



EUROPEAN MEDICINES AGENCY  
SCIENCE MEDICINES HEALTH

09 June 2017  
EMA/379727/2017  
Procedure Management and Committees Support

## List of nationally authorised medicinal products

Active substance: hydrochlorothiazide / olmesartan

Procedure no.: PSUSA/00002209/201610



<b>Product Name (in authorisation country)</b>	<b>MRP/DCP Authorisation number</b>	<b>National Authorisation Number</b>	<b>MAH of product in the member state</b>	<b>Member State where product is authorised</b>
PLAUNAZIDE 20 mg/25 mg compresse rivestite con film	DE/H/0524/002	037108204	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
PLAUNAZIDE 20 mg/25 mg compresse rivestite con film	DE/H/0524/002	037108127	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
PLAUNAZIDE 20 mg/12,5 mg compresse rivestite con film	DE/H/0524/001	037108103	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
PLAUNAZIDE 20 mg/12,5 mg compresse rivestite con film	DE/H/0524/001	037108014	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
PLAUNAZIDE 20 mg/12,5 mg compresse rivestite con film	DE/H/0524/001	037108091	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
PLAUNAZIDE 20 mg/12,5 mg compresse rivestite con film	DE/H/0524/001	037108038	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
PLAUNAZIDE 20 mg/12,5 mg compresse rivestite con film	DE/H/0524/001	037108115	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
PLAUNAZIDE 20 mg/12,5 mg compresse rivestite con film	DE/H/0524/001	037108077	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
PLAUNAZIDE 20 mg/25 mg compresse rivestite con film	DE/H/0524/002	037108141	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT

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PLAUNAZIDE 20 mg/25 mg compresse rivestite con film	DE/H/0524/002	037108216	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
PLAUNAZIDE 20 mg/25 mg compresse rivestite con film	DE/H/0524/002	037108139	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
PLAUNAZIDE 20 mg/25 mg compresse rivestite con film	DE/H/0524/002	037108166	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
PLAUNAZIDE 20 mg/25 mg compresse rivestite con film	DE/H/0524/002	037108178	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
PLAUNAZIDE 20 mg/25 mg compresse rivestite con film	DE/H/0524/002	037108228	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
PLAUNAZIDE 20 mg/25 mg compresse rivestite con film	DE/H/0524/002	037108154	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
PLAUNAZIDE 20 mg/25 mg compresse rivestite con film	DE/H/0524/002	037108180	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
PLAUNAZIDE 20 mg/12,5 mg compresse rivestite con film	DE/H/0524/001	037108065	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
PLAUNAZIDE 20 mg/25 mg compresse rivestite con film	DE/H/0524/002	037108192	MENARINI INTERNATIONAL OPERATIONS	IT

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			LUXEMBOURG S.A.	
PLAUNAZIDE 20 mg/12,5 mg compresse rivestite con film	DE/H/0524/001	037108026	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
PLAUNAZIDE 20 mg/12,5 mg compresse rivestite con film	DE/H/0524/001	037108089	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
PLAUNAZIDE 20 mg/12,5 mg compresse rivestite con film	DE/H/0524/001	037108040	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
PLAUNAZIDE 20 mg/12,5 mg compresse rivestite con film	DE/H/0524/001	037108053	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
Benetor Comp 20 mg/12,5 mg filmdragerade tabletter	DE/H/0525/001	21334	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	FI
Benetor Comp 20 mg/25 mg filmdragerade tabletter	DE/H/0525/002	21335	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	FI
BELSAR Plus 20mg/25mg filmomhulde tabletten	DE/H/0525/002	BE282791	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	BE
BELSAR Plus 20mg/12,5mg filmomhulde tabletten	DE/H/0525/001	BE282782	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	BE
Belsar Plus 20 mg/12,5 mg Filmtabletten	DE/H/0525/001	BE282782	MENARINI INTERNATIONAL	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
			OPERATIONS LUXEMBOURG S.A.	
Belsar Plus 20 mg/25 mg Filmtabletten	DE/H/0525/002	BE282791	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	BE
Olartan-Plus® 20 mg/12,5 mg επικαλυμμένο με λεπτό υμένιο δισκία	DE/H/0525/001	4744	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	GR
Olartan-Plus® 20 mg/25 mg επικαλυμμένο με λεπτό υμένιο δισκία	DE/H/0525/002	4745	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	GR
Co-Tensiol 20 mg/12,5 mg filmsko obložene tablete	DE/H/0525/001	H/06/00426/010	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	SI
Co-Tensiol 20 mg/25 mg filmsko obložene tablete	DE/H/0525/002	H/06/00426/015	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	SI
Co-Tensiol 20 mg/12,5 mg filmsko obložene tablete	DE/H/0525/001	H/06/00426/009	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	SI
Co-Tensiol 20 mg/25 mg filmsko obložene tablete	DE/H/0525/002	H/06/00426/016	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	SI
Co-Tensiol 20 mg/12,5 mg filmsko obložene tablete	DE/H/0525/001	H/06/00426/003	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	SI
Co-Tensiol 20 mg/12,5 mg	DE/H/0525/001	H/06/00426/006	MENARINI	SI

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filmsko obložene tablete			INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	
Co-Tensiol 20 mg/12,5 mg filmsko obložene tablete	DE/H/0525/001	H/06/00426/007	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	SI
Co-Tensiol 20 mg/12,5 mg filmsko obložene tablete	DE/H/0525/001	H/06/00426/005	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	SI
Co-Tensiol 20 mg/25 mg filmsko obložene tablete	DE/H/0525/002	H/06/00426/017	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	SI
Co-Tensiol 20 mg/25 mg filmsko obložene tablete	DE/H/0525/002	H/06/00426/014	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	SI
Co-Tensiol 20 mg/12,5 mg filmsko obložene tablete	DE/H/0525/001	H/06/00426/011	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	SI
Co-Tensiol 20 mg/25 mg filmsko obložene tablete	DE/H/0525/002	H/06/00426/018	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	SI
Co-Tensiol 20 mg/25 mg filmsko obložene tablete	DE/H/0525/002	H/06/00426/019	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	SI
Co-Tensiol 20 mg/12,5 mg filmsko obložene tablete	DE/H/0525/001	H/06/00426/004	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	SI

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Co-Tensiol 20 mg/12,5 mg filmsko obložene tablete	DE/H/0525/001	H/06/00426/008	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	SI
Co-Tensiol 20 mg/25 mg filmsko obložene tablete	DE/H/0525/002	H/06/00426/021	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	SI
Co-Tensiol 20 mg/25 mg filmsko obložene tablete	DE/H/0525/002	H/06/00426/022	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	SI
Co-Tensiol 20 mg/25 mg filmsko obložene tablete	DE/H/0525/002	H/06/00426/020	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	SI
BELSAR PLUS 20 mg/25 mg comprimés pelliculés	DE/H/0525/002	BE282791	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	BE
Mencord® Plus 20 mg/12,5 mg Filmtabletten	DE/H/0525/001	1-26160	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	AT
Mencord® Plus 20 mg/25 mg Filmtabletten	DE/H/0525/002	1-26161	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	AT
BELSAR PLUS 20 mg/12,5 mg comprimés pelliculés	DE/H/0525/001	BE282782	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	BE
Olartan-Plus® 20 mg/25 mg επικαλυμμένο με λεπτό υμένιο δισκία	DE/H/0525/002	20334	MENARINI INTERNATIONAL OPERATIONS	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
			LUXEMBOURG S.A.	
Sarten Plus H 20/25 mg potahované tablety	DE/H/0525/002	58/459/05-C	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	CZ
Benetor Comp 20 mg/25 mg kalvopäällysteiset tabletit	DE/H/0525/002	21335	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	FI
Benetor Comp, filmovertukne tablettar	DE/H/0525/002	38152	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	DK
Mesar plus, 20 mg/12,5 mg õhukese polümeerikattega tabletid	DE/H/0525/001	501105	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	EE
Benetor Comp, filmovertukne tablettar	DE/H/0525/001	38151	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	DK
Olartan-Plus® 20 mg/12,5 mg επικαλυμμένο με λεπτό υμένιο δισκί	DE/H/0525/001	20333	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	CY
Mesar plus, 20 mg/25 mg õhukese polümeerikattega tabletid	DE/H/0525/002	501005	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	EE
Sarten Plus H 20/12,5 mg potahované tablety	DE/H/0525/001	58/458/05-C	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	CZ
Benetor Comp 20 mg/12,5 mg kalvopäällysteiset tabletit	DE/H/0525/001	21334	MENARINI INTERNATIONAL	FI



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			OPERATIONS LUXEMBOURG S.A.	
Laresin Plus 20 mg/25 mg filmtabletta	DE/H/0525/002	OGYI-T-20333/04	BERLIN-CHEMIE AG	HU
Laresin Plus 20 mg/25 mg filmtabletta	DE/H/0525/002	OGYI-T-20333/03	BERLIN-CHEMIE AG	HU
Laresin Plus 20 mg/12,5 mg filmtabletta	DE/H/0525/001	OGYI-T-20333/02	BERLIN-CHEMIE AG	HU
Laresin Plus 20 mg/12,5 mg filmtabletta	DE/H/0525/001	OGYI-T-20333/01	BERLIN-CHEMIE AG	HU
Benetor Comp 20 mg/12,5 mg filmuhúðaðar töflur	DE/H/0525/001	IS/1/05/048/01	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IS
Benetor Comp 20 mg/25 mg filmuhúðaðar töflur	DE/H/0525/002	IS/1/05/048/02	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IS
ALTEISDUO 20 mg/25 mg, comprimé pelliculé	DE/H/0525/002	34009 372 221 9 8	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	FR
ALTEISDUO 20 mg/25 mg, comprimé pelliculé	DE/H/0525/002	34009 372 222 5 9	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	FR
ALTEISDUO 20 mg/12,5 mg, comprimé pelliculé	DE/H/0525/001	34009 372 219 4 8	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	FR
ALTEISDUO 20 mg/25 mg, comprimé pelliculé	DE/H/0525/002	34009 567 674 2 7	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	FR
Votum® plus 20 mg/12,5 mg	DE/H/0525/001	58597.00.00	BERLIN-CHEMIE AG	DE

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Filmtabletten				
ALTEISDUO 20 mg/12,5 mg, comprimé pelliculé	DE/H/0525/001	34009 372 220 2 0	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	FR
ALTEISDUO 20 mg/12,5 mg, comprimé pelliculé	DE/H/0525/001	34009 567 673 6 6	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	FR
Votum® plus 20 mg/25 mg Filmtabletten	DE/H/0525/002	58597.01.00	BERLIN-CHEMIE AG	DE
Mesar Plus 20 mg/25 mg apvalkotās tabletes	DE/H/0525/002	05-0632	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	LV
Mesar Plus 20 mg/12,5 mg apvalkotās tabletes	DE/H/0525/001	05-0631	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	LV
OLPREZIDE 20 mg/25 mg compresse rivestite con film	DE/H/0525/002	037109182	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
OLPREZIDE 20 mg/12,5 mg compresse rivestite con film	DE/H/0525/001	037109081	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
OLPREZIDE 20 mg/12,5 mg compresse rivestite con film	DE/H/0525/001	037109030	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
OLPREZIDE 20 mg/12,5 mg compresse rivestite con film	DE/H/0525/001	037109117	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT

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Omesar Plus 20 mg /25 mg film-coated tablets	DE/H/0525/002	PA 865/14/2	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IE
OLPREZIDE 20 mg/12,5 mg compresse rivestite con film	DE/H/0525/001	037109055	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
OLPREZIDE 20 mg/12,5 mg compresse rivestite con film	DE/H/0525/001	037109016	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
Omesar Plus 20 mg /12.5 mg film-coated tablets	DE/H/0525/001	PA 865/14/1	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IE
OLPREZIDE 20 mg/12,5 mg compresse rivestite con film	DE/H/0525/001	037109079	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
OLPREZIDE 20 mg/12,5 mg compresse rivestite con film	DE/H/0525/001	037109042	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
OLPREZIDE 20 mg/12,5 mg compresse rivestite con film	DE/H/0525/001	037109028	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
OLPREZIDE 20 mg/12,5 mg compresse rivestite con film	DE/H/0525/001	037109093	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
OLPREZIDE 20 mg/12,5 mg compresse rivestite con film	DE/H/0525/001	037109105	MENARINI INTERNATIONAL OPERATIONS	IT

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			LUXEMBOURG S.A.	
OLPREZIDE 20 mg/12,5 mg compresse rivestite con film	DE/H/0525/001	037109067	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
OLPREZIDE 20 mg/25 mg compresse rivestite con film	DE/H/0525/002	037109206	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
OLPREZIDE 20 mg/25 mg compresse rivestite con film	DE/H/0525/002	037109131	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
OLPREZIDE 20 mg/25 mg compresse rivestite con film	DE/H/0525/002	037109129	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
OLPREZIDE 20 mg/25 mg compresse rivestite con film	DE/H/0525/002	037109143	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
OLPREZIDE 20 mg/25 mg compresse rivestite con film	DE/H/0525/002	037109194	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
OLPREZIDE 20 mg/25 mg compresse rivestite con film	DE/H/0525/002	037109156	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
OLPREZIDE 20 mg/25 mg compresse rivestite con film	DE/H/0525/002	037109220	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
OLPREZIDE 20 mg/25 mg compresse rivestite con film	DE/H/0525/002	037109218	MENARINI INTERNATIONAL	IT

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			OPERATIONS LUXEMBOURG S.A.	
OLPREZIDE 20 mg/25 mg compresse rivestite con film	DE/H/0525/002	037109168	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
Mesar plus 20 mg/12,5 mg plévele dengtos tabletès	DE/H/0525/001	LT/1/05/0436/037	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	LT
Mesar plus 20 mg/12,5 mg plévele dengtos tabletès	DE/H/0525/001	LT/1/05/0436/002	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	LT
Mesar plus 20 mg/12,5 mg plévele dengtos tabletès	DE/H/0525/001	LT/1/05/0436/001	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	LT
OLPREZIDE 20 mg/25 mg compresse rivestite con film	DE/H/0525/002	037109170	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
Mesar plus 20 mg/25 mg plévele dengtos tabletès	DE/H/0525/002	LT/1/05/0436/044	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	LT
Mesar plus 20 mg/12,5 mg plévele dengtos tabletès	DE/H/0525/001	LT/1/05/0436/036	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	LT
Mesar plus 20 mg/12,5 mg plévele dengtos tabletès	DE/H/0525/001	LT/1/05/0436/004	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	LT
Mesar plus 20 mg/12,5 mg	DE/H/0525/001	LT/1/05/0436/003	MENARINI	LT

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plèvele dengtos tabletès			INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	
Mesar plus 20 mg/25 mg plèvele dengtos tabletès	DE/H/0525/002	LT/1/05/0436/007	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	LT
Mesar plus 20 mg/25 mg plèvele dengtos tabletès	DE/H/0525/002	LT/1/05/0436/046	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	LT
Mesar plus 20 mg/25 mg plèvele dengtos tabletès	DE/H/0525/002	LT/1/05/0436/041	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	LT
Mesar plus 20 mg/25 mg plèvele dengtos tabletès	DE/H/0525/002	LT/1/05/0436/045	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	LT
Mesar plus 20 mg/25 mg plèvele dengtos tabletès	DE/H/0525/002	LT/1/05/0436/042	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	LT
BELSAR PLUS 20 mg/25 mg comprimés pelliculés	DE/H/0525/002	0951/11111330	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	LU
Benetor Comp 20 mg/12,5 mg tableter, filmdrasjerte	DE/H/0525/001	05-3515	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	NO
Omesar Plus 20 mg /25 mg film-coated tablets	DE/H/0525/002	MA204/00305	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	MT

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Mesar plus 20 mg/12,5 mg plévele dengtos tabletés	DE/H/0525/001	LT/1/05/0436/038	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	LT
Mesar plus 20 mg/12,5 mg plévele dengtos tabletés	DE/H/0525/001	LT/1/05/0436/033	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	LT
Mesar plus 20 mg/12,5 mg plévele dengtos tabletés	DE/H/0525/001	LT/1/05/0436/035	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	LT
Mesar plus 20 mg/12,5 mg plévele dengtos tabletés	DE/H/0525/001	LT/1/05/0436/034	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	LT
Mesar plus 20 mg/12,5 mg plévele dengtos tabletés	DE/H/0525/001	LT/1/05/0436/039	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	LT
Mesar plus 20 mg/25 mg plévele dengtos tabletés	DE/H/0525/002	LT/1/05/0436/005	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	LT
Mesar plus 20 mg/25 mg plévele dengtos tabletés	DE/H/0525/002	LT/1/05/0436/006	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	LT
Mesar plus 20 mg/25 mg plévele dengtos tabletés	DE/H/0525/002	LT/1/05/0436/040	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	LT
Mesar plus 20 mg/25 mg plévele dengtos tabletés	DE/H/0525/002	LT/1/05/0436/008	MENARINI INTERNATIONAL OPERATIONS	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
			LUXEMBOURG S.A.	
Mesar plus 20 mg/25 mg plévele dengtos tabletés	DE/H/0525/002	LT/1/05/0436/043	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	LT
Revival Plus, 20 mg + 25 mg, tabletki powlekane	DE/H/0525/002	15495	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	PL
Omesar Plus 20 mg /12.5 mg film-coated tablets	DE/H/0525/001	MA204/00304	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	MT
Benetor Comp 20 mg/25 mg tabletter, filmdrasjerte	DE/H/0525/002	05-3516	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	NO
Revival Plus, 20 mg + 12,5 mg, tabletki powlekane	DE/H/0525/001	15494	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	PL
Tenzar Plus 20 mg/12,5 mg filmom obalené tablety	DE/H/0525/001	58/0048/06-S	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	SK
Olsar Plus 20 mg/12,5 mg, comprimidos revestidos por película	DE/H/0525/001	5911987	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	PT
Olsar Plus 20 mg/12,5 mg, comprimidos revestidos por película	DE/H/0525/001	5911888	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	PT
Olsar Plus 20 mg/12,5 mg, comprimidos revestidos por	DE/H/0525/001	5911789	MENARINI INTERNATIONAL	PT



<b>Product Name (in authorisation country)</b>	<b>MRP/DCP Authorisation number</b>	<b>National Authorisation Number</b>	<b>MAH of product in the member state</b>	<b>Member State where product is authorised</b>
película			OPERATIONS LUXEMBOURG S.A.	
Olsar Plus 20 mg/12,5 mg, comprimidos revestidos por película	DE/H/0525/001	5912084	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	PT
Olsar Plus 20 mg/25 mg, comprimidos revestidos por película	DE/H/0525/002	5912480	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	PT
Olsar Plus 20 mg/25 mg, comprimidos revestidos por película	DE/H/0525/002	5912282	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	PT
Olsar Plus 20 mg/25 mg, comprimidos revestidos por película	DE/H/0525/002	5912381	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	PT
BELSAR PLUS 20 mg/12,5 mg comprimés pelliculés	DE/H/0525/001	0951/11111329	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	LU
Olsar Plus 20 mg/25 mg, comprimidos revestidos por película	DE/H/0525/002	5912183	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	PT
Ixia Plus 20 mg /25 mg comprimidos recubiertos con película	DE/H/0525/002	67.749	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	ES
Ixia Plus 20 mg /12,5 mg comprimidos recubiertos con película	DE/H/0525/001	67.748	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	ES
Co-Tensiol 20 mg/25 mg	DE/H/0525/002	H/06/00426/013	MENARINI	SI

<b>Product Name (in authorisation country)</b>	<b>MRP/DCP Authorisation number</b>	<b>National Authorisation Number</b>	<b>MAH of product in the member state</b>	<b>Member State where product is authorised</b>
filmsko obložene tablete			INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	
Co-Tensiol 20 mg/25 mg filmsko obložene tablete	DE/H/0525/002	H/06/00426/012	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	SI
Tenzar Plus 20 mg/25 mg filmom obalené tablety	DE/H/0525/002	58/0049/06-S	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	SK
Co-Tensiol 20 mg/12,5 mg filmsko obložene tablete	DE/H/0525/001	H/06/00426/001	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	SI
Co-Tensiol 20 mg/12,5 mg filmsko obložene tablete	DE/H/0525/001	H/06/00426/002	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	SI
Benetor Comp 40 mg/12,5 mg filmdragerade tabletter	DE/H/0525/003	27137	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	FI
Benetor Comp 40 mg/25 mg filmdragerade tabletter	DE/H/0525/004	27138	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	FI
BELSAR PLUS 40mg/25mg filmomhulde tabletten	DE/H/0525/004	BE368706	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	BE
BELSAR PLUS 40mg/12,5mg filmomhulde tabletten	DE/H/0525/003	BE368697	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	BE

<b>Product Name (in authorisation country)</b>	<b>MRP/DCP Authorisation number</b>	<b>National Authorisation Number</b>	<b>MAH of product in the member state</b>	<b>Member State where product is authorised</b>
BELSAR PLUS 40 mg/12,5 mg Filmtabletten	DE/H/0525/003	BE368697	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	BE
BELSAR PLUS 40 mg/25 mg Filmtabletten	DE/H/0525/004	BE368706	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	BE
Co-Tensiol 40 mg/25 mg filmsko obložene tablete	DE/H/0525/004	H/06/00426/036	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	SI
Co-Tensiol 40 mg/25 mg filmsko obložene tablete	DE/H/0525/004	H/06/00426/042	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	SI
Co-Tensiol 40 mg/25 mg filmsko obložene tablete	DE/H/0525/004	H/06/00426/043	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	SI
Co-Tensiol 40 mg/25 mg filmsko obložene tablete	DE/H/0525/004	H/06/00426/037	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	SI
Co-Tensiol 40 mg/25 mg filmsko obložene tablete	DE/H/0525/004	H/06/00426/045	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	SI
Co-Tensiol 40 mg/25 mg filmsko obložene tablete	DE/H/0525/004	H/06/00426/046	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	SI
Co-Tensiol 40 mg/25 mg filmsko obložene tablete	DE/H/0525/004	H/06/00426/041	MENARINI INTERNATIONAL OPERATIONS	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
			LUXEMBOURG S.A.	
Co-Tensiol 40 mg/25 mg filmsko obložene tablete	DE/H/0525/004	H/06/00426/038	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	SI
Co-Tensiol 40 mg/25 mg filmsko obložene tablete	DE/H/0525/004	H/06/00426/040	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	SI
Co-Tensiol 40 mg/25 mg filmsko obložene tablete	DE/H/0525/004	H/06/00426/044	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	SI
Co-Tensiol 40 mg/25 mg filmsko obložene tablete	DE/H/0525/004	H/06/00426/039	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	SI
Co-Tensiol 40 mg/12,5 mg filmsko obložene tablete	DE/H/0525/003	H/06/00426/026	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	SI
Co-Tensiol 40 mg/12,5 mg filmsko obložene tablete	DE/H/0525/003	H/06/00426/025	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	SI
Co-Tensiol 40 mg/12,5 mg filmsko obložene tablete	DE/H/0525/003	H/06/00426/024	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	SI
Co-Tensiol 40 mg/12,5 mg filmsko obložene tablete	DE/H/0525/003	H/06/00426/033	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	SI
Co-Tensiol 40 mg/12,5 mg filmsko obložene tablete	DE/H/0525/003	H/06/00426/034	MENARINI INTERNATIONAL	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
			OPERATIONS LUXEMBOURG S.A.	
Co-Tensiol 40 mg/12,5 mg filmsko obložene tablete	DE/H/0525/003	H/06/00426/030	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	SI
Co-Tensiol 40 mg/12,5 mg filmsko obložene tablete	DE/H/0525/003	H/06/00426/027	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	SI
Co-Tensiol 40 mg/12,5 mg filmsko obložene tablete	DE/H/0525/003	H/06/00426/028	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	SI
Co-Tensiol 40 mg/12,5 mg filmsko obložene tablete	DE/H/0525/003	H/06/00426/031	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	SI
Co-Tensiol 40 mg/12,5 mg filmsko obložene tablete	DE/H/0525/003	H/06/00426/029	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	SI
Co-Tensiol 40 mg/12,5 mg filmsko obložene tablete	DE/H/0525/003	H/06/00426/032	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	SI
Mencord Plus 40 mg/12,5 mg Filmtabletten	DE/H/0525/003	1-29039	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	AT
Mencord Plus 40 mg/25 mg Filmtabletten	DE/H/0525/004	1-29040	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	AT
BELSAR PLUS 40 mg/25 mg	DE/H/0525/004	BE368706	MENARINI	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
comprimés pelliculés			INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	
Benetor Comp 40 mg/25 mg kalvopäällysteiset tabletit	DE/H/0525/004	27138	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	FI
Benetor Comp 40 mg/12,5 mg kalvopäällysteiset tabletit	DE/H/0525/003	27137	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	FI
Mesar plus, 40 mg/25 mg õhukese polümeerikattega tabletid	DE/H/0525/004	670810	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	EE
Benetor Comp, filmovertrukne tabletter	DE/H/0525/003	44540	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	DK
Mesar plus, 40 mg/12,5 mg õhukese polümeerikattega tabletid	DE/H/0525/003	670910	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	EE
Benetor Comp, filmovertrukne tabletter	DE/H/0525/004	44541	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	DK
BELSAR PLUS 40 mg/12,5 mg comprimés pelliculés	DE/H/0525/003	BE368697	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	BE
Sarten Plus H 40 mg/25 mg, potahované tablety	DE/H/0525/004	58/884/10-C	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	CZ

<b>Product Name (in authorisation country)</b>	<b>MRP/DCP Authorisation number</b>	<b>National Authorisation Number</b>	<b>MAH of product in the member state</b>	<b>Member State where product is authorised</b>
Sarten Plus H 40 mg/12,5 mg, potahované tablety	DE/H/0525/003	58/883/10-C	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	CZ
Laresin Plus 40 mg/12,5 mg filmtabletta	DE/H/0525/003	OGYI-T-20333/05	BERLIN-CHEMIE AG	HU
Laresin Plus 40 mg/12,5 mg filmtabletta	DE/H/0525/003	OGYI-T-20333/06	BERLIN-CHEMIE AG	HU
Laresin Plus 40 mg/25 mg filmtabletta	DE/H/0525/004	OGYI-T-20333/08	BERLIN-CHEMIE AG	HU
Laresin Plus 40 mg/25 mg filmtabletta	DE/H/0525/004	OGYI-T-20333/07	BERLIN-CHEMIE AG	HU
Benetor Comp 40 mg/25 mg filmhúðaðar töflur	DE/H/0525/004	IS/1/09/051/02	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IS
Benetor Comp 40 mg/12,5 mg filmhúðaðar töflur	DE/H/0525/003	IS/1/09/051/01	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IS
ALTEISDUO 40 mg/12,5 mg, comprimé pelliculé	DE/H/0525/003	34009 576 792 4 8	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	FR
ALTEISDUO 40 mg/12,5 mg, comprimé pelliculé	DE/H/0525/003	34009 350 238 6 5	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	FR
ALTEISDUO 40 mg/25 mg, comprimé pelliculé	DE/H/0525/004	34009 576 793 0 9	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	FR
ALTEISDUO 40 mg/25 mg, comprimé pelliculé	DE/H/0525/004	34009 350 242 3 7	MENARINI INTERNATIONAL OPERATIONS	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
			LUXEMBOURG S.A.	
ALTEISDUO 40 mg/12,5 mg, comprimé pelliculé	DE/H/0525/003	34009 350 239 2 6	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	FR
Votum® plus 40 mg/25 mg Filmtabletten	DE/H/0525/004	58597.03.00	BERLIN-CHEMIE AG	DE
Votum® plus 40 mg/12,5 mg Filmtabletten	DE/H/0525/003	58597.02.00	BERLIN-CHEMIE AG	DE
ALTEISDUO 40 mg/25 mg, comprimé pelliculé	DE/H/0525/004	34009 350 244 6 6	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	FR
Olartan-Plus® 40 mg/25 mg επικαλυμμένο με λεπτό υμένιο δισκίο	DE/H/0525/004	4747	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	GR
Olartan-Plus® 40 mg/12,5 mg επικαλυμμένο με λεπτό υμένιο δισκίο	DE/H/0525/003	4746	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	GR
Mesar Plus 40 mg/12,5 mg apvalkotās tabletes	DE/H/0525/003	10-0145	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	LV
Mesar Plus 40 mg/25 mg apvalkotās tabletes	DE/H/0525/004	10-0146	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	LV
OLPREZIDE 40 mg/12,5 mg compresse rivestite con film	DE/H/0525/003	037109319	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
OLPREZIDE 40 mg/12,5 mg compresse rivestite con film	DE/H/0525/003	037109295	MENARINI INTERNATIONAL	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
			OPERATIONS LUXEMBOURG S.A.	
OLPREZIDE 40 mg/25 mg compresse rivestite con film	DE/H/0525/004	037109422	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
OLPREZIDE 40 mg/25 mg compresse rivestite con film	DE/H/0525/004	037109372	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
OLPREZIDE 40 mg/25 mg compresse rivestite con film	DE/H/0525/004	037109358	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
OLPREZIDE 40 mg/12,5 mg compresse rivestite con film	DE/H/0525/003	037109232	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
OLPREZIDE 40 mg/12,5 mg compresse rivestite con film	DE/H/0525/003	037109321	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
OLPREZIDE 40 mg/12,5 mg compresse rivestite con film	DE/H/0525/003	037109271	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
OLPREZIDE 40 mg/25 mg compresse rivestite con film	DE/H/0525/004	037109446	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
OLPREZIDE 40 mg/25 mg compresse rivestite con film	DE/H/0525/004	037109461	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
OLPREZIDE 40 mg/25 mg	DE/H/0525/004	037109360	MENARINI	IT

<b>Product Name (in authorisation country)</b>	<b>MRP/DCP Authorisation number</b>	<b>National Authorisation Number</b>	<b>MAH of product in the member state</b>	<b>Member State where product is authorised</b>
compresse rivestite con film			INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	
OLPREZIDE 40 mg/25 mg compresse rivestite con film	DE/H/0525/004	037109459	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
OLPREZIDE 40 mg/25 mg compresse rivestite con film	DE/H/0525/004	037109408	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
OLPREZIDE 40 mg/25 mg compresse rivestite con film	DE/H/0525/004	037109396	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
OLPREZIDE 40 mg/25 mg compresse rivestite con film	DE/H/0525/004	037109384	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
OLPREZIDE 40 mg/25 mg compresse rivestite con film	DE/H/0525/004	037109434	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
OLPREZIDE 40 mg/25 mg compresse rivestite con film	DE/H/0525/004	037109410	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
Omesar Plus 40 mg/25 mg film-coated tablets	DE/H/0525/004	PA 865/14/4	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IE
Omesar Plus 40 mg/12.5 mg film-coated tablets	DE/H/0525/003	PA 865/14/3	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IE

<b>Product Name (in authorisation country)</b>	<b>MRP/DCP Authorisation number</b>	<b>National Authorisation Number</b>	<b>MAH of product in the member state</b>	<b>Member State where product is authorised</b>
OLPREZIDE 40 mg/12,5 mg compresse rivestite con film	DE/H/0525/003	037109307	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
OLPREZIDE 40 mg/12,5 mg compresse rivestite con film	DE/H/0525/003	037109345	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
OLPREZIDE 40 mg/12,5 mg compresse rivestite con film	DE/H/0525/003	037109244	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
OLPREZIDE 40 mg/12,5 mg compresse rivestite con film	DE/H/0525/003	037109257	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
OLPREZIDE 40 mg/12,5 mg compresse rivestite con film	DE/H/0525/003	037109333	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
OLPREZIDE 40 mg/12,5 mg compresse rivestite con film	DE/H/0525/003	037109269	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
OLPREZIDE 40 mg/12,5 mg compresse rivestite con film	DE/H/0525/003	037109283	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
Mesar plus 40 mg/12,5 mg plévele dengtos tabletés	DE/H/0525/003	LT/1/05/0436/019	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	LT
Mesar plus 40 mg/12,5 mg plévele dengtos tabletés	DE/H/0525/003	LT/1/05/0436/018	MENARINI INTERNATIONAL OPERATIONS	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
			LUXEMBOURG S.A.	
Mesar plus 40 mg/12,5 mg plévele dengtos tabletés	DE/H/0525/003	LT/1/05/0436/016	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	LT
Mesar plus 40 mg/12,5 mg plévele dengtos tabletés	DE/H/0525/003	LT/1/05/0436/009	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	LT
Mesar plus 40 mg/12,5 mg plévele dengtos tabletés	DE/H/0525/003	LT/1/05/0436/017	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	LT
Mesar plus 40 mg/12,5 mg plévele dengtos tabletés	DE/H/0525/003	LT/1/05/0436/015	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	LT
Mesar plus 40 mg/12,5 mg plévele dengtos tabletés	DE/H/0525/003	LT/1/05/0436/010	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	LT
BELSAR PLUS 40 mg/12,5 mg comprimés pelliculés	DE/H/0525/003	0951/10080007	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	LU
Mesar plus 40 mg/25 mg plévele dengtos tabletés	DE/H/0525/004	LT/1/05/0436/029	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	LT
Mesar plus 40 mg/25 mg plévele dengtos tabletés	DE/H/0525/004	LT/1/05/0436/030	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	LT
Mesar plus 40 mg/25 mg plévele dengtos tabletés	DE/H/0525/004	LT/1/05/0436/028	MENARINI INTERNATIONAL	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
			OPERATIONS LUXEMBOURG S.A.	
Mesar plus 40 mg/12,5 mg plêvele dengtos tabletês	DE/H/0525/003	LT/1/05/0436/020	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	LT
Mesar plus 40 mg/12,5 mg plêvele dengtos tabletês	DE/H/0525/003	LT/1/05/0436/013	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	LT
Mesar plus 40 mg/12,5 mg plêvele dengtos tabletês	DE/H/0525/003	LT/1/05/0436/012	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	LT
Omesar Plus 40 mg/25 mg film-coated tablets	DE/H/0525/004	MA204/00307	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	MT
BELSAR PLUS 40 mg/25 mg comprimês pelliculês	DE/H/0525/004	0951/10080008	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	LU
Mesar plus 40 mg/12,5 mg plêvele dengtos tabletês	DE/H/0525/003	LT/1/05/0436/014	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	LT
Mesar plus 40 mg/12,5 mg plêvele dengtos tabletês	DE/H/0525/003	LT/1/05/0436/011	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	LT
Benetor Comp 40 mg/12,5 mg tableter, filmdrasjerte	DE/H/0525/003	08-6467	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	NO
Mesar plus 40 mg/25 mg	DE/H/0525/004	LT/1/05/0436/021	MENARINI	LT

<b>Product Name (in authorisation country)</b>	<b>MRP/DCP Authorisation number</b>	<b>National Authorisation Number</b>	<b>MAH of product in the member state</b>	<b>Member State where product is authorised</b>
plèvele dengtos tabletès			INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	
Benetor Comp 40 mg/25 mg tableter, filmdrasjerte	DE/H/0525/004	08-6468	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	NO
Omesar Plus 40 mg/12.5 mg film-coated tablets	DE/H/0525/003	MA204/00306	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	MT
Olsar Plus 40 mg + 12,5 mg, comprimidos revestidos por película	DE/H/0525/003	5350160	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	PT
Olsar Plus 40 mg + 12,5 mg, comprimidos revestidos por película	DE/H/0525/003	5350178	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	PT
Olsar Plus 40 mg + 25 mg, comprimidos revestidos por película	DE/H/0525/004	5350210	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	PT
Olsar Plus 40 mg + 25 mg, comprimidos revestidos por película	DE/H/0525/004	5350202	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	PT
Mesar plus 40 mg/25 mg plèvele dengtos tabletès	DE/H/0525/004	LT/1/05/0436/024	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	LT
Mesar plus 40 mg/25 mg plèvele dengtos tabletès	DE/H/0525/004	LT/1/05/0436/025	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	LT

<b>Product Name (in authorisation country)</b>	<b>MRP/DCP Authorisation number</b>	<b>National Authorisation Number</b>	<b>MAH of product in the member state</b>	<b>Member State where product is authorised</b>
Mesar plus 40 mg/25 mg plèvele dengtos tabletės	DE/H/0525/004	LT/1/05/0436/032	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	LT
Mesar plus 40 mg/25 mg plèvele dengtos tabletės	DE/H/0525/004	LT/1/05/0436/027	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	LT
Mesar plus 40 mg/25 mg plèvele dengtos tabletės	DE/H/0525/004	LT/1/05/0436/031	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	LT
Mesar plus 40 mg/25 mg plèvele dengtos tabletės	DE/H/0525/004	LT/1/05/0436/022	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	LT
Mesar plus 40 mg/25 mg plèvele dengtos tabletės	DE/H/0525/004	LT/1/05/0436/026	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	LT
Mesar plus 40 mg/25 mg plèvele dengtos tabletės	DE/H/0525/004	LT/1/05/0436/023	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	LT
Ixia Plus 40 mg /25 mg comprimidos recubiertos con película	DE/H/0525/004	72.698	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	ES
Tenzar Plus 40 mg/12,5 mg filmom obalené tablety	DE/H/0525/003	58/0653/10-S	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	SK
Co-Tensiol 40 mg/25 mg filmsko obložene tablete	DE/H/0525/004	H/06/00426/035	MENARINI INTERNATIONAL OPERATIONS	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
			LUXEMBOURG S.A.	
Ixia Plus 40 mg /12,5 mg comprimidos recubiertos con película	DE/H/0525/003	72.696	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	ES
Co-Tensiol 40 mg/12,5 mg filmsko obložene tablete	DE/H/0525/003	H/06/00426/023	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	SI
Tenzar Plus 40 mg/25 mg filmom obalené tablety	DE/H/0525/004	58/0654/10-S	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	SK
Olartan-Plus® 40 mg/25 mg επικαλυμμένο με λεπτό υμένιο δισκίο	DE/H/0525/004	21786	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	CY
Olartan-Plus® 40 mg/12,5 mg επικαλυμμένο με λεπτό υμένιο δισκί	DE/H/0525/003	21785	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	CY
Olmesartan HCT Daiichi Sankyo 20 mg/12,5 mg Filmtabletten	DE/H/4604/001	96199.00.00	DAIICHI SANKYO EUROPE GMBH	DE
Olmesartan HCT Daiichi Sankyo 20 mg/25 mg Filmtabletten	DE/H/4604/002	96200.00.00	DAIICHI SANKYO EUROPE GMBH	DE
Olmesartan HCT Daiichi Sankyo 40 mg/12,5 mg Filmtabletten	DE/H/4604/003	96201.00.00	DAIICHI SANKYO EUROPE GMBH	DE
Olmesartan HCT Daiichi Sankyo 40 mg/25 mg Filmtabletten	DE/H/4604/004	96202.00.00	DAIICHI SANKYO EUROPE GMBH	DE
OLMETEC PLUS 20 mg/12,5 mg comprimés pelliculés	DE/H/0523/001	BE 282746	DAIICHI SANKYO BELGIUM S.A	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
OLMETEC PLUS 20 mg/25 mg comprimés pelliculés	DE/H/0523/002	BE 282764	DAIICHI SANKYO BELGIUM S.A	BE
Olmotec Plus 20 mg/12,5 mg Filmtabletten	DE/H/0523/001	1-26158	DAIICHI SANKYO AUSTRIA GMBH	AT
Olmotec Plus 20 mg/25 mg Filmtabletten	DE/H/0523/002	1-26159	DAIICHI SANKYO AUSTRIA GMBH	AT
OLMETEC PLUS 20mg/12,5mg filmomhulde tabletten	DE/H/0523/001	BE 282746	DAIICHI SANKYO BELGIUM S.A	BE
OLMETEC PLUS 20mg/25mg filmomhulde tabletten	DE/H/0523/002	BE 282764	DAIICHI SANKYO BELGIUM S.A	BE
Olmotec Plus 20 mg/12,5 mg tabletti, kalvopäällysteinen	DE/H/0523/001	21332	DAIICHI SANKYO EUROPE GMBH	FI
Olmotec Plus 20 mg/25 mg tabletti, kalvopäällysteinen	DE/H/0523/002	21333	DAIICHI SANKYO EUROPE GMBH	FI
Olmotec Plus 20 mg/12,5 mg filmuhúðaðar töflur	DE/H/0523/001	IS/1/05/047/01	DAIICHI SANKYO EUROPE GMBH	IS
Olmotec Plus 20 mg/25 mg filmuhúðaðar töflur	DE/H/0523/002	IS/1/05/047/02	DAIICHI SANKYO EUROPE GMBH	IS
Benetor Plus 20 mg /12.5 mg film-coated tablets	DE/H/0523/001	PA1595/2/1	DAIICHI SANKYO IRELAND LIMITED	IE
Benetor Plus 20 mg /25 mg film-coated tablets	DE/H/0523/002	PA1595/2/2	DAIICHI SANKYO IRELAND LIMITED	IE
OLMEGAN 20 mg/12,5 mg compresse rivestite con film	DE/H/0523/001	037110018	DAIICHI SANKYO ITALIA S.P.A	IT
OLMEGAN 20 mg/12,5 mg compresse rivestite con film	DE/H/0523/001	037110020	DAIICHI SANKYO ITALIA S.P.A	IT
OLMEGAN 20 mg/12,5 mg compresse rivestite con film	DE/H/0523/001	037110032	DAIICHI SANKYO ITALIA S.P.A	IT
OLMEGAN 20 mg/12,5 mg compresse rivestite con film	DE/H/0523/001	037110044	DAIICHI SANKYO ITALIA S.P.A	IT
OLMEGAN 20 mg/12,5 mg compresse rivestite con film	DE/H/0523/001	037110119	DAIICHI SANKYO ITALIA S.P.A	IT
OLMEGAN 20 mg/12,5 mg compresse rivestite con film	DE/H/0523/001	037110057	DAIICHI SANKYO ITALIA S.P.A	IT

<b>Product Name (in authorisation country)</b>	<b>MRP/DCP Authorisation number</b>	<b>National Authorisation Number</b>	<b>MAH of product in the member state</b>	<b>Member State where product is authorised</b>
OLMEGAN 20 mg/12,5 mg compresse rivestite con film	DE/H/0523/001	037110069	DAIICHI SANKYO ITALIA S.P.A	IT
OLMEGAN 20 mg/12,5 mg compresse rivestite con film	DE/H/0523/001	037110071	DAIICHI SANKYO ITALIA S.P.A	IT
OLMEGAN 20 mg/12,5 mg compresse rivestite con film	DE/H/0523/001	037110083	DAIICHI SANKYO ITALIA S.P.A	IT
OLMEGAN 20 mg/12,5 mg compresse rivestite con film	DE/H/0523/001	037110095	DAIICHI SANKYO ITALIA S.P.A	IT
OLMEGAN 20 mg/12,5 mg compresse rivestite con film	DE/H/0523/001	037110107	DAIICHI SANKYO ITALIA S.P.A	IT
OLMEGAN 20 mg/25 mg compresse rivestite con film	DE/H/0523/002	037110121	DAIICHI SANKYO ITALIA S.P.A	IT
OLMEGAN 20 mg/25 mg compresse rivestite con film	DE/H/0523/002	037110133	DAIICHI SANKYO ITALIA S.P.A	IT
OLMEGAN 20 mg/25 mg compresse rivestite con film	DE/H/0523/002	037110145	DAIICHI SANKYO ITALIA S.P.A	IT
OLMEGAN 20 mg/25 mg compresse rivestite con film	DE/H/0523/002	037110158	DAIICHI SANKYO ITALIA S.P.A	IT
OLMEGAN 20 mg/25 mg compresse rivestite con film	DE/H/0523/002	037110160	DAIICHI SANKYO ITALIA S.P.A	IT
OLMEGAN 20 mg/25 mg compresse rivestite con film	DE/H/0523/002	037110172	DAIICHI SANKYO ITALIA S.P.A	IT
OLMEGAN 20 mg/25 mg compresse rivestite con film	DE/H/0523/002	037110184	DAIICHI SANKYO ITALIA S.P.A	IT
OLMEGAN 20 mg/25 mg compresse rivestite con film	DE/H/0523/002	037110196	DAIICHI SANKYO ITALIA S.P.A	IT
OLMEGAN 20 mg/25 mg compresse rivestite con film	DE/H/0523/002	037110208	DAIICHI SANKYO ITALIA S.P.A	IT
OLMEGAN 20 mg/25 mg compresse rivestite con film	DE/H/0523/002	037110210	DAIICHI SANKYO ITALIA S.P.A	IT
OLMEGAN 20 mg/25 mg compresse rivestite con film	DE/H/0523/002	037110222	DAIICHI SANKYO ITALIA S.P.A	IT
Olmetec Comp 20 mg/12,5 mg tableter, filmdrasjerte	DE/H/0523/001	05-3517	DAIICHI SANKYO EUROPE GMBH	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
Olmotec Comp 20 mg/25 mg tabletter, filmdrasjerte	DE/H/0523/002	05-3518	DAIICHI SANKYO EUROPE GMBH	NO
CoOLMETEC 20 mg/25 mg, comprimé pelliculé	DE/H/0523/002	372 213-6	DAIICHI SANKYO FRANCE SAS	FR
CoOLMETEC 20 mg/25 mg, comprimé pelliculé	DE/H/0523/002	372 214-2	DAIICHI SANKYO FRANCE SAS	FR
CoOLMETEC 20 mg/25 mg, comprimé pelliculé	DE/H/0523/002	567 669-9	DAIICHI SANKYO FRANCE SAS	FR
CoOLMETEC 20 mg/12,5 mg, comprimé pelliculé	DE/H/0523/001	372 210-7	DAIICHI SANKYO FRANCE SAS	FR
CoOLMETEC 20 mg/12,5 mg, comprimé pelliculé	DE/H/0523/001	372 211-3	DAIICHI SANKYO FRANCE SAS	FR
CoOLMETEC 20 mg/12,5 mg, comprimé pelliculé	DE/H/0523/001	567 668-2	DAIICHI SANKYO FRANCE SAS	FR
OLMETEC PLUS 20 mg/12,5 mg comprimés pelliculés	DE/H/0523/001	1413/06020005	DAIICHI SANKYO BELGIUM S.A	LU
OLMETEC PLUS 20 mg/25 mg comprimés pelliculés	DE/H/0523/002	1413/06020004	DAIICHI SANKYO BELGIUM S.A	LU
Olmotec Plus 20 mg+12,5 mg comprimidos revestidos por película	DE/H/0523/001	5910989	DAIICHI SANKYO PORTUGAL,LDA	PT
Olmotec Plus 20 mg+12,5 mg comprimidos revestidos por película	DE/H/0523/001	5911086	DAIICHI SANKYO PORTUGAL,LDA	PT
Olmotec Plus 20 mg+12,5 mg comprimidos revestidos por película	DE/H/0523/001	5911185	DAIICHI SANKYO PORTUGAL,LDA	PT
Olmotec Plus 20 mg+12,5 mg comprimidos revestidos por película	DE/H/0523/001	5911284	DAIICHI SANKYO PORTUGAL,LDA	PT
Olmotec Plus 20 mg+25 mg comprimidos revestidos por película	DE/H/0523/002	5911383	DAIICHI SANKYO PORTUGAL,LDA	PT
Olmotec Plus 20 mg+25 mg comprimidos revestidos por película	DE/H/0523/002	5911482	DAIICHI SANKYO PORTUGAL,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
película				
Olmetec Plus 20 mg+25 mg comprimidos revestidos por película	DE/H/0523/002	5911581	DAIICHI SANKYO PORTUGAL,LDA	PT
Olmetec Plus 20 mg+25 mg comprimidos revestidos por película	DE/H/0523/002	5911680	DAIICHI SANKYO PORTUGAL,LDA	PT
Olmetec Plus 20 mg/12,5 mg Filmtabletten	DE/H/0523/001	58589.00.00	DAIICHI SANKYO EUROPE GMBH	DE
Olmetec Plus 20 mg/25 mg Filmtabletten	DE/H/0523/002	58589.01.00	DAIICHI SANKYO EUROPE GMBH	DE
Olmetec Plus 20 mg /12.5 mg film-coated tablets	DE/H/0523/001	PL 08265/0022	DAIICHI SANKYO UK LTD	UK
Olmetec Plus 20 mg /25 mg film-coated tablets	DE/H/0523/002	PL 08265/0023	DAIICHI SANKYO UK LTD	UK
Olmetec Plus 20 mg/12,5 mg comprimidos recubiertos con película	DE/H/0523/001	67.672	DAIICHI SANKYO ESPAÑA, S.A.	ES
Olmetec Plus 20 mg/25 mg comprimidos recubiertos con película	DE/H/0523/002	67.679	DAIICHI SANKYO ESPAÑA, S.A.	ES
Olmetec HCTZ 20/12,5, filmomhulde tabletten 20 mg / 12,5 mg	DE/H/0523/001	RVG 32740	DAIICHI SANKYO NEDERLAND B.V.	NL
Olmetec HCTZ 20/25, filmomhulde tabletten 20 mg / 25 mg	DE/H/0523/002	RVG 32741	DAIICHI SANKYO NEDERLAND B.V.	NL
Olmetec Plus, filmovertrukne tabletter 20 mg/12,5 mg	DE/H/0523/001	38146	DAIICHI SANKYO EUROPE GMBH	DK
Olmetec Plus, filmovertrukne tabletter 20 mg/25 mg	DE/H/0523/002	38147	DAIICHI SANKYO EUROPE GMBH	DK
OLMETEC PLUS® 20 mg/25 mg επικαλυμμένο με λεπτό υμένιο δισκίο	DE/H/0523/002	20163/27-3-2006	PFIZER HELLAS, A.E.	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
OLMETEC PLUS® 20 mg/12,5 mg επικαλυμμένο με λεπτό υμένιο δισκίο	DE/H/0523/001	20162/27-3-2006	PFIZER HELLAS, A.E.	GR
PLAUNAZIDE 40 mg/12,5 mg compresse rivestite con film	DE/H/0524/003	037108230	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
PLAUNAZIDE 40 mg/12,5 mg compresse rivestite con film	DE/H/0524/003	037108255	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
PLAUNAZIDE 40 mg/12,5 mg compresse rivestite con film	DE/H/0524/003	037108305	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
PLAUNAZIDE 40 mg/25 mg compresse rivestite con film	DE/H/0524/004	037108382	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
PLAUNAZIDE 40 mg/25 mg compresse rivestite con film	DE/H/0524/004	037108457	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
PLAUNAZIDE 40 mg/12,5 mg compresse rivestite con film	DE/H/0524/003	037108343	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
PLAUNAZIDE 40 mg/12,5 mg compresse rivestite con film	DE/H/0524/003	037108331	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
PLAUNAZIDE 40 mg/25 mg compresse rivestite con film	DE/H/0524/004	037108394	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT

<b>Product Name (in authorisation country)</b>	<b>MRP/DCP Authorisation number</b>	<b>National Authorisation Number</b>	<b>MAH of product in the member state</b>	<b>Member State where product is authorised</b>
PLAUNAZIDE 40 mg/25 mg compresse rivestite con film	DE/H/0524/004	037108406	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
PLAUNAZIDE 40 mg/12,5 mg compresse rivestite con film	DE/H/0524/003	037108281	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
PLAUNAZIDE 40 mg/12,5 mg compresse rivestite con film	DE/H/0524/003	037108293	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
PLAUNAZIDE 40 mg/25 mg compresse rivestite con film	DE/H/0524/004	037108356	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
PLAUNAZIDE 40 mg/25 mg compresse rivestite con film	DE/H/0524/004	037108420	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
PLAUNAZIDE 40 mg/25 mg compresse rivestite con film	DE/H/0524/004	037108370	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
PLAUNAZIDE 40 mg/25 mg compresse rivestite con film	DE/H/0524/004	037108368	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
PLAUNAZIDE 40 mg/25 mg compresse rivestite con film	DE/H/0524/004	037108469	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
PLAUNAZIDE 40 mg/25 mg compresse rivestite con film	DE/H/0524/004	037108432	MENARINI INTERNATIONAL OPERATIONS	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
			LUXEMBOURG S.A.	
PLAUNAZIDE 40 mg/12,5 mg compresse rivestite con film	DE/H/0524/003	037108329	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
PLAUNAZIDE 40 mg/25 mg compresse rivestite con film	DE/H/0524/004	037108444	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
PLAUNAZIDE 40 mg/25 mg compresse rivestite con film	DE/H/0524/004	037108418	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
PLAUNAZIDE 40 mg/12,5 mg compresse rivestite con film	DE/H/0524/003	037108242	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
PLAUNAZIDE 40 mg/12,5 mg compresse rivestite con film	DE/H/0524/003	037108317	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
PLAUNAZIDE 40 mg/12,5 mg compresse rivestite con film	DE/H/0524/003	037108279	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
PLAUNAZIDE 40 mg/12,5 mg compresse rivestite con film	DE/H/0524/003	037108267	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
Olmes Plus 40 mg/12,5 mg Filmtabletten	DE/H/0524/003	58593.02.00	DAIICHI SANKYO EUROPE GMBH	DE
Olmes Plus 40 mg/25 mg Filmtabletten	DE/H/0524/004	58593.03.00	DAIICHI SANKYO EUROPE GMBH	DE
PLAUNAZIDE 40 mg/12,5 mg compresse rivestite con film	DE/H/0524/003	037108230	MENARINI INTERNATIONAL	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
			OPERATIONS LUXEMBOURG S.A.	
PLAUNAZIDE 40 mg/12,5 mg compresse rivestite con film	DE/H/0524/003	037108255	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
PLAUNAZIDE 40 mg/12,5 mg compresse rivestite con film	DE/H/0524/003	037108305	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
PLAUNAZIDE 40 mg/25 mg compresse rivestite con film	DE/H/0524/004	037108382	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
PLAUNAZIDE 40 mg/25 mg compresse rivestite con film	DE/H/0524/004	037108457	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
PLAUNAZIDE 40 mg/12,5 mg compresse rivestite con film	DE/H/0524/003	037108343	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
PLAUNAZIDE 40 mg/12,5 mg compresse rivestite con film	DE/H/0524/003	037108331	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
PLAUNAZIDE 40 mg/25 mg compresse rivestite con film	DE/H/0524/004	037108394	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
PLAUNAZIDE 40 mg/25 mg compresse rivestite con film	DE/H/0524/004	037108406	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
PLAUNAZIDE 40 mg/12,5 mg	DE/H/0524/003	037108281	MENARINI	IT



<b>Product Name (in authorisation country)</b>	<b>MRP/DCP Authorisation number</b>	<b>National Authorisation Number</b>	<b>MAH of product in the member state</b>	<b>Member State where product is authorised</b>
compresse rivestite con film			INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	
PLAUNAZIDE 40 mg/12,5 mg compresse rivestite con film	DE/H/0524/003	037108293	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
PLAUNAZIDE 40 mg/25 mg compresse rivestite con film	DE/H/0524/004	037108356	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
PLAUNAZIDE 40 mg/25 mg compresse rivestite con film	DE/H/0524/004	037108420	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
PLAUNAZIDE 40 mg/25 mg compresse rivestite con film	DE/H/0524/004	037108370	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
PLAUNAZIDE 40 mg/25 mg compresse rivestite con film	DE/H/0524/004	037108368	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
PLAUNAZIDE 40 mg/25 mg compresse rivestite con film	DE/H/0524/004	037108469	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
PLAUNAZIDE 40 mg/25 mg compresse rivestite con film	DE/H/0524/004	037108432	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
PLAUNAZIDE 40 mg/12,5 mg compresse rivestite con film	DE/H/0524/003	037108329	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT

<b>Product Name (in authorisation country)</b>	<b>MRP/DCP Authorisation number</b>	<b>National Authorisation Number</b>	<b>MAH of product in the member state</b>	<b>Member State where product is authorised</b>
PLAUNAZIDE 40 mg/25 mg compresse rivestite con film	DE/H/0524/004	037108444	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
PLAUNAZIDE 40 mg/25 mg compresse rivestite con film	DE/H/0524/004	037108418	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
PLAUNAZIDE 40 mg/12,5 mg compresse rivestite con film	DE/H/0524/003	037108242	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
PLAUNAZIDE 40 mg/12,5 mg compresse rivestite con film	DE/H/0524/003	037108317	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
PLAUNAZIDE 40 mg/12,5 mg compresse rivestite con film	DE/H/0524/003	037108279	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
PLAUNAZIDE 40 mg/12,5 mg compresse rivestite con film	DE/H/0524/003	037108267	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
Openvas Plus 40 mg /25 mg comprimidos recubiertos con película	DE/H/0524/004	72694	PFIZER, S.L.	ES
Openvas Plus 40 mg /12,5 mg comprimidos recubiertos con película	DE/H/0524/003	72693	PFIZER, S.L.	ES
Benetor Comp 20 mg/12,5 mg filmdragerade tabletter	DE/H/0525/001	21334	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	FI
BELSAR Plus 20mg/12,5mg	DE/H/0525/001	BE282782	MENARINI	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
filmomhulde tabletten			INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	
Belsar Plus 20 mg/12,5 mg Filmtabletten	DE/H/0525/001	BE282782	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	BE
Olartan-Plus® 20 mg/12,5 mg επικαλυμμένο με λεπτό υμένιο δισκία	DE/H/0525/001	4744	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	GR
Co-Tensiol 20 mg/12,5 mg filmsko obložene tablete	DE/H/0525/001	H/06/00426/010	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	SI
Co-Tensiol 20 mg/12,5 mg filmsko obložene tablete	DE/H/0525/001	H/06/00426/009	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	SI
Co-Tensiol 20 mg/12,5 mg filmsko obložene tablete	DE/H/0525/001	H/06/00426/003	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	SI
Co-Tensiol 20 mg/12,5 mg filmsko obložene tablete	DE/H/0525/001	H/06/00426/006	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	SI
Co-Tensiol 20 mg/12,5 mg filmsko obložene tablete	DE/H/0525/001	H/06/00426/007	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	SI
Co-Tensiol 20 mg/12,5 mg filmsko obložene tablete	DE/H/0525/001	H/06/00426/005	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	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-Tensiol 20 mg/12,5 mg filmsko obložene tablete	DE/H/0525/001	H/06/00426/011	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	SI
Co-Tensiol 20 mg/12,5 mg filmsko obložene tablete	DE/H/0525/001	H/06/00426/004	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	SI
Co-Tensiol 20 mg/12,5 mg filmsko obložene tablete	DE/H/0525/001	H/06/00426/008	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	SI
Olmes Plus 20 mg/12,5 mg Filmtabletten	DE/H/0524/001	58593.00.00	DAIICHI SANKYO EUROPE GMBH	DE
Olmes Plus 20 mg/25 mg Filmtabletten	DE/H/0524/002	58593.01.00	DAIICHI SANKYO EUROPE GMBH	DE
Mencord® Plus 20 mg/12,5 mg Filmtabletten	DE/H/0525/001	1-26160	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	AT
BELSAR PLUS 20 mg/12,5 mg comprimés pelliculés	DE/H/0525/001	BE282782	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	BE
Mesar plus, 20 mg/12,5 mg õhukese polümeerikattega tabletid	DE/H/0525/001	501105	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	EE
Benetor Comp, filmovertrukne tableter	DE/H/0525/001	38151	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	DK
Olartan-Plus® 20 mg/12,5 mg επικαλυμμένο με λεπτό υμένιο δισκί	DE/H/0525/001	20333	MENARINI INTERNATIONAL OPERATIONS	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
			LUXEMBOURG S.A.	
Sarten Plus H 20/12,5 mg potahované tablety	DE/H/0525/001	58/458/05-C	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	CZ
Benetor Comp 20 mg/12,5 mg kalvopäällysteiset tabletit	DE/H/0525/001	21334	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	FI
Laresin Plus 20 mg/12,5 mg filmtabletta	DE/H/0525/001	OGYI-T-20333/02	BERLIN-CHEMIE AG	HU
Laresin Plus 20 mg/12,5 mg filmtabletta	DE/H/0525/001	OGYI-T-20333/01	BERLIN-CHEMIE AG	HU
Benetor Comp 20 mg/12,5 mg filmuhúðaðar töflur	DE/H/0525/001	IS/1/05/048/01	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IS
ALTEISDUO 20 mg/12,5 mg, comprimé pelliculé	DE/H/0525/001	34009 372 219 4 8	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	FR
Votum® plus 20 mg/12,5 mg Filmtabletten	DE/H/0525/001	58597.00.00	BERLIN-CHEMIE AG	DE
ALTEISDUO 20 mg/12,5 mg, comprimé pelliculé	DE/H/0525/001	34009 372 220 2 0	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	FR
ALTEISDUO 20 mg/12,5 mg, comprimé pelliculé	DE/H/0525/001	34009 567 673 6 6	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	FR
Mesar Plus 20 mg/12,5 mg apvalkotās tabletes	DE/H/0525/001	05-0631	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	LV

<b>Product Name (in authorisation country)</b>	<b>MRP/DCP Authorisation number</b>	<b>National Authorisation Number</b>	<b>MAH of product in the member state</b>	<b>Member State where product is authorised</b>
PLAUNAZIDE 20 mg/25 mg compresse rivestite con film	DE/H/0524/002	037108204	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
PLAUNAZIDE 20 mg/25 mg compresse rivestite con film	DE/H/0524/002	037108127	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
OLPREZIDE 20 mg/12,5 mg compresse rivestite con film	DE/H/0525/001	037109081	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
OLPREZIDE 20 mg/12,5 mg compresse rivestite con film	DE/H/0525/001	037109030	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
OLPREZIDE 20 mg/12,5 mg compresse rivestite con film	DE/H/0525/001	037109117	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
PLAUNAZIDE 20 mg/12,5 mg compresse rivestite con film	DE/H/0524/001	037108103	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
PLAUNAZIDE 20 mg/12,5 mg compresse rivestite con film	DE/H/0524/001	037108014	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
PLAUNAZIDE 20 mg/12,5 mg compresse rivestite con film	DE/H/0524/001	037108091	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
PLAUNAZIDE 20 mg/12,5 mg compresse rivestite con film	DE/H/0524/001	037108038	MENARINI INTERNATIONAL OPERATIONS	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
			LUXEMBOURG S.A.	
PLAUNAZIDE 20 mg/12,5 mg compresse rivestite con film	DE/H/0524/001	037108115	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
PLAUNAZIDE 20 mg/12,5 mg compresse rivestite con film	DE/H/0524/001	037108077	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
PLAUNAZIDE 20 mg/25 mg compresse rivestite con film	DE/H/0524/002	037108141	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
PLAUNAZIDE 20 mg/25 mg compresse rivestite con film	DE/H/0524/002	037108216	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
PLAUNAZIDE 20 mg/25 mg compresse rivestite con film	DE/H/0524/002	037108139	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
PLAUNAZIDE 20 mg/25 mg compresse rivestite con film	DE/H/0524/002	037108166	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
PLAUNAZIDE 20 mg/25 mg compresse rivestite con film	DE/H/0524/002	037108178	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
PLAUNAZIDE 20 mg/25 mg compresse rivestite con film	DE/H/0524/002	037108228	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
PLAUNAZIDE 20 mg/25 mg compresse rivestite con film	DE/H/0524/002	037108154	MENARINI INTERNATIONAL	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
			OPERATIONS LUXEMBOURG S.A.	
PLAUNAZIDE 20 mg/25 mg compresse rivestite con film	DE/H/0524/002	037108180	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
OLPREZIDE 20 mg/12,5 mg compresse rivestite con film	DE/H/0525/001	037109055	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
OLPREZIDE 20 mg/12,5 mg compresse rivestite con film	DE/H/0525/001	037109016	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
Omesar Plus 20 mg /12.5 mg film-coated tablets	DE/H/0525/001	PA 865/14/1	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IE
PLAUNAZIDE 20 mg/12,5 mg compresse rivestite con film	DE/H/0524/001	037108065	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
PLAUNAZIDE 20 mg/25 mg compresse rivestite con film	DE/H/0524/002	037108192	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
PLAUNAZIDE 20 mg/12,5 mg compresse rivestite con film	DE/H/0524/001	037108026	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
PLAUNAZIDE 20 mg/12,5 mg compresse rivestite con film	DE/H/0524/001	037108089	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
PLAUNAZIDE 20 mg/12,5 mg	DE/H/0524/001	037108040	MENARINI	IT



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comprese rivestite con film			INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	
OLPREZIDE 20 mg/12,5 mg compresse rivestite con film	DE/H/0525/001	037109079	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
OLPREZIDE 20 mg/12,5 mg compresse rivestite con film	DE/H/0525/001	037109042	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
OLPREZIDE 20 mg/12,5 mg compresse rivestite con film	DE/H/0525/001	037109028	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
OLPREZIDE 20 mg/12,5 mg compresse rivestite con film	DE/H/0525/001	037109093	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
OLPREZIDE 20 mg/12,5 mg compresse rivestite con film	DE/H/0525/001	037109105	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
OLPREZIDE 20 mg/12,5 mg compresse rivestite con film	DE/H/0525/001	037109067	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
PLAUNAZIDE 20 mg/12,5 mg compresse rivestite con film	DE/H/0524/001	037108053	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	IT
Mesar plus 20 mg/12,5 mg plévele dengtos tabletés	DE/H/0525/001	LT/1/05/0436/037	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	LT

<b>Product Name (in authorisation country)</b>	<b>MRP/DCP Authorisation number</b>	<b>National Authorisation Number</b>	<b>MAH of product in the member state</b>	<b>Member State where product is authorised</b>
Mesar plus 20 mg/12,5 mg plèvele dengtos tabletės	DE/H/0525/001	LT/1/05/0436/002	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	LT
Mesar plus 20 mg/12,5 mg plèvele dengtos tabletės	DE/H/0525/001	LT/1/05/0436/001	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	LT
Mesar plus 20 mg/12,5 mg plèvele dengtos tabletės	DE/H/0525/001	LT/1/05/0436/036	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	LT
Mesar plus 20 mg/12,5 mg plèvele dengtos tabletės	DE/H/0525/001	LT/1/05/0436/004	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	LT
Mesar plus 20 mg/12,5 mg plèvele dengtos tabletės	DE/H/0525/001	LT/1/05/0436/003	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	LT
Benetor Comp 20 mg/12,5 mg tableter, filmdrasjerte	DE/H/0525/001	05-3515	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	NO
Mesar plus 20 mg/12,5 mg plèvele dengtos tabletės	DE/H/0525/001	LT/1/05/0436/038	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	LT
Mesar plus 20 mg/12,5 mg plèvele dengtos tabletės	DE/H/0525/001	LT/1/05/0436/033	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	LT
Mesar plus 20 mg/12,5 mg plèvele dengtos tabletės	DE/H/0525/001	LT/1/05/0436/035	MENARINI INTERNATIONAL OPERATIONS	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
			LUXEMBOURG S.A.	
Mesar plus 20 mg/12,5 mg plévele dengtos tabletés	DE/H/0525/001	LT/1/05/0436/034	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	LT
Mesar plus 20 mg/12,5 mg plévele dengtos tabletés	DE/H/0525/001	LT/1/05/0436/039	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	LT
Omesar Plus 20 mg /12.5 mg film-coated tablets	DE/H/0525/001	MA204/00304	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	MT
Revival Plus, 20 mg + 12,5 mg, tabletki powlekane	DE/H/0525/001	15494	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	PL
Tenzar Plus 20 mg/12,5 mg filmom obalené tablety	DE/H/0525/001	58/0048/06-S	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	SK
Olsar Plus 20 mg/12,5 mg, comprimidos revestidos por película	DE/H/0525/001	5911987	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	PT
Olsar Plus 20 mg/12,5 mg, comprimidos revestidos por película	DE/H/0525/001	5911888	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	PT
Olsar Plus 20 mg/12,5 mg, comprimidos revestidos por película	DE/H/0525/001	5911789	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	PT
Olsar Plus 20 mg/12,5 mg, comprimidos revestidos por	DE/H/0525/001	5912084	MENARINI INTERNATIONAL	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
película			OPERATIONS LUXEMBOURG S.A.	
BELSAR PLUS 20 mg/12,5 mg comprimés pelliculés	DE/H/0525/001	0951/11111329	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	LU
Ixia Plus 20 mg /12,5 mg comprimidos recubiertos con película	DE/H/0525/001	67.748	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	ES
Co-Tensiol 20 mg/12,5 mg filmsko obložene tablete	DE/H/0525/001	H/06/00426/001	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	SI
Co-Tensiol 20 mg/12,5 mg filmsko obložene tablete	DE/H/0525/001	H/06/00426/002	MENARINI INTERNATIONAL OPERATIONS LUXEMBOURG S.A.	SI
Openvas Plus 20 mg /25 mg comprimidos recubiertos con película	DE/H/0524/002	67.678	PFIZER, S.L.	ES
Openvas Plus 20 mg /12,5 mg comprimidos recubiertos con película	DE/H/0524/001	67.680	PFIZER, S.L.	ES
Olmesartan medoxomil/Hydrochlorothiazide 20 mg/12.5 mg Film-coated Tablets	PT/H/1155/001	PA 0577/175/001	GENERICS [UK] LIMITED	IE
Olmesartan medoxomil/Hydrochlorothiazide 20 mg/25 mg Film-coated Tablets	PT/H/1155/002	PA 0577/175/002	MCDERMOTT LABORATORIES LTD	IE
Olmesartan medoxomil/Hydrochlorothiazide 40 mg/12.5 mg Film-coated	PT/H/1155/003	PA 0577/175/003	MCDERMOTT LABORATORIES LTD	IE

Product Name (in authorisation country)	MRP/DCP Authorisation number	National Authorisation Number	MAH of product in the member state	Member State where product is authorised
Tablets				
Olmesartan medoxomil/Hydrochlorothiazide 40 mg/25 mg Film-coated Tablets	PT/H/1155/004	PA 0577/175/004	MCDERMOTT LABORATORIES LTD	IE
Olmesartan medoxomilo + Hidroclorotiazida Mylan 20 mg + 12,5 mg comprimidos revestidos por película	PT/H/1155/001	5645155	MYLAN, LDA	PT
Olmesartan medoxomilo + Hidroclorotiazida Mylan 20 mg + 12,5 mg comprimidos revestidos por película	PT/H/1155/001	5645163	MYLAN, LDA	PT
Olmesartan medoxomilo + Hidroclorotiazida Mylan 20 mg + 25 mg comprimidos revestidos por película	PT/H/1155/002	5645205	MYLAN, LDA	PT
Olmesartan medoxomilo + Hidroclorotiazida Mylan 20 mg + 25 mg comprimidos revestidos por película	PT/H/1155/002	5645213	MYLAN, LDA	PT
Olmesartan medoxomilo + Hidroclorotiazida Mylan 40 mg + 12,5 mg comprimidos revestidos por película	PT/H/1155/003	5645221	MYLAN, LDA	PT
Olmesartan medoxomilo + Hidroclorotiazida Mylan 40 mg + 12,5 mg comprimidos revestidos por película	PT/H/1155/003	5645239	MYLAN, LDA	PT
Olmesartan medoxomilo + Hidroclorotiazida Mylan 40 mg + 25 mg comprimidos revestidos por película	PT/H/1155/004	5645247	MYLAN, 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
Olmesartan medoxomilo + Hidroclorotiazida Mylan 40 mg + 25 mg comprimidos revestidos por película	PT/H/1155/004	5645254	MYLAN, LDA	PT
Olmesartan/Hidroclorotiazida Mylan 20 mg/12,5 mg comprimidos recubiertos con película EFG	PT/H/1155/001	80358	MYLAN PHARMACEUTICALS S.L.	ES
Olmesartan/Hidroclorotiazida Mylan 20 mg/25 mg comprimidos recubiertos con película EFG	PT/H/1155/002	80359	MYLAN PHARMACEUTICALS S.L.	ES
Olmesartan + HCTZ / Mylan 20 mg/12,5 mg Επικαλυμμένα με λεπτό υμένιο δισκία	PT/H/1155/001	89249/16-12-2015	GENERICS PHARMA HELLAS LTD	GR
Olmesartan + HCTZ / Mylan 20 mg/25 mg Επικαλυμμένα με λεπτό υμένιο δισκία	PT/H/1155/002	89250/16-12-2015	GENERICS PHARMA HELLAS LTD	GR
Olmesartan + HCTZ / Mylan 40 mg/12,5 mg Επικαλυμμένα με λεπτό υμένιο δισκία	PT/H/1155/003	89251/16-12-2015	GENERICS PHARMA HELLAS LTD	GR
Olmesartan + HCTZ / Mylan 40 mg/25 mg Επικαλυμμένα με λεπτό υμένιο δισκία	PT/H/1155/004	89252/16-12-2015	GENERICS PHARMA HELLAS LTD	GR
Olmesartan/Hidroclorotiazida Mylan 40 mg/12,5 mg comprimidos recubiertos con película EFG	PT/H/1155/003	80361	MYLAN, LDA	ES
Olmesartan/Hidroclorotiazida Mylan 40 mg/25 mg comprimidos recubiertos con película EFG	PT/H/1155/004	80362	MYLAN, LDA	ES
Olmetec Plus 40 mg/12,5 mg comprimés pelliculés	DE/H/0523/003	BE368645	DAIICHI SANKYO BELGIUM S.A	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
Olmotec Plus 40 mg/25 mg comprimés pelliculés	DE/H/0523/004	BE368654	DAIICHI SANKYO BELGIUM S.A	BE
Olmotec Plus 40 mg/12,5 mg Filmtabletten	DE/H/0523/003	1-29037	DAIICHI SANKYO AUSTRIA GMBH	AT
Olmotec Plus 40 mg/25 mg Filmtabletten	DE/H/0523/004	1-29038	DAIICHI SANKYO AUSTRIA GMBH	AT
Olmotec Plus 40mg/12,5mg filmomhulde tabletten	DE/H/0523/003	BE368645	DAIICHI SANKYO BELGIUM S.A	BE
Olmotec Plus 40mg/25mg filmomhulde tabletten	DE/H/0523/004	BE368654	DAIICHI SANKYO BELGIUM S.A	BE
Olmotec Plus, filmovertrukne tabletter 40 mg/12,5mg	DE/H/0523/003	44538	DAIICHI SANKYO EUROPE GMBH	DK
Olmotec Plus, filmovertrukne tabletter 40 mg/25mg	DE/H/0523/004	44539	DAIICHI SANKYO EUROPE GMBH	DK
Olmotec Plus 40 mg/12,5 mg tabletti, kalvopäällysteinen	DE/H/0523/003	27135	DAIICHI SANKYO EUROPE GMBH	FI
Olmotec Plus 40 mg/25 mg tabletti, kalvopäällysteinen	DE/H/0523/004	27136	DAIICHI SANKYO EUROPE GMBH	FI
Olmotec Plus 40 mg/12,5 mg filmuhúðaðar töflur	DE/H/0523/003	IS/1/09/050/01	DAIICHI SANKYO EUROPE GMBH	IS
Olmotec Plus 40 mg/25 mg filmuhúðaðar töflur	DE/H/0523/004	IS/1/09/050/02	DAIICHI SANKYO EUROPE GMBH	IS
Benetor Plus 40 mg/12.5 mg film-coated tablets	DE/H/0523/003	PA1595/2/3	DAIICHI SANKYO IRELAND LIMITED	IE
Benetor Plus 40 mg/25 mg film-coated tablets	DE/H/0523/004	PA1595/2/4	DAIICHI SANKYO IRELAND LIMITED	IE
Olmotec Comp 40 mg/12,5 mg tabletter, filmdrasjerte	DE/H/0523/003	08-6469	DAIICHI SANKYO EUROPE GMBH	NO
Olmotec Comp 40 mg/25 mg tabletter, filmdrasjerte	DE/H/0523/004	08-6470	DAIICHI SANKYO EUROPE GMBH	NO
OLMEGAN 40 mg/12,5 mg compresse rivestite con film	DE/H/0523/003	037110234	DAIICHI SANKYO ITALIA S.P.A	IT
OLMEGAN 40 mg/12,5 mg compresse rivestite con film	DE/H/0523/003	037110246	DAIICHI SANKYO ITALIA S.P.A	IT

<b>Product Name (in authorisation country)</b>	<b>MRP/DCP Authorisation number</b>	<b>National Authorisation Number</b>	<b>MAH of product in the member state</b>	<b>Member State where product is authorised</b>
OLMEGAN 40 mg/12,5 mg compresse rivestite con film	DE/H/0523/003	037110259	DAIICHI SANKYO ITALIA S.P.A	IT
OLMEGAN 40 mg/12,5 mg compresse rivestite con film	DE/H/0523/003	037110261	DAIICHI SANKYO ITALIA S.P.A	IT
OLMEGAN 40 mg/12,5 mg compresse rivestite con film	DE/H/0523/003	037110273	DAIICHI SANKYO ITALIA S.P.A	IT
OLMEGAN 40 mg/12,5 mg compresse rivestite con film	DE/H/0523/003	037110285	DAIICHI SANKYO ITALIA S.P.A	IT
OLMEGAN 40 mg/12,5 mg compresse rivestite con film	DE/H/0523/003	037110297	DAIICHI SANKYO ITALIA S.P.A	IT
OLMEGAN 40 mg/12,5 mg compresse rivestite con film	DE/H/0523/003	037110309	DAIICHI SANKYO ITALIA S.P.A	IT
OLMEGAN 40 mg/12,5 mg compresse rivestite con film	DE/H/0523/003	037110311	DAIICHI SANKYO ITALIA S.P.A	IT
OLMEGAN 40 mg/12,5 mg compresse rivestite con film	DE/H/0523/003	037110323	DAIICHI SANKYO ITALIA S.P.A	IT
OLMEGAN 40 mg/12,5 mg compresse rivestite con film	DE/H/0523/003	037110335	DAIICHI SANKYO ITALIA S.P.A	IT
OLMEGAN 40 mg/12,5 mg compresse rivestite con film	DE/H/0523/003	037110347	DAIICHI SANKYO ITALIA S.P.A	IT
OLMEGAN 40 mg/25 mg compresse rivestite con film	DE/H/0523/004	037110436	DAIICHI SANKYO ITALIA S.P.A	IT
OLMEGAN 40 mg/25 mg compresse rivestite con film	DE/H/0523/004	037110350	DAIICHI SANKYO ITALIA S.P.A	IT
OLMEGAN 40 mg/25 mg compresse rivestite con film	DE/H/0523/004	037110362	DAIICHI SANKYO ITALIA S.P.A	IT
OLMEGAN 40 mg/25 mg compresse rivestite con film	DE/H/0523/004	037110374	DAIICHI SANKYO ITALIA S.P.A	IT
OLMEGAN 40 mg/25 mg compresse rivestite con film	DE/H/0523/004	037110386	DAIICHI SANKYO ITALIA S.P.A	IT
OLMEGAN 40 mg/25 mg compresse rivestite con film	DE/H/0523/004	037110398	DAIICHI SANKYO ITALIA S.P.A	IT
OLMEGAN 40 mg/25 mg compresse rivestite con film	DE/H/0523/004	037110400	DAIICHI SANKYO ITALIA S.P.A	IT



<b>Product Name (in authorisation country)</b>	<b>MRP/DCP Authorisation number</b>	<b>National Authorisation Number</b>	<b>MAH of product in the member state</b>	<b>Member State where product is authorised</b>
OLMEGAN 40 mg/25 mg compresse rivestite con film	DE/H/0523/004	037110412	DAIICHI SANKYO ITALIA S.P.A	IT
OLMEGAN 40 mg/25 mg compresse rivestite con film	DE/H/0523/004	037110424	DAIICHI SANKYO ITALIA S.P.A	IT
OLMEGAN 40 mg/25 mg compresse rivestite con film	DE/H/0523/004	037110448	DAIICHI SANKYO ITALIA S.P.A	IT
OLMEGAN 40 mg/25 mg compresse rivestite con film	DE/H/0523/004	037110451	DAIICHI SANKYO ITALIA S.P.A	IT
OLMEGAN 40 mg/25 mg compresse rivestite con film	DE/H/0523/004	037110463	DAIICHI SANKYO ITALIA S.P.A	IT
Olmetec Plus 40 mg/12,5 mg comprimés pelliculés	DE/H/0523/003	1851/10080021	DAIICHI SANKYO BELGIUM S.A	LU
Olmetec Plus 40 mg/25 mg comprimés pelliculés	DE/H/0523/004	1851/10080022	DAIICHI SANKYO BELGIUM S.A	LU
Olmetec Plus 40 mg + 12,5 mg, comprimidos revestidos por película	DE/H/0523/003	5350228	DAIICHI SANKYO PORTUGAL,LDA	PT
Olmetec Plus 40 mg + 12,5 mg, comprimidos revestidos por película	DE/H/0523/003	5350236	DAIICHI SANKYO PORTUGAL,LDA	PT
Olmetec Plus 40 mg + 25 mg, comprimidos revestidos por película	DE/H/0523/004	5350244	DAIICHI SANKYO PORTUGAL,LDA	PT
Olmetec Plus 40 mg + 25 mg, comprimidos revestidos por película	DE/H/0523/004	5350251	DAIICHI SANKYO PORTUGAL,LDA	PT
CoOLMETEC 40 mg/25 mg, comprimé pelliculé	DE/H/0523/004	576 795-3	DAIICHI SANKYO FRANCE SAS	FR
CoOLMETEC 40 mg/25 mg, comprimé pelliculé	DE/H/0523/004	350 248-1	DAIICHI SANKYO FRANCE SAS	FR
CoOLMETEC 40 mg/25 mg, comprimé pelliculé	DE/H/0523/004	350 247-5	DAIICHI SANKYO FRANCE SAS	FR
Olmetec Plus 40 mg/12,5 mg Filmtabletten	DE/H/0523/003	58589.02.00	DAIICHI SANKYO EUROPE 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
Olmetec Plus 40 mg/25 mg Filmtabletten	DE/H/0523/004	58589.03.00	DAIICHI SANKYO EUROPE GMBH	DE
Olmetec Plus 40 mg/12,5 mg comprimidos recubiertos con película	DE/H/0523/003	72.695	DAIICHI SANKYO ESPAÑA, S.A.	ES
Olmetec Plus 40 mg /25 mg comprimidos recubiertos con película	DE/H/0523/004	72.697	DAIICHI SANKYO ESPAÑA, S.A.	ES
OLMETEC HCTZ 40 mg/12,5 mg filmomhulde tabletten	DE/H/0523/003	RVG 104240	DAIICHI SANKYO NEDERLAND B.V.	NL
OLMETEC HCTZ 40 mg/25 mg filmomhulde tabletten	DE/H/0523/004	RVG 104241	DAIICHI SANKYO NEDERLAND B.V.	NL
CoOLMETEC 40 mg/12,5 mg, comprimé pelliculé	DE/H/0523/003	350 245-2	DAIICHI SANKYO FRANCE SAS	FR
CoOLMETEC 40 mg/12,5 mg, comprimé pelliculé	DE/H/0523/003	576 794-7	DAIICHI SANKYO FRANCE SAS	FR
CoOLMETEC 40 mg/12,5 mg, comprimé pelliculé	DE/H/0523/003	350 246-9	DAIICHI SANKYO FRANCE SAS	FR
Olmetec Plus 40 mg/12.5 mg film-coated tablets	DE/H/0523/003	PL 08265/0029	DAIICHI SANKYO UK LTD	UK
Olmetec Plus 40 mg/25 mg film-coated tablets	DE/H/0523/004	PL 08265/0030	DAIICHI SANKYO UK LTD	UK
OLMETEC PLUS® 40 mg/25 mg επικαλυμμένο με λεπτό υμένιο δισκίο	DE/H/0523/004	77204/14-12-2010	PFIZER HELLAS, A.E.	GR
OLMETEC PLUS® 40 mg/12,5 mg επικαλυμμένο με λεπτό υμένιο δισκίο	DE/H/0523/003	77203/14-12-2010	PFIZER HELLAS, A.E.	GR