELÍCULA	SE/H/0566/001	62.562	NOVARTIS FARMACÉUTICA S.A.	ES
KALPRESS PLUS 80MG/12,5 MG COMPRIMIDOS RECUBIERTOS CON PELÍCULA	SE/H/0566/001	62.562	NOVARTIS FARMACÉUTICA S.A.	ES
KALPRESS PLUS 80MG/12,5 MG COMPRIMIDOS RECUBIERTOS CON PELÍCULA	SE/H/0566/001	62.562	NOVARTIS FARMACÉUTICA S.A.	ES
KALPRESS PLUS FORTE 160 MG/25 MG COMPRIMIDOS RECUBIERTOS CON PELÍCULA	SE/H/0566/003	66.744	NOVARTIS FARMACÉUTICA S.A.	ES
KALPRESS PLUS FORTE 160 MG/25 MG COMPRIMIDOS RECUBIERTOS CON PELÍCULA	SE/H/0566/003	66.744	NOVARTIS FARMACÉUTICA S.A.	ES
KALPRESS PLUS FORTE 160 MG/25 MG COMPRIMIDOS RECUBIERTOS CON PELÍCULA	SE/H/0566/003	66.744	NOVARTIS FARMACÉUTICA S.A.	ES
Kalpress Plus Forte 320 mg/25 mg comprimidos recubiertos con película	SE/H/0566/005	70.060	NOVARTIS FARMACÉUTICA S.A.	ES

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
Kalpress Plus Forte 320 mg/25 mg comprimidos recubiertos con película	SE/H/0566/005	70.060	NOVARTIS FARMACÉUTICA S.A.	ES
Kalpress Plus Forte 320 mg/25 mg comprimidos recubiertos con película	SE/H/0566/005	70.060	NOVARTIS FARMACÉUTICA S.A.	ES
Miten 160 mg comprimidos recubiertos con película	SE/H/0408/002	64.963	NOVARTIS FARMACÉUTICA S.A.	ES
Miten 160 mg comprimidos recubiertos con película	SE/H/0408/002	64.963	NOVARTIS FARMACÉUTICA S.A.	ES
Miten 160 mg comprimidos recubiertos con película	SE/H/0408/002	64.963	NOVARTIS FARMACÉUTICA S.A.	ES
Miten 160 mg comprimidos recubiertos con película	SE/H/0408/002	64.963	NOVARTIS FARMACÉUTICA S.A.	ES
Miten 160 mg comprimidos recubiertos con película	SE/H/0408/002	64.963	NOVARTIS FARMACÉUTICA S.A.	ES
Miten 320 mg comprimidos recubiertos con película	SE/H/0408/004	69.102	NOVARTIS FARMACÉUTICA S.A.	ES
Miten 320 mg comprimidos recubiertos con película	SE/H/0408/004	69.102	NOVARTIS FARMACÉUTICA S.A.	ES
Miten 320 mg comprimidos recubiertos con película	SE/H/0408/004	69.102	NOVARTIS FARMACÉUTICA S.A.	ES
Miten 320 mg comprimidos recubiertos con película	SE/H/0408/004	69.102	NOVARTIS FARMACÉUTICA S.A.	ES

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
Miten 320 mg comprimidos recubiertos con película	SE/H/0408/004	69.102	NOVARTIS FARMACÉUTICA S.A.	ES
Miten 80 mg comprimidos recubiertos con película	SE/H/0408/001	64.961	NOVARTIS FARMACÉUTICA S.A.	ES
Miten 80 mg comprimidos recubiertos con película	SE/H/0408/001	64.961	NOVARTIS FARMACÉUTICA S.A.	ES
Miten 80 mg comprimidos recubiertos con película	SE/H/0408/001	64.961	NOVARTIS FARMACÉUTICA S.A.	ES
Miten 80 mg comprimidos recubiertos con película	SE/H/0408/001	64.961	NOVARTIS FARMACÉUTICA S.A.	ES
Miten 80 mg comprimidos recubiertos con película	SE/H/0408/001	64.961	NOVARTIS FARMACÉUTICA S.A.	ES
Miten Cardio 40 mg comprimidos recubiertos con película	SE/H/0408/003	67.301	NOVARTIS FARMACÉUTICA S.A.	ES
Miten Cardio 40 mg comprimidos recubiertos con película	SE/H/0408/003	67.301	NOVARTIS FARMACÉUTICA S.A.	ES
Miten Cardio 40 mg comprimidos recubiertos con película	SE/H/0408/003	67.301	NOVARTIS FARMACÉUTICA S.A.	ES
Miten Cardio 40 mg comprimidos recubiertos con película	SE/H/0408/003	67.301	NOVARTIS FARMACÉUTICA S.A.	ES
Miten Cardio 40 mg comprimidos recubiertos con película	SE/H/0408/003	67.301	NOVARTIS FARMACÉUTICA S.A.	ES

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
Miten Plus 160 mg/12,5 mg comprimidos recubiertos con película	SE/H/0567/001	65.897	NOVARTIS FARMACÉUTICA S.A.	ES
Miten Plus 160 mg/12,5 mg comprimidos recubiertos con película	SE/H/0567/001	65.897	NOVARTIS FARMACÉUTICA S.A.	ES
Miten Plus 80mg/12,5 mg comprimidos recubiertos con película	SE/H/0567/003	64.233	NOVARTIS FARMACÉUTICA S.A.	ES
Miten Plus 80mg/12,5 mg comprimidos recubiertos con película	SE/H/0567/003	64.233	NOVARTIS FARMACÉUTICA S.A.	ES
Miten Plus Forte 160 mg/25 mg comprimidos recubiertos con película	SE/H/0567/002	66.760	NOVARTIS FARMACÉUTICA S.A.	ES
Miten Plus Forte 160 mg/25 mg comprimidos recubiertos con película	SE/H/0567/002	66.760	NOVARTIS FARMACÉUTICA S.A.	ES
NISIS 160 mg, comprimé pelliculé	not available	34009 356 946 2 1	IPSEN PHARMA	FR
NISIS 160 mg, comprimé pelliculé	not available	34009 371 494 1 9	IPSEN PHARMA	FR
NISIS 160 mg, comprimé pelliculé	not available	34009 356 947 9 9	IPSEN PHARMA	FR
NISIS 160 mg, comprimé pelliculé	not available	34009 371 495 8 7	IPSEN PHARMA	FR
NISIS 160 mg, comprimé pelliculé	not available	34009 371 496 4 8	IPSEN PHARMA	FR
NISIS 160 mg, comprimé pelliculé	not available	34009 356 948 5 0	IPSEN PHARMA	FR
NISIS 160 mg, comprimé pelliculé	not available	34009 356 958 0 2	IPSEN PHARMA	FR
NISIS 160 mg,	not available	34009 371 497 0 9	IPSEN PHARMA	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
comprimé pelliculé				
NISIS 160 mg, comprimé pelliculé	not available	34009 356 959 7 0	IPSEN PHARMA	FR
NISIS 160 mg, comprimé pelliculé	not available	34009 373 892 4 2	IPSEN PHARMA	FR
NISIS 160 mg, comprimé pelliculé	not available	34009 371 498 7 7	IPSEN PHARMA	FR
NISIS 160 mg, comprimé pelliculé	not available	34009 379 068 1 4	IPSEN PHARMA	FR
NISIS 160 mg, comprimé pelliculé	not available	34009 379 069 8 2	IPSEN PHARMA	FR
NISIS 160 mg, comprimé pelliculé	not available	34009 379 070 6 4	IPSEN PHARMA	FR
NISIS 160 mg, comprimé pelliculé	not available	34009 379 071 2 5	IPSEN PHARMA	FR
NISIS 160 mg, comprimé pelliculé	not available	34009 379 072 9 3	IPSEN PHARMA	FR
NISIS 160 mg, comprimé pelliculé	not available	34009 379 073 5 4	IPSEN PHARMA	FR
NISIS 160 mg, comprimé pelliculé	not available	34009 356 945 6 0	IPSEN PHARMA	FR
NISIS 40 mg, comprimé pelliculé sécable	not available	356 932-1	IPSEN PHARMA	FR
NISIS 40 mg, comprimé pelliculé sécable	not available	372 292-3	IPSEN PHARMA	FR
NISIS 40 mg, comprimé pelliculé sécable	not available	356 933-8	IPSEN PHARMA	FR
NISIS 40 mg, comprimé pelliculé sécable	not available	371 509-9	IPSEN PHARMA	FR
NISIS 40 mg, comprimé pelliculé sécable	not available	372 295-2	IPSEN PHARMA	FR
NISIS 40 mg, comprimé pelliculé sécable	not available	356 934-4	IPSEN PHARMA	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
NISIS 40 mg, comprimé pelliculé sécable	not available	356 935-0	IPSEN PHARMA	FR
NISIS 40 mg, comprimé pelliculé sécable	not available	372 294-6	IPSEN PHARMA	FR
NISIS 40 mg, comprimé pelliculé sécable	not available	356 936-7	IPSEN PHARMA	FR
NISIS 40 mg, comprimé pelliculé sécable	not available	373 890-1	IPSEN PHARMA	FR
NISIS 40 mg, comprimé pelliculé sécable	not available	372 296-9	IPSEN PHARMA	FR
NISIS 40 mg, comprimé pelliculé sécable	not available	379 054-0	IPSEN PHARMA	FR
NISIS 40 mg, comprimé pelliculé sécable	not available	379 055-7	IPSEN PHARMA	FR
NISIS 40 mg, comprimé pelliculé sécable	not available	379 056-3	IPSEN PHARMA	FR
NISIS 40 mg, comprimé pelliculé sécable	not available	379 058-6	IPSEN PHARMA	FR
NISIS 40 mg, comprimé pelliculé sécable	not available	379 059-2	IPSEN PHARMA	FR
NISIS 40 mg, comprimé pelliculé sécable	not available	379 060-0	IPSEN PHARMA	FR
NISIS 40 mg, comprimé pelliculé sécable	not available	356 931-5	IPSEN PHARMA	FR
NISIS 80 mg, comprimé pelliculé	not available	34009 356 939 6 9	IPSEN PHARMA	FR
NISIS 80 mg, comprimé pelliculé	not available	34009 372 290 0 5	IPSEN PHARMA	FR
NISIS 80 mg, comprimé pelliculé	not available	34009 356 940 4 1	IPSEN PHARMA	FR
NISIS 80 mg, comprimé pelliculé	not available	34009 371 510 7 8	IPSEN PHARMA	FR
NISIS 80 mg, comprimé	not available	34009 371 511 3 9	IPSEN PHARMA	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
pelliculé				
NISIS 80 mg, comprimé pelliculé	not available	34009 356 941 0 2	IPSEN PHARMA	FR
NISIS 80 mg, comprimé pelliculé	not available	34009 356 942 7 0	IPSEN PHARMA	FR
NISIS 80 mg, comprimé pelliculé	not available	34009 372 291 7 3	IPSEN PHARMA	FR
NISIS 80 mg, comprimé pelliculé	not available	34009 356 943 3 1	IPSEN PHARMA	FR
NISIS 80 mg, comprimé pelliculé	not available	34009 373 891 8 1	IPSEN PHARMA	FR
NISIS 80 mg, comprimé pelliculé	not available	34009 371 513 6 8	IPSEN PHARMA	FR
NISIS 80 mg, comprimé pelliculé	not available	34009 379 061 7 3	IPSEN PHARMA	FR
NISIS 80 mg, comprimé pelliculé	not available	34009 379 062 3 4	IPSEN PHARMA	FR
NISIS 80 mg, comprimé pelliculé	not available	34009 379 064 6 3	IPSEN PHARMA	FR
NISIS 80 mg, comprimé pelliculé	not available	34009 379 065 2 4	IPSEN PHARMA	FR
NISIS 80 mg, comprimé pelliculé	not available	34009 379 067 5 3	IPSEN PHARMA	FR
NISIS 80 mg, comprimé pelliculé	not available	34009 379 066 9 2	IPSEN PHARMA	FR
NISIS 80 mg, comprimé pelliculé	not available	34009 356 937 3 0	IPSEN PHARMA	FR
NISISCO 160 mg/12,5 mg, comprimé pelliculé	not available	34009 371 515 9 7	IPSEN PHARMA	FR
NISISCO 160 mg/12,5 mg, comprimé pelliculé	not available	34009 564 979 7 3	IPSEN PHARMA	FR
NISISCO 160 mg/12,5 mg, comprimé pelliculé	not available	34009 564 980 5 5	IPSEN PHARMA	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
NISISCO 160 mg/12,5 mg, comprimé pelliculé	not available	34009 371 514 2 9	IPSEN PHARMA	FR
NISISCO 160 mg/12,5 mg, comprimé pelliculé	not available	34009 371 516 5 8	IPSEN PHARMA	FR
NISISCO 160 mg/12,5 mg, comprimé pelliculé	not available	34009 564 981 1 6	IPSEN PHARMA	FR
NISISCO 160 mg/12,5 mg, comprimé pelliculé	not available	34009 379 039 1 2	IPSEN PHARMA	FR
NISISCO 160 mg/12,5 mg, comprimé pelliculé	not available	34009 379 041 6 2	IPSEN PHARMA	FR
NISISCO 160 mg/12,5 mg, comprimé pelliculé	not available	34009 379 042 2 3	IPSEN PHARMA	FR
NISISCO 160 mg/12,5 mg, comprimé pelliculé	not available	34009 379 043 9 1	IPSEN PHARMA	FR
NISISCO 160 mg/12,5 mg, comprimé pelliculé	not available	34009 379 044 5 2	IPSEN PHARMA	FR
NISISCO 160 mg/12,5 mg, comprimé pelliculé	not available	34009 379 045 1 3	IPSEN PHARMA	FR
NISISCO 160 mg/12,5 mg, comprimé pelliculé	not available	34009 572 099 2 6	IPSEN PHARMA	FR
NISISCO 160 mg/12,5 mg, comprimé pelliculé	not available	34009 363 108 9 6	IPSEN PHARMA	FR
NISISCO 160 mg/25 mg, comprimé pelliculé	not available	34009 371 492 9 7	IPSEN PHARMA	FR
NISISCO 160 mg/25 mg, comprimé pelliculé	not available	34009 563 687 2 3	IPSEN PHARMA	FR
NISISCO 160 mg/25 mg, comprimé pelliculé	not available	34009 563 688 9 1	IPSEN PHARMA	FR
NISISCO 160 mg/25 mg, comprimé pelliculé	not available	34009 373 893 0 3	IPSEN PHARMA	FR
NISISCO 160 mg/25 mg, comprimé pelliculé	not available	34009 371 493 5 8	IPSEN PHARMA	FR
NISISCO 160 mg/25 mg, comprimé pelliculé	not available	34009 563 689 5 2	IPSEN PHARMA	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
comprimé pelliculé				
NISISCO 160 mg/25 mg, comprimé pelliculé	not available	34009 379 047 4 2	IPSEN PHARMA	FR
NISISCO 160 mg/25 mg, comprimé pelliculé	not available	34009 379 048 0 3	IPSEN PHARMA	FR
NISISCO 160 mg/25 mg, comprimé pelliculé	not available	34009 379 049 7 1	IPSEN PHARMA	FR
NISISCO 160 mg/25 mg, comprimé pelliculé	not available	34009 379 050 5 3	IPSEN PHARMA	FR
NISISCO 160 mg/25 mg, comprimé pelliculé	not available	34009 379 051 1 4	IPSEN PHARMA	FR
NISISCO 160 mg/25 mg, comprimé pelliculé	not available	34009 379 052 8 2	IPSEN PHARMA	FR
NISISCO 160 mg/25 mg, comprimé pelliculé	not available	34009 572 100 0 7	IPSEN PHARMA	FR
NISISCO 160 mg/25 mg, comprimé pelliculé	not available	34009 359 237 2 1	IPSEN PHARMA	FR
NISISCO 80 mg/12,5 mg, comprimé pelliculé	not available	34009 371 518 8 7	IPSEN PHARMA	FR
NISISCO 80 mg/12,5 mg, comprimé pelliculé	not available	34009 345 415 0 6	IPSEN PHARMA	FR
NISISCO 80 mg/12,5 mg, comprimé pelliculé	not available	34009 345 416 7 4	IPSEN PHARMA	FR
NISISCO 80 mg/12,5 mg, comprimé pelliculé	not available	34009 371 517 1 9	IPSEN PHARMA	FR
NISISCO 80 mg/12,5 mg, comprimé pelliculé	not available	34009 371 519 4 8	IPSEN PHARMA	FR
NISISCO 80 mg/12,5 mg, comprimé pelliculé	not available	34009 345 417 3 5	IPSEN PHARMA	FR
NISISCO 80 mg/12,5 mg, comprimé pelliculé	not available	34009 379 031 0 3	IPSEN PHARMA	FR
NISISCO 80 mg/12,5 mg, comprimé pelliculé	not available	34009 379 032 7 1	IPSEN PHARMA	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
NISISCO 80 mg/12,5 mg, comprimé pelliculé	not available	34009 379 033 3 2	IPSEN PHARMA	FR
NISISCO 80 mg/12,5 mg, comprimé pelliculé	not available	34009 379 035 6 1	IPSEN PHARMA	FR
NISISCO 80 mg/12,5 mg, comprimé pelliculé	not available	34009 379 036 2 2	IPSEN PHARMA	FR
NISISCO 80 mg/12,5 mg, comprimé pelliculé	not available	34009 379 037 9 0	IPSEN PHARMA	FR
NISISCO 80 mg/12,5 mg, comprimé pelliculé	not available	34009 572 101 7 5	IPSEN PHARMA	FR
NISISCO 80 mg/12,5 mg, comprimé pelliculé	not available	34009 345 414 4 5	IPSEN PHARMA	FR
Provas 160 mg Filmtabletten	SE/H/0408/002	50720.01.00	NOVARTIS PHARMA GMBH	DE
Provas 160 mg Filmtabletten	SE/H/0408/002	50720.01.00	NOVARTIS PHARMA GMBH	DE
Provas 160 mg Filmtabletten	SE/H/0408/002	50720.01.00	NOVARTIS PHARMA GMBH	DE
Provas 160 mg Filmtabletten	SE/H/0408/002	50720.01.00	NOVARTIS PHARMA GMBH	DE
Provas 160 mg Filmtabletten	SE/H/0408/002	50720.01.00	NOVARTIS PHARMA GMBH	DE
Provas 320 mg Filmtabletten	SE/H/0408/004	67300.00.00	NOVARTIS PHARMA GMBH	DE
Provas 320 mg Filmtabletten	SE/H/0408/004	67300.00.00	NOVARTIS PHARMA GMBH	DE
Provas 320 mg Filmtabletten	SE/H/0408/004	67300.00.00	NOVARTIS PHARMA GMBH	DE
Provas 320 mg Filmtabletten	SE/H/0408/004	67300.00.00	NOVARTIS PHARMA GMBH	DE
Provas 40 mg	SE/H/0408/003	50720.02.00	NOVARTIS 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
Filmdabletten				
Provas 40 mg Filmdabletten	SE/H/0408/003	50720.02.00	NOVARTIS PHARMA GMBH	DE
Provas 40 mg Filmdabletten	SE/H/0408/003	50720.02.00	NOVARTIS PHARMA GMBH	DE
Provas 40 mg Filmdabletten	SE/H/0408/003	50720.02.00	NOVARTIS PHARMA GMBH	DE
Provas 40 mg Filmdabletten	SE/H/0408/003	50720.02.00	NOVARTIS PHARMA GMBH	DE
Provas 80 mg Filmdabletten	SE/H/0408/001	50720.00.00	NOVARTIS PHARMA GMBH	DE
Provas 80 mg Filmdabletten	SE/H/0408/001	50720.00.00	NOVARTIS PHARMA GMBH	DE
Provas 80 mg Filmdabletten	SE/H/0408/001	50720.00.00	NOVARTIS PHARMA GMBH	DE
Provas 80 mg Filmdabletten	SE/H/0408/001	50720.00.00	NOVARTIS PHARMA GMBH	DE
Provas 80 mg Filmdabletten	SE/H/0408/001	50720.00.00	NOVARTIS PHARMA GMBH	DE
Provas® 160 comp 160 mg/12,5 mg Filmdabletten	IT/H/0244/002	52890.00.00	NOVARTIS PHARMA GMBH	DE
Provas® 160 maxx 160 mg/25 mg Filmdabletten	IT/H/0244/003	52891.00.00	NOVARTIS PHARMA GMBH	DE
Provas® 320 comp 320 mg/12,5 mg Filmdabletten	SE/H/0567/004	69624.00.00	NOVARTIS PHARMA GMBH	DE
Provas® 320 comp 320 mg/12,5 mg Filmdabletten	SE/H/0567/004	69624.00.00	NOVARTIS PHARMA GMBH	DE
Provas® 320 maxx 320 mg/25 mg Filmdabletten	SE/H/0567/005	69625.00.00	NOVARTIS PHARMA GMBH	DE
Provas® 320 maxx 320	SE/H/0567/005	69625.00.00	NOVARTIS 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
mg/25 mg Filmdabletten				
Provas® 80 comp 80 mg/12,5 mg Filmdabletten	IT/H/0244/001	44775.00.00	NOVARTIS PHARMA GMBH	DE
RIXIL	SE/H/0407/005	034776361	SANDOZ S.P.A.	IT
RIXIL	SE/H/0407/005	034776361	SANDOZ S.P.A.	IT
RIXIL	SE/H/0407/005	034776361	SANDOZ S.P.A.	IT
RIXIL	SE/H/0407/005	034776361	SANDOZ S.P.A.	IT
RIXIL	SE/H/0407/005	034776361	SANDOZ S.P.A.	IT
RIXIL	SE/H/0407/005	034776361	SANDOZ S.P.A.	IT
RIXIL	SE/H/0407/005	034776361	SANDOZ S.P.A.	IT
TAREG 160 mg compresse rivestite con film	SE/H/0406/004	033178144	NOVARTIS EUROPHARM LIMITED	IT
TAREG 160 mg compresse rivestite con film	SE/H/0406/004	033178157	NOVARTIS EUROPHARM LIMITED	IT
TAREG 160 mg compresse rivestite con film	SE/H/0406/004	033178169	NOVARTIS EUROPHARM LIMITED	IT
TAREG 160 mg compresse rivestite con film	SE/H/0406/004	033178171	NOVARTIS EUROPHARM LIMITED	IT
TAREG 160 mg compresse rivestite con film	SE/H/0406/004	033178183	NOVARTIS EUROPHARM LIMITED	IT
TAREG 160 mg compresse rivestite con film	SE/H/0406/004	033178753	NOVARTIS EUROPHARM LIMITED	IT
TAREG 160 mg compresse rivestite con film	SE/H/0406/004	033178765	NOVARTIS EUROPHARM LIMITED	IT
TAREG 160 mg	SE/H/0406/004	033178777	NOVARTIS EUROPHARM	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			LIMITED	
TAREG 160 mg compresse rivestite con film	SE/H/0406/004	033178789	NOVARTIS EUROPHARM LIMITED	IT
TAREG 160 mg compresse rivestite con film	SE/H/0406/004	033178791	NOVARTIS EUROPHARM LIMITED	IT
TAREG 160 mg compresse rivestite con film	SE/H/0406/004	033178803	NOVARTIS EUROPHARM LIMITED	IT
TAREG 160 mg compresse rivestite con film	SE/H/0406/004	033178815	NOVARTIS EUROPHARM LIMITED	IT
TAREG 160 mg compresse rivestite con film	SE/H/0406/004	033178827	NOVARTIS EUROPHARM LIMITED	IT
TAREG 160 mg compresse rivestite con film	SE/H/0406/004	033178839	NOVARTIS EUROPHARM LIMITED	IT
TAREG 160 mg compresse rivestite con film	SE/H/0406/004	033178841	NOVARTIS EUROPHARM LIMITED	IT
TAREG 160 mg compresse rivestite con film	SE/H/0406/004	033178930	NOVARTIS EUROPHARM LIMITED	IT
TAREG 160 mg compresse rivestite con film	SE/H/0406/004	033178942	NOVARTIS EUROPHARM LIMITED	IT
TAREG 160 mg compresse rivestite con film	SE/H/0406/004	033178955	NOVARTIS EUROPHARM LIMITED	IT
TAREG 160 mg	SE/H/0406/004	033178967	NOVARTIS EUROPHARM	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			LIMITED	
TAREG 160 mg compresse rivestite con film	SE/H/0406/004	033178979	NOVARTIS EUROPHARM LIMITED	IT
Tareg 160 mg comprimidos revestidos por película	SE/H/0407/002	3845187	LABORATORIO NORMAL-PRODUTOS FARMACEUTICOS, LDA	PT
Tareg 160 mg comprimidos revestidos por película	SE/H/0407/002	3845286	LABORATORIO NORMAL-PRODUTOS FARMACEUTICOS, LDA	PT
Tareg 160 mg comprimidos revestidos por película	SE/H/0407/002	3845385	LABORATORIO NORMAL-PRODUTOS FARMACEUTICOS, LDA	PT
Tareg 160 mg comprimidos revestidos por película	SE/H/0407/002	5317482	LABORATORIO NORMAL-PRODUTOS FARMACEUTICOS, LDA	PT
Tareg 160 mg comprimidos revestidos por película	SE/H/0407/002	5920889	LABORATORIO NORMAL-PRODUTOS FARMACEUTICOS, LDA	PT
Tareg 160 mg comprimidos revestidos por película	SE/H/0407/002	3845187	LABORATORIO NORMAL-PRODUTOS FARMACEUTICOS, LDA	PT
Tareg 160 mg comprimidos revestidos por película	SE/H/0407/002	3845286	LABORATORIO NORMAL-PRODUTOS FARMACEUTICOS, LDA	PT
Tareg 160 mg comprimidos revestidos por película	SE/H/0407/002	3845385	LABORATORIO NORMAL-PRODUTOS FARMACEUTICOS, LDA	PT
Tareg 160 mg comprimidos revestidos por película	SE/H/0407/002	5317482	LABORATORIO NORMAL-PRODUTOS FARMACEUTICOS, LDA	PT
Tareg 160 mg	SE/H/0407/002	5920889	LABORATORIO NORMAL-	PT

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
comprimidos revestidos por película			PRODUTOS FARMACEUTICOS, LDA	
Tareg 160 mg comprimidos revestidos por película	SE/H/0407/002	3845187	LABORATORIO NORMAL-PRODUTOS FARMACEUTICOS, LDA	PT
Tareg 160 mg comprimidos revestidos por película	SE/H/0407/002	3845286	LABORATORIO NORMAL-PRODUTOS FARMACEUTICOS, LDA	PT
Tareg 160 mg comprimidos revestidos por película	SE/H/0407/002	3845385	LABORATORIO NORMAL-PRODUTOS FARMACEUTICOS, LDA	PT
Tareg 160 mg comprimidos revestidos por película	SE/H/0407/002	5317482	LABORATORIO NORMAL-PRODUTOS FARMACEUTICOS, LDA	PT
Tareg 160 mg comprimidos revestidos por película	SE/H/0407/002	5920889	LABORATORIO NORMAL-PRODUTOS FARMACEUTICOS, LDA	PT
Tareg 160 mg comprimidos revestidos por película	SE/H/0407/002	3845187	LABORATORIO NORMAL-PRODUTOS FARMACEUTICOS, LDA	PT
Tareg 160 mg comprimidos revestidos por película	SE/H/0407/002	3845286	LABORATORIO NORMAL-PRODUTOS FARMACEUTICOS, LDA	PT
Tareg 160 mg comprimidos revestidos por película	SE/H/0407/002	3845385	LABORATORIO NORMAL-PRODUTOS FARMACEUTICOS, LDA	PT
Tareg 160 mg comprimidos revestidos por película	SE/H/0407/002	5317482	LABORATORIO NORMAL-PRODUTOS FARMACEUTICOS, LDA	PT
Tareg 160 mg comprimidos revestidos por película	SE/H/0407/002	5920889	LABORATORIO NORMAL-PRODUTOS FARMACEUTICOS, LDA	PT
Tareg 160 mg	SE/H/0407/002	3845187	LABORATORIO NORMAL-	PT

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
comprimidos revestidos por película			PRODUTOS FARMACEUTICOS, LDA	
Tareg 160 mg comprimidos revestidos por película	SE/H/0407/002	3845286	LABORATORIO NORMAL-PRODUTOS FARMACEUTICOS, LDA	PT
Tareg 160 mg comprimidos revestidos por película	SE/H/0407/002	3845385	LABORATORIO NORMAL-PRODUTOS FARMACEUTICOS, LDA	PT
Tareg 160 mg comprimidos revestidos por película	SE/H/0407/002	5317482	LABORATORIO NORMAL-PRODUTOS FARMACEUTICOS, LDA	PT
Tareg 160 mg comprimidos revestidos por película	SE/H/0407/002	5920889	LABORATORIO NORMAL-PRODUTOS FARMACEUTICOS, LDA	PT
Tareg 160 mg comprimidos revestidos por película	SE/H/0407/002	3845187	LABORATORIO NORMAL-PRODUTOS FARMACEUTICOS, LDA	PT
Tareg 160 mg comprimidos revestidos por película	SE/H/0407/002	3845286	LABORATORIO NORMAL-PRODUTOS FARMACEUTICOS, LDA	PT
Tareg 160 mg comprimidos revestidos por película	SE/H/0407/002	3845385	LABORATORIO NORMAL-PRODUTOS FARMACEUTICOS, LDA	PT
Tareg 160 mg comprimidos revestidos por película	SE/H/0407/002	5317482	LABORATORIO NORMAL-PRODUTOS FARMACEUTICOS, LDA	PT
Tareg 160 mg comprimidos revestidos por película	SE/H/0407/002	5920889	LABORATORIO NORMAL-PRODUTOS FARMACEUTICOS, LDA	PT
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

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
TAREG 160 mg, comprimé pelliculé	SE/H/0406/004	34009 365 907 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
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 3 mg/ml soluzione orale	SE/H/0406/007	033178423	NOVARTIS EUROPHARM LIMITED	IT
Tareg 3 mg/ml, solution buvable	SE/H/0406/007	3400949147489	NOVARTIS PHARMA S.A.S.	FR
Tareg 320 mg compresse rivestite con film	SE/H/0406/006	033178334	NOVARTIS EUROPHARM LIMITED	IT
Tareg 320 mg compresse rivestite con film	SE/H/0406/006	033178346	NOVARTIS EUROPHARM LIMITED	IT
Tareg 320 mg compresse rivestite con film	SE/H/0406/006	033178359	NOVARTIS EUROPHARM LIMITED	IT
Tareg 320 mg compresse rivestite con film	SE/H/0406/006	033178361	NOVARTIS EUROPHARM LIMITED	IT
Tareg 320 mg compresse rivestite con film	SE/H/0406/006	033178373	NOVARTIS EUROPHARM LIMITED	IT
Tareg 320 mg compresse rivestite con film	SE/H/0406/006	033178385	NOVARTIS EUROPHARM LIMITED	IT
Tareg 320 mg compresse	SE/H/0406/006	033178397	NOVARTIS EUROPHARM	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			LIMITED	
Tareg 320 mg compresse rivestite con film	SE/H/0406/006	033178409	NOVARTIS EUROPHARM LIMITED	IT
Tareg 320 mg compresse rivestite con film	SE/H/0406/006	033178411	NOVARTIS EUROPHARM LIMITED	IT
Tareg 320 mg compresse rivestite con film	SE/H/0406/006	033178690	NOVARTIS EUROPHARM LIMITED	IT
Tareg 320 mg compresse rivestite con film	SE/H/0406/006	033178702	NOVARTIS EUROPHARM LIMITED	IT
Tareg 320 mg compresse rivestite con film	SE/H/0406/006	033178714	NOVARTIS EUROPHARM LIMITED	IT
Tareg 320 mg compresse rivestite con film	SE/H/0406/006	033178726	NOVARTIS EUROPHARM LIMITED	IT
Tareg 320 mg compresse rivestite con film	SE/H/0406/006	033178738	NOVARTIS EUROPHARM LIMITED	IT
Tareg 320 mg compresse rivestite con film	SE/H/0406/006	033178740	NOVARTIS EUROPHARM LIMITED	IT
TAREG 40 mg compresse rivestite con film	SE/H/0406/005	033178195	NOVARTIS EUROPHARM LIMITED	IT
TAREG 40 mg compresse rivestite con film	SE/H/0406/005	033178219	NOVARTIS EUROPHARM LIMITED	IT
TAREG 40 mg compresse rivestite con film	SE/H/0406/005	033178221	NOVARTIS EUROPHARM LIMITED	IT
TAREG 40 mg compresse rivestite con film	SE/H/0406/005	033178233	NOVARTIS EUROPHARM LIMITED	IT
TAREG 40 mg compresse rivestite con film	SE/H/0406/005	033178245	NOVARTIS EUROPHARM LIMITED	IT
TAREG 40 mg compresse rivestite con film	SE/H/0406/005	033178258	NOVARTIS EUROPHARM LIMITED	IT
TAREG 40 mg compresse rivestite con film	SE/H/0406/005	033178260	NOVARTIS EUROPHARM LIMITED	IT
TAREG 40 mg compresse rivestite con film	SE/H/0406/005	033178272	NOVARTIS EUROPHARM LIMITED	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
TAREG 40 mg compresse rivestite con film	SE/H/0406/005	033178207	NOVARTIS EUROPHARM LIMITED	IT
TAREG 40 mg compresse rivestite con film	SE/H/0406/005	033178284	NOVARTIS EUROPHARM LIMITED	IT
TAREG 40 mg compresse rivestite con film	SE/H/0406/005	033178296	NOVARTIS EUROPHARM LIMITED	IT
TAREG 40 mg compresse rivestite con film	SE/H/0406/005	033178308	NOVARTIS EUROPHARM LIMITED	IT
TAREG 40 mg compresse rivestite con film	SE/H/0406/005	033178310	NOVARTIS EUROPHARM LIMITED	IT
TAREG 40 mg compresse rivestite con film	SE/H/0406/005	033178322	NOVARTIS EUROPHARM LIMITED	IT
TAREG 40 mg compresse rivestite con film	SE/H/0406/005	033178435	NOVARTIS EUROPHARM LIMITED	IT
TAREG 40 mg compresse rivestite con film	SE/H/0406/005	033178447	NOVARTIS EUROPHARM LIMITED	IT
TAREG 40 mg compresse rivestite con film	SE/H/0406/005	033178450	NOVARTIS EUROPHARM LIMITED	IT
TAREG 40 mg compresse rivestite con film	SE/H/0406/005	033178462	NOVARTIS EUROPHARM LIMITED	IT
TAREG 40 mg compresse rivestite con film	SE/H/0406/005	033178474	NOVARTIS EUROPHARM LIMITED	IT
TAREG 40 mg compresse rivestite con film	SE/H/0406/005	033178486	NOVARTIS EUROPHARM LIMITED	IT
TAREG 40 mg compresse rivestite con film	SE/H/0406/005	033178498	NOVARTIS EUROPHARM LIMITED	IT
TAREG 40 mg compresse rivestite con film	SE/H/0406/005	033178500	NOVARTIS EUROPHARM LIMITED	IT
TAREG 40 mg compresse rivestite con film	SE/H/0406/005	033178512	NOVARTIS EUROPHARM LIMITED	IT
TAREG 40 mg compresse rivestite con film	SE/H/0406/005	033178536	NOVARTIS EUROPHARM LIMITED	IT
TAREG 40 mg compresse	SE/H/0406/005	033178548	NOVARTIS EUROPHARM	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			LIMITED	
TAREG 40 mg compresse rivestite con film	SE/H/0406/005	033178551	NOVARTIS EUROPHARM LIMITED	IT
TAREG 40 mg compresse rivestite con film	SE/H/0406/005	033178563	NOVARTIS EUROPHARM LIMITED	IT
TAREG 40 mg compresse rivestite con film	SE/H/0406/005	033178575	NOVARTIS EUROPHARM LIMITED	IT
TAREG 40 mg compresse rivestite con film	SE/H/0406/005	033178587	NOVARTIS EUROPHARM LIMITED	IT
TAREG 40 mg compresse rivestite con film	SE/H/0406/005	033178599	NOVARTIS EUROPHARM LIMITED	IT
TAREG 40 mg compresse rivestite con film	SE/H/0406/005	033178601	NOVARTIS EUROPHARM LIMITED	IT
TAREG 40 mg compresse rivestite con film	SE/H/0406/005	033178613	NOVARTIS EUROPHARM LIMITED	IT
Tareg 40 mg comprimidos revestidos por película	SE/H/0407/003	5002472	LABORATORIO NORMAL-PRODUTOS FARMACEUTICOS, LDA	PT
Tareg 40 mg comprimidos revestidos por película	SE/H/0407/003	5002506	LABORATORIO NORMAL-PRODUTOS FARMACEUTICOS, LDA	PT
Tareg 40 mg comprimidos revestidos por película	SE/H/0407/003	5002514	LABORATORIO NORMAL-PRODUTOS FARMACEUTICOS, LDA	PT
Tareg 40 mg comprimidos revestidos por película	SE/H/0407/003	5002522	LABORATORIO NORMAL-PRODUTOS FARMACEUTICOS, LDA	PT
Tareg 40 mg comprimidos revestidos por película	SE/H/0407/003	5002530	LABORATORIO NORMAL-PRODUTOS FARMACEUTICOS, LDA	PT
Tareg 40 mg comprimidos revestidos por película	SE/H/0407/003	5002472	LABORATORIO NORMAL-PRODUTOS FARMACEUTICOS, LDA	PT

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
Tareg 40 mg comprimidos revestidos por película	SE/H/0407/003	5002506	LABORATORIO NORMAL-PRODUTOS FARMACEUTICOS, LDA	PT
Tareg 40 mg comprimidos revestidos por película	SE/H/0407/003	5002514	LABORATORIO NORMAL-PRODUTOS FARMACEUTICOS, LDA	PT
Tareg 40 mg comprimidos revestidos por película	SE/H/0407/003	5002522	LABORATORIO NORMAL-PRODUTOS FARMACEUTICOS, LDA	PT
Tareg 40 mg comprimidos revestidos por película	SE/H/0407/003	5002530	LABORATORIO NORMAL-PRODUTOS FARMACEUTICOS, LDA	PT
Tareg 40 mg comprimidos revestidos por película	SE/H/0407/003	5002472	LABORATORIO NORMAL-PRODUTOS FARMACEUTICOS, LDA	PT
Tareg 40 mg comprimidos revestidos por película	SE/H/0407/003	5002506	LABORATORIO NORMAL-PRODUTOS FARMACEUTICOS, LDA	PT
Tareg 40 mg comprimidos revestidos por película	SE/H/0407/003	5002514	LABORATORIO NORMAL-PRODUTOS FARMACEUTICOS, LDA	PT
Tareg 40 mg comprimidos revestidos por película	SE/H/0407/003	5002522	LABORATORIO NORMAL-PRODUTOS FARMACEUTICOS, LDA	PT
Tareg 40 mg comprimidos revestidos por película	SE/H/0407/003	5002530	LABORATORIO NORMAL-PRODUTOS FARMACEUTICOS, LDA	PT
Tareg 40 mg comprimidos revestidos por película	SE/H/0407/003	5002472	LABORATORIO NORMAL-PRODUTOS FARMACEUTICOS, LDA	PT
Tareg 40 mg comprimidos revestidos por película	SE/H/0407/003	5002506	LABORATORIO NORMAL-PRODUTOS FARMACEUTICOS, LDA	PT

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
Tareg 40 mg comprimidos revestidos por película	SE/H/0407/003	5002514	LABORATORIO NORMAL-PRODUTOS FARMACEUTICOS, LDA	PT
Tareg 40 mg comprimidos revestidos por película	SE/H/0407/003	5002522	LABORATORIO NORMAL-PRODUTOS FARMACEUTICOS, LDA	PT
Tareg 40 mg comprimidos revestidos por película	SE/H/0407/003	5002530	LABORATORIO NORMAL-PRODUTOS FARMACEUTICOS, LDA	PT
Tareg 40 mg comprimidos revestidos por película	SE/H/0407/003	5002472	LABORATORIO NORMAL-PRODUTOS FARMACEUTICOS, LDA	PT
Tareg 40 mg comprimidos revestidos por película	SE/H/0407/003	5002506	LABORATORIO NORMAL-PRODUTOS FARMACEUTICOS, LDA	PT
Tareg 40 mg comprimidos revestidos por película	SE/H/0407/003	5002514	LABORATORIO NORMAL-PRODUTOS FARMACEUTICOS, LDA	PT
Tareg 40 mg comprimidos revestidos por película	SE/H/0407/003	5002522	LABORATORIO NORMAL-PRODUTOS FARMACEUTICOS, LDA	PT
Tareg 40 mg comprimidos revestidos por película	SE/H/0407/003	5002530	LABORATORIO NORMAL-PRODUTOS FARMACEUTICOS, LDA	PT
Tareg 40 mg comprimidos revestidos por película	SE/H/0407/003	5002472	LABORATORIO NORMAL-PRODUTOS FARMACEUTICOS, LDA	PT
Tareg 40 mg comprimidos revestidos por película	SE/H/0407/003	5002506	LABORATORIO NORMAL-PRODUTOS FARMACEUTICOS, LDA	PT
Tareg 40 mg comprimidos revestidos por película	SE/H/0407/003	5002514	LABORATORIO NORMAL-PRODUTOS FARMACEUTICOS, LDA	PT

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
Tareg 40 mg comprimidos revestidos por película	SE/H/0407/003	5002522	LABORATORIO NORMAL-PRODUTOS FARMACEUTICOS, LDA	PT
Tareg 40 mg comprimidos revestidos por película	SE/H/0407/003	5002530	LABORATORIO NORMAL-PRODUTOS FARMACEUTICOS, LDA	PT
TAREG 40 mg, comprimé pelliculé sécable	SE/H/0406/005	3400936958302	NOVARTIS PHARMA S.A.S.	FR
TAREG 40 mg, comprimé pelliculé sécable	SE/H/0406/005	3400936958470	NOVARTIS PHARMA S.A.S.	FR
TAREG 40 mg, comprimé pelliculé sécable	SE/H/0406/005	3400936958531	NOVARTIS PHARMA S.A.S.	FR
TAREG 40 mg, comprimé pelliculé sécable	SE/H/0406/005	3400937138062	NOVARTIS PHARMA S.A.S.	FR
TAREG 40 mg, comprimé pelliculé sécable	SE/H/0406/005	3400937138123	NOVARTIS PHARMA S.A.S.	FR
TAREG 40 mg, comprimé pelliculé sécable	SE/H/0406/005	3400938153811	NOVARTIS PHARMA S.A.S.	FR
TAREG 40 mg, comprimé pelliculé sécable	SE/H/0406/005	3400938153989	NOVARTIS PHARMA S.A.S.	FR
TAREG 40 mg, comprimé pelliculé sécable	SE/H/0406/005	3400938154061	NOVARTIS PHARMA S.A.S.	FR
TAREG 40 mg, comprimé pelliculé sécable	SE/H/0406/005	3400938154122	NOVARTIS PHARMA S.A.S.	FR
TAREG 40 mg, comprimé pelliculé sécable	SE/H/0406/005	3400938154351	NOVARTIS PHARMA S.A.S.	FR
Tareg 80 mg compresse rivestite con film	SE/H/0406/003	033178094	NOVARTIS EUROPHARM LIMITED	IT
Tareg 80 mg compresse rivestite con film	SE/H/0406/003	033178106	NOVARTIS EUROPHARM LIMITED	IT
Tareg 80 mg compresse rivestite con film	SE/H/0406/003	033178120	NOVARTIS EUROPHARM LIMITED	IT
Tareg 80 mg compresse	SE/H/0406/003	033178118	NOVARTIS EUROPHARM	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			LIMITED	
Tareg 80 mg compresse rivestite con film	SE/H/0406/003	033178625	NOVARTIS EUROPHARM LIMITED	IT
Tareg 80 mg compresse rivestite con film	SE/H/0406/003	033178132	NOVARTIS EUROPHARM LIMITED	IT
Tareg 80 mg compresse rivestite con film	SE/H/0406/003	033178649	NOVARTIS EUROPHARM LIMITED	IT
Tareg 80 mg compresse rivestite con film	SE/H/0406/003	033178637	NOVARTIS EUROPHARM LIMITED	IT
Tareg 80 mg compresse rivestite con film	SE/H/0406/003	033178652	NOVARTIS EUROPHARM LIMITED	IT
Tareg 80 mg compresse rivestite con film	SE/H/0406/003	033178664	NOVARTIS EUROPHARM LIMITED	IT
Tareg 80 mg compresse rivestite con film	SE/H/0406/003	033178676	NOVARTIS EUROPHARM LIMITED	IT
Tareg 80 mg compresse rivestite con film	SE/H/0406/003	033178688	NOVARTIS EUROPHARM LIMITED	IT
Tareg 80 mg compresse rivestite con film	SE/H/0406/003	033178878	NOVARTIS EUROPHARM LIMITED	IT
Tareg 80 mg compresse rivestite con film	SE/H/0406/003	033178866	NOVARTIS EUROPHARM LIMITED	IT
Tareg 80 mg compresse rivestite con film	SE/H/0406/003	033178854	NOVARTIS EUROPHARM LIMITED	IT
Tareg 80 mg compresse rivestite con film	SE/H/0406/003	033178892	NOVARTIS EUROPHARM LIMITED	IT
Tareg 80 mg compresse rivestite con film	SE/H/0406/003	033178880	NOVARTIS EUROPHARM LIMITED	IT
Tareg 80 mg compresse rivestite con film	SE/H/0406/003	033178916	NOVARTIS EUROPHARM LIMITED	IT
Tareg 80 mg compresse rivestite con film	SE/H/0406/003	033178904	NOVARTIS EUROPHARM LIMITED	IT
Tareg 80 mg compresse rivestite con film	SE/H/0406/003	033178928	NOVARTIS EUROPHARM LIMITED	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
Tareg 80 mg Comprimidos revestidos por película	SE/H/0407/001	3844883	LABORATORIO NORMAL- PRODUTOS FARMACEUTICOS, LDA	PT
Tareg 80 mg Comprimidos revestidos por película	SE/H/0407/001	3844982	LABORATORIO NORMAL- PRODUTOS FARMACEUTICOS, LDA	PT
Tareg 80 mg Comprimidos revestidos por película	SE/H/0407/001	3845088	LABORATORIO NORMAL- PRODUTOS FARMACEUTICOS, LDA	PT
Tareg 80 mg Comprimidos revestidos por película	SE/H/0407/001	5317383	LABORATORIO NORMAL- PRODUTOS FARMACEUTICOS, LDA	PT
Tareg 80 mg Comprimidos revestidos por película	SE/H/0407/001	5920780	LABORATORIO NORMAL- PRODUTOS FARMACEUTICOS, LDA	PT
Tareg 80 mg Comprimidos revestidos por película	SE/H/0407/001	3844883	LABORATORIO NORMAL- PRODUTOS FARMACEUTICOS, LDA	PT
Tareg 80 mg Comprimidos revestidos por película	SE/H/0407/001	3844982	LABORATORIO NORMAL- PRODUTOS FARMACEUTICOS, LDA	PT
Tareg 80 mg Comprimidos revestidos por película	SE/H/0407/001	3845088	LABORATORIO NORMAL- PRODUTOS FARMACEUTICOS, LDA	PT
Tareg 80 mg Comprimidos revestidos por película	SE/H/0407/001	5317383	LABORATORIO NORMAL- PRODUTOS FARMACEUTICOS, LDA	PT
Tareg 80 mg Comprimidos revestidos por película	SE/H/0407/001	5920780	LABORATORIO NORMAL- PRODUTOS FARMACEUTICOS, LDA	PT
Tareg 80 mg Comprimidos revestidos por película	SE/H/0407/001	3844883	LABORATORIO NORMAL- PRODUTOS FARMACEUTICOS, LDA	PT

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
Tareg 80 mg Comprimidos revestidos por película	SE/H/0407/001	3844982	LABORATORIO NORMAL-PRODUTOS FARMACEUTICOS, LDA	PT
Tareg 80 mg Comprimidos revestidos por película	SE/H/0407/001	3845088	LABORATORIO NORMAL-PRODUTOS FARMACEUTICOS, LDA	PT
Tareg 80 mg Comprimidos revestidos por película	SE/H/0407/001	5317383	LABORATORIO NORMAL-PRODUTOS FARMACEUTICOS, LDA	PT
Tareg 80 mg Comprimidos revestidos por película	SE/H/0407/001	5920780	LABORATORIO NORMAL-PRODUTOS FARMACEUTICOS, LDA	PT
Tareg 80 mg Comprimidos revestidos por película	SE/H/0407/001	3844883	LABORATORIO NORMAL-PRODUTOS FARMACEUTICOS, LDA	PT
Tareg 80 mg Comprimidos revestidos por película	SE/H/0407/001	3844982	LABORATORIO NORMAL-PRODUTOS FARMACEUTICOS, LDA	PT
Tareg 80 mg Comprimidos revestidos por película	SE/H/0407/001	3845088	LABORATORIO NORMAL-PRODUTOS FARMACEUTICOS, LDA	PT
Tareg 80 mg Comprimidos revestidos por película	SE/H/0407/001	5317383	LABORATORIO NORMAL-PRODUTOS FARMACEUTICOS, LDA	PT
Tareg 80 mg Comprimidos revestidos por película	SE/H/0407/001	5920780	LABORATORIO NORMAL-PRODUTOS FARMACEUTICOS, LDA	PT
Tareg 80 mg Comprimidos revestidos por película	SE/H/0407/001	3844883	LABORATORIO NORMAL-PRODUTOS FARMACEUTICOS, LDA	PT
Tareg 80 mg Comprimidos revestidos por película	SE/H/0407/001	3844982	LABORATORIO NORMAL-PRODUTOS FARMACEUTICOS, LDA	PT

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
Tareg 80 mg Comprimidos revestidos por película	SE/H/0407/001	3845088	LABORATORIO NORMAL-PRODUTOS FARMACEUTICOS, LDA	PT
Tareg 80 mg Comprimidos revestidos por película	SE/H/0407/001	5317383	LABORATORIO NORMAL-PRODUTOS FARMACEUTICOS, LDA	PT
Tareg 80 mg Comprimidos revestidos por película	SE/H/0407/001	5920780	LABORATORIO NORMAL-PRODUTOS FARMACEUTICOS, LDA	PT
Tareg 80 mg Comprimidos revestidos por película	SE/H/0407/001	3844883	LABORATORIO NORMAL-PRODUTOS FARMACEUTICOS, LDA	PT
Tareg 80 mg Comprimidos revestidos por película	SE/H/0407/001	3844982	LABORATORIO NORMAL-PRODUTOS FARMACEUTICOS, LDA	PT
Tareg 80 mg Comprimidos revestidos por película	SE/H/0407/001	3845088	LABORATORIO NORMAL-PRODUTOS FARMACEUTICOS, LDA	PT
Tareg 80 mg Comprimidos revestidos por película	SE/H/0407/001	5317383	LABORATORIO NORMAL-PRODUTOS FARMACEUTICOS, LDA	PT
Tareg 80 mg Comprimidos revestidos por película	SE/H/0407/001	5920780	LABORATORIO NORMAL-PRODUTOS FARMACEUTICOS, LDA	PT
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	SE/H/0406/003	34009 371 386 4 2	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
pelliculé				
TAREG 80 mg, comprimé pelliculé	SE/H/0406/003	3400938154412	NOVARTIS PHARMA S.A.S.	FR
TAREG 80 mg, comprimé pelliculé	SE/H/0406/003	3400938154580	NOVARTIS PHARMA S.A.S.	FR
TAREG 80 mg, comprimé pelliculé	SE/H/0406/003	3400938154641	NOVARTIS PHARMA S.A.S.	FR
TAREG 80 mg, comprimé pelliculé	SE/H/0406/003	3400938154702	NOVARTIS PHARMA S.A.S.	FR
TAREG 80 mg, comprimé pelliculé	SE/H/0406/003	3400938154931	NOVARTIS PHARMA S.A.S.	FR
Troval 160 mg Film-coated tablets	not available	ML 21334	DELORBIS PHARMACEUTICALS LTD	CY
Troval 80 mg Film-coated tablets	not available	ML 21333	DELORBIS PHARMACEUTICALS LTD	CY
Valpression 160 mg capsule	not available	033119025	A. MENARINI - INDUSTRIE FARMACEUTICHE RIUNITE - S.R.L.	IT
Valpression 160 mg compresse rivestite con film	not available	033119090	A. MENARINI - INDUSTRIE FARMACEUTICHE RIUNITE - S.R.L.	IT
Valpression 160 mg compresse rivestite con film	not available	033119102	A. MENARINI - INDUSTRIE FARMACEUTICHE RIUNITE - S.R.L.	IT
Valpression 160 mg compresse rivestite con film	not available	033119114	A. MENARINI - INDUSTRIE FARMACEUTICHE RIUNITE - S.R.L.	IT
Valpression 160 mg compresse rivestite con film	not available	033119126	A. MENARINI - INDUSTRIE FARMACEUTICHE RIUNITE - S.R.L.	IT
Valpression 160 mg compresse rivestite con film	not available	033119088	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 3 mg/ml soluzione orale	not available	033119429	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 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 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 320 mg compresse rivestite con film	not available	033119355	A. MENARINI - INDUSTRIE FARMACEUTICHE RIUNITE - S.R.L.	IT
Valpression 40 mg compresse rivestite con film	not available	033119239	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 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 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
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 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 con film	not available	033119191	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 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 80 mg compresse rivestite con film	not available	033119037	A. MENARINI - INDUSTRIE FARMACEUTICHE RIUNITE - S.R.L.	IT
Valpression 80 mg compresse rivestite con film	not available	033119049	A. MENARINI - INDUSTRIE FARMACEUTICHE RIUNITE - S.R.L.	IT
Valpression 80 mg compresse rivestite con film	not available	033119064	A. MENARINI - INDUSTRIE FARMACEUTICHE RIUNITE - S.R.L.	IT
Valpression 80 mg compresse rivestite con film	not available	033119076	A. MENARINI - INDUSTRIE FARMACEUTICHE RIUNITE - S.R.L.	IT
Valpression 80 mg compresse rivestite con film	not available	033119052	A. MENARINI - INDUSTRIE FARMACEUTICHE RIUNITE - S.R.L.	IT
VALS 160 mg comprimidos recubiertos con película	not available	64.494	LABORATORIOS DR. ESTEVE S.A.	ES
VALS 320 mg comprimidos recubiertos con película	not available	69.345	LABORATORIOS DR. ESTEVE S.A.	ES
VALS 80 mg comprimidos recubiertos con película	not available	64.495	LABORATORIOS DR. ESTEVE S.A.	ES

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 ratiopharm 120mg/12,5mg Filmtabletten	AT/H/0514/001	135567	RATIOPHARM ARZNEIMITTEL VERTRIEBS-GMBH	AT
Valsartan / HCT TEVA 120mg/12,5mg Filmtabletten	AT/H/0515/001	135568	RATIOPHARM ARZNEIMITTEL VERTRIEBS-GMBH	AT
Valsartan + Hydrochlorothiazide Aurovitas, 160 mg + 12,5 mg, tabletki powlekane	PT/H/1594/002	23888	AUROVITAS UNIPESOAL, LDA.	PL
Valsartan + Hydrochlorothiazide Aurovitas, 320 mg + 12,5 mg, tabletki powlekane	PT/H/1594/004	23890	AUROVITAS UNIPESOAL, LDA.	PL
Valsartan + Hydrochlorothiazide Aurovitas, 80 mg + 12,5 mg, tabletki powlekane	PT/H/1594/001	23887	AUROVITAS UNIPESOAL, LDA.	PL
Valsartan 160 mg Capsules	not available	PL 04416/1273	SANDOZ LTD	UK
Valsartan 160 mg film-coated tablets	not available	PL 17907/0362	BRISTOL LABORATORIES LTD (BERKHAMSTED)	UK
Valsartan 320 mg film-coated tablets	not available	PL 17907/0363	BRISTOL LABORATORIES LTD (BERKHAMSTED)	UK
Valsartan 40 mg Capsules	not available	PL 04416/1271	SANDOZ LTD	UK
Valsartan 40 mg Film-coated Tablets	DK/H/1517/001	PL 00289/1235	TEVA UK LIMITED	UK
Valsartan 40 mg film-coated tablets	not available	PL 17907/0360	BRISTOL LABORATORIES LTD (BERKHAMSTED)	UK

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 80 mg Capsules	not available	PL 04416/1272	SANDOZ LTD	UK
Valsartan 80 mg film-coated tablets	not available	PL 17907/0361	BRISTOL LABORATORIES LTD (BERKHAMSTED)	UK
Valsartan AbZ 120 mg Filmtabletten	SE/H/0746/003	70447.00.00	ABZ-PHARMA GMBH	DE
Valsartan Actavis 40 mg plėvele dengtos tabletės	IS/H/0105/001	LT/1/08/1220/001-012	ACTAVIS GROUP PTC EHF.	LT
Valsartan comp. AbZ 120 mg/12,5 mg Filmtabletten	AT/H/0515/001	90714.00.00	ABZ-PHARMA GMBH	DE
Valsartan HCT 160/12,5mg Actavis plevele dengtos tabletės	IS/H/0194/002	LT/1/11/2601/015-028	ACTAVIS GROUP PTC EHF.	LT
Valsartan HCT 160/25mg Actavis plevele dengtos tabletės	IS/H/0194/003	LT/1/11/2601/029-042	ACTAVIS GROUP PTC EHF.	LT
Valsartan HCT Actavis 320 mg/12,5 mg plėvele dengtos tabletės	FI/H/0857/001	LT/1/13/3193/001	ACTAVIS GROUP PTC EHF.	LT
Valsartan HCT Actavis 320 mg/12,5 mg plėvele dengtos tabletės	FI/H/0857/001	LT/1/13/3193/002	ACTAVIS GROUP PTC EHF.	LT
Valsartan HCT Actavis 320 mg/12,5 mg plėvele dengtos tabletės	FI/H/0857/001	LT/1/13/3193/003	ACTAVIS GROUP PTC EHF.	LT
Valsartan HCT Actavis 320 mg/12,5 mg plėvele dengtos tabletės	FI/H/0857/001	LT/1/13/3193/004	ACTAVIS GROUP PTC EHF.	LT
Valsartan HCT Actavis 320 mg/12,5 mg plėvele dengtos tabletės	FI/H/0857/001	LT/1/13/3193/005	ACTAVIS GROUP PTC EHF.	LT
Valsartan HCT Actavis	FI/H/0857/001	LT/1/13/3193/006	ACTAVIS GROUP PTC EHF.	LT

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
320 mg/12,5 mg plêvele dengtos tabletės				
Valsartan HCT Actavis 320 mg/12,5 mg plêvele dengtos tabletės	FI/H/0857/001	LT/1/13/3193/007	ACTAVIS GROUP PTC EHF.	LT
Valsartan HCT Actavis 320 mg/12,5 mg plêvele dengtos tabletės	FI/H/0857/001	LT/1/13/3193/008	ACTAVIS GROUP PTC EHF.	LT
Valsartan HCT Actavis 320 mg/12,5 mg plêvele dengtos tabletės	FI/H/0857/001	LT/1/13/3193/009	ACTAVIS GROUP PTC EHF.	LT
Valsartan HCT Actavis 320 mg/12,5 mg plêvele dengtos tabletės	FI/H/0857/001	LT/1/13/3193/010	ACTAVIS GROUP PTC EHF.	LT
Valsartan HCT Actavis 320 mg/12,5 mg plêvele dengtos tabletės	FI/H/0857/001	LT/1/13/3193/011	ACTAVIS GROUP PTC EHF.	LT
Valsartan HCT Actavis 320 mg/25 mg plêvele dengtos tabletės	FI/H/0857/002	LT/1/13/3193/012	ACTAVIS GROUP PTC EHF.	LT
Valsartan HCT Actavis 320 mg/25 mg plêvele dengtos tabletės	FI/H/0857/002	LT/1/13/3193/013	ACTAVIS GROUP PTC EHF.	LT
Valsartan HCT Actavis 320 mg/25 mg plêvele dengtos tabletės	FI/H/0857/002	LT/1/13/3193/015	ACTAVIS GROUP PTC EHF.	LT
Valsartan HCT Actavis 320 mg/25 mg plêvele dengtos tabletės	FI/H/0857/002	LT/1/13/3193/014	ACTAVIS GROUP PTC EHF.	LT
Valsartan HCT Actavis 320 mg/25 mg plêvele dengtos tabletės	FI/H/0857/002	LT/1/13/3193/016	ACTAVIS GROUP PTC EHF.	LT
Valsartan HCT Actavis	FI/H/0857/002	LT/1/13/3193/017	ACTAVIS GROUP PTC EHF.	LT

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
320 mg/25 mg plévele dengtos tabletés				
Valsartan HCT Actavis 320 mg/25 mg plévele dengtos tabletés	FI/H/0857/002	LT/1/13/3193/018	ACTAVIS GROUP PTC EHF.	LT
Valsartan HCT Actavis 320 mg/25 mg plévele dengtos tabletés	FI/H/0857/002	LT/1/13/3193/019	ACTAVIS GROUP PTC EHF.	LT
Valsartan HCT Actavis 320 mg/25 mg plévele dengtos tabletés	FI/H/0857/002	LT/1/13/3193/020	ACTAVIS GROUP PTC EHF.	LT
Valsartan HCT Actavis 320 mg/25 mg plévele dengtos tabletés	FI/H/0857/002	LT/1/13/3193/021	ACTAVIS GROUP PTC EHF.	LT
Valsartan HCT Actavis 320 mg/25 mg plévele dengtos tabletés	FI/H/0857/002	LT/1/13/3193/022	ACTAVIS GROUP PTC EHF.	LT
Valsartan Novartis 320 mg	SE/H/0408/004	22842	NOVARTIS SVERIGE AB	SE
Valsartan Novartis 320 mg	SE/H/0408/004	22842	NOVARTIS SVERIGE AB	SE
Valsartan Novartis 320 mg	SE/H/0408/004	22842	NOVARTIS SVERIGE AB	SE
Valsartan Novartis 320 mg	SE/H/0408/004	22842	NOVARTIS SVERIGE AB	SE
Valsartan Novartis 320 mg	SE/H/0408/004	22842	NOVARTIS SVERIGE AB	SE
Valsartan Novartis 40 mg filmdragerad tablett	SE/H/0408/003	20466	NOVARTIS SVERIGE AB	SE
Valsartan Novartis 40 mg filmdragerad tablett	SE/H/0408/003	20466	NOVARTIS SVERIGE AB	SE
Valsartan Novartis 40 mg filmdragerad tablett	SE/H/0408/003	20466	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 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 80 mg filmdragerade tabletter	SE/H/0408/001	19957	NOVARTIS SVERIGE AB	SE
Valsartan Novartis 80 mg filmdragerade tabletter	SE/H/0408/001	19957	NOVARTIS SVERIGE AB	SE
Valsartan Novartis 80 mg filmdragerade tabletter	SE/H/0408/001	19957	NOVARTIS SVERIGE AB	SE
Valsartan Novartis 80 mg filmdragerade tabletter	SE/H/0408/001	19957	NOVARTIS SVERIGE AB	SE
Valsartan ratiopharm 120 mg Filmtabletten	SE/H/0747/003	1-29314	RATIOPHARM ARZNEIMITTEL VERTRIEBS-GMBH	AT
Valsartan ratiopharm 40 mg film-coated tablets	SE/H/0747/001	26039	RATIOPHARM GMBH	SE
Valsartan ratiopharm, 120 mg filmdragerad tablett	SE/H/0747/003	26041	RATIOPHARM GMBH	SE
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

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 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 rivestite con film	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
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 Teva Pharma 120 mg filmdragerad tablett	SE/H/0748/003	26045	TEVA PHARMA B.V.	SE
Valsartan//Hydroklortiazid Novartis 160 mg/25 mg filmdragerade tabletter	SE/H/0567/002	22461	NOVARTIS SVERIGE AB	SE
Valsartan//Hydroklortiazid Novartis 160 mg/25 mg filmdragerade tabletter	SE/H/0567/002	22461	NOVARTIS SVERIGE AB	SE
Valsartan/Hydrochlorothi azid Aurovitas 320 mg/12,5 mg potahované tablety	PT/H/1594/004	58/805/15-C	AUROVITAS UNIPessoal, LDA.	CZ
Valsartan/Hydroklortiazid Novartis 160 mg/12,5 mg filmdragerade tabletter	SE/H/0567/001	22460	NOVARTIS SVERIGE AB	SE
Valsartan/Hydroklortiazid Novartis 160 mg/12,5 mg filmdragerade	SE/H/0567/001	22460	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
tabletter				
Valsartan/Hydroklortiazid Novartis 320 mg/12,5 mg filmdragerade tabletter	SE/H/0567/004	23645	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 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
Valsartan/Hydroklortiazid Novartis 80 mg/12,5 mg filmdragerade tabletter	SE/H/0567/003	22462	NOVARTIS SVERIGE AB	SE
Valsartan-CT 120 mg Filmtabletten	SE/H/0748/003	70455.00.00	ABZ-PHARMA GMBH	DE
Valsartan-HCT-Teva 120 mg/12,5 mg filmtabletta	AT/H/0514/001	OGYI-T-20990/06	TEVA GYÓGYSZERGYÁR ZRT	HU
Valsartan-HCT-Teva 120 mg/12,5 mg filmtabletta	AT/H/0514/001	OGYI-T-20990/07	TEVA GYÓGYSZERGYÁR ZRT	HU
Valsartan-HCT-Teva 120 mg/12,5 mg filmtabletta	AT/H/0514/001	OGYI-T-20990/05	TEVA GYÓGYSZERGYÁR ZRT	HU
Valsartan-HCT-Teva 120 mg/12,5 mg filmtabletta	AT/H/0514/001	OGYI-T-20990/08	TEVA GYÓGYSZERGYÁR ZRT	HU
Valsartan-Novartis 160 mg filmdragerade	SE/H/0408/002	19958	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
tabletter				
Valsartan-Novartis 160 mg filmdragerade tabletter	SE/H/0408/002	19958	NOVARTIS SVERIGE AB	SE
Valsartan-Novartis 160 mg filmdragerade tabletter	SE/H/0408/002	19958	NOVARTIS SVERIGE AB	SE
Valsartan-Novartis 160 mg filmdragerade tabletter	SE/H/0408/002	19958	NOVARTIS SVERIGE AB	SE
Valsartan-Novartis 160 mg filmdragerade tabletter	SE/H/0408/002	19958	NOVARTIS SVERIGE AB	SE
Valsartan-ratiopharm comp. 120 mg/12,5 mg Filmdragerade tabletter	AT/H/0514/001	90713.00.00	RATIOPHARM GMBH	DE
Valsartan-ratiopharm® 120 mg Filmdragerade tabletter	SE/H/0747/003	70451.00.00	RATIOPHARM GMBH	DE
Valsartan-ratiopharm® 160 mg Filmdragerade tabletter	SE/H/0747/004	0687/10100058	RATIOPHARM GMBH	LU
Valsartan-ratiopharm® 40 mg Filmdragerade tabletter	SE/H/0747/001	0687/10100054	RATIOPHARM GMBH	LU
Valsartan-ratiopharm® 80 mg Filmdragerade tabletter	SE/H/0747/002	0687/10100057	RATIOPHARM GMBH	LU
Valsocard 160 mg fimtablett	not available	OGYI-T-21874/15-21	ACTAVIS GROUP PTC EHF.	HU
Valsocard 160 mg fimtablett	not available	OGYI-T-21874/15-21	ACTAVIS GROUP PTC EHF.	HU
Valsocard 40 mg fimtablett	OGYI-T-21874/01-07	OGYI-T-21874/01-07	ACTAVIS GROUP PTC EHF.	HU
Valsocard 40 mg fimtablett	OGYI-T-21874/01-07	OGYI-T-21874/01-07	ACTAVIS GROUP PTC EHF.	HU
Valsocard 80 mg	OGYI-T-21874/08-14	OGYI-T-21874/08-14	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
fimtabletta				
Valsocard 80 mg fimtabletta	OGYI-T-21874/08-14	OGYI-T-21874/08-14	ACTAVIS GROUP PTC EHF.	HU
Valsocard HCT 160 mg/12.5 mg fimtabletta	HU/N/0001	OGYI-T-21875/08-14	ACTAVIS GROUP PTC EHF.	HU
Valsocard HCT 160 mg/25 mg fimtabletta	HU/N/0001	OGYI-T-21875/15-21	ACTAVIS GROUP PTC EHF.	HU
Valsocard HCT 80 mg/12.5 mg fimtabletta	HU/N/0001	OGYI-T-21875/01-07	ACTAVIS GROUP PTC EHF.	HU
Valtan HCT 320 mg/12,5 mg filmom obalené tablety	SE/H/0994/001	58/0176/11-S	RATIOPHARM GMBH	SK
Valtan HCT 320 mg/25 mg filmom obalené tablety	SE/H/0994/002	58/0177/11-S	RATIOPHARM GMBH	SK
Valtsu 120 mg filmdragerade tabletter	SE/H/0746/003	26037	TEVA SWEDEN AB	SE
ДИОВАН 160 MG ФИЛМИРАНИ ТАБЛЕТКИ	SE/H/0406/004	20020028	NOVARTIS PHARMA GMBH	BG
ДИОВАН 3 MG/ML ПЕРОРАЛЕН РАЗТВОР	SE/H/0406/007	20100328	NOVARTIS PHARMA GMBH	BG
ДИОВАН 40 MG ФИЛМИРАНИ ТАБЛЕТКИ	SE/H/0406/005	20050206	NOVARTIS PHARMA GMBH	BG
КО-ДИОВАН 160 MG/12,5 MG ФИЛМИРАНИ ТАБЛЕТКИ	SE/H/0565/002	20020026	NOVARTIS PHARMA GMBH	BG
КО-ДИОВАН 160 MG/12,5 MG ФИЛМИРАНИ ТАБЛЕТКИ	SE/H/0565/002	20020026	NOVARTIS PHARMA GMBH	BG
КО-ДИОВАН 160 MG/12,5 MG ФИЛМИРАНИ ТАБЛЕТКИ	SE/H/0565/002	20020026	NOVARTIS PHARMA GMBH	BG
КО-ДИОВАН 160	SE/H/0565/002	20020026	NOVARTIS PHARMA GMBH	BG

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
MG/12,5 MG ФИЛМИРАНИ ТАБЛЕТКИ				
КО-ДИОВАН 160 MG/25 MG ФИЛМИРАНИ ТАБЛЕТКИ	SE/H/0565/003	20040085	NOVARTIS PHARMA GMBH	BG
КО-ДИОВАН 160 MG/25 MG ФИЛМИРАНИ ТАБЛЕТКИ	SE/H/0565/003	20040085	NOVARTIS PHARMA GMBH	BG
КО-ДИОВАН 160 MG/25 MG ФИЛМИРАНИ ТАБЛЕТКИ	SE/H/0565/003	20040085	NOVARTIS PHARMA GMBH	BG
КО-ДИОВАН 160 MG/25 MG ФИЛМИРАНИ ТАБЛЕТКИ	SE/H/0565/003	20040085	NOVARTIS PHARMA GMBH	BG