



EUROPEAN MEDICINES AGENCY
SCIENCE MEDICINES HEALTH

05 June 2025
EMA/193200/2025
Human Medicines Division

List of nationally authorised medicinal products

Active substance(s): letrozole

Procedure No. PSUSA/00001842/202410



Product Name (in authorisation country)	MRP/DCP Authorisation number	National Authorisation Number	MAH of product in the member state	Member State where product is authorised
Letrozole 2.5 mg Film-coated tablets	not available	PL 20416/0414	CRESCENT PHARMA LIMITED	XI
Letrozole 2.5 mg Film-coated tablets	not available	PL 25258/0198	GLENMARK PHARMACEUTICALS EUROPE LIMITED	XI
Letrozole 2.5mg film-coated tablets	not available	PL 14251/0083	MANX HEALTHCARE LTD	XI
Femara 2,5 mg επικαλυμμένα με λεπτό υμένιο δισκία	FR/H/0110/001	233170101	NOVARTIS (HELLAS) S.A.C.I.	GR
Фемара 2,5 mg филмирани таблетки	FR/H/0110/001	20020268	NOVARTIS EUROPHARM LIMITED	BG
Femara 2,5 mg filmsko obložene tablete	FR/H/0110/001	H/03/00606/001	NOVARTIS EUROPHARM LIMITED	SI
Femara 2,5 mg filmsko obložene tablete	FR/H/0110/001	H/03/00606/002	NOVARTIS EUROPHARM LIMITED	SI
Femara 2,5 mg filmsko obložene tablete	FR/H/0110/001	H/03/00606/003	NOVARTIS EUROPHARM LIMITED	SI
Femara 2,5 mg filmsko obložene tablete	FR/H/0110/001	H/03/00606/004	NOVARTIS EUROPHARM LIMITED	SI
Femara 2,5 mg filmsko obložene tablete	FR/H/0110/001	H/03/00606/005	NOVARTIS EUROPHARM LIMITED	SI
Femara 2,5 mg comprimidos revestidos por película	FR/H/0110/001	2499986	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Femara 2,5 mg comprimidos revestidos por película	FR/H/0110/001	2500080	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Femara 2,5 mg comprimidos revestidos por película	FR/H/0110/001	5123583	NOVARTIS FARMA - PRODUTOS FARMACÊUTICOS S.A.	PT
Femara 2,5 mg compresse rivestite con film	FR/H/0110/001	033242037	NOVARTIS FARMA S.P.A.	IT
Femara 2,5 mg compresse rivestite con	FR/H/0110/001	033242049	NOVARTIS FARMA S.P.A.	IT

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film				
Femara 2,5 mg compresse rivestite con film	FR/H/0110/001	033242052	NOVARTIS FARMA S.P.A.	IT
Femara 2,5 mg compresse rivestite con film	FR/H/0110/001	033242013	NOVARTIS FARMA S.P.A.	IT
Femara 2,5 mg compresse rivestite con film	FR/H/0110/001	033242025	NOVARTIS FARMA S.P.A.	IT
Loxifan 2,5 mg comprimidos recubiertos con película	not available	62.079	NOVARTIS FARMACÉUTICA S.A.	ES
Femara 2,5 mg comprimidos recubiertos con película	FR/H/0110/001	61.628	NOVARTIS FARMACÉUTICA S.A.	ES
Femar 2,5 mg tabletti, kalvopäällysteinen	FR/H/0110/001	12452	NOVARTIS FINLAND OY	FI
Femar 2,5 mg filmdragerad tablett	FR/H/0110/001	12452	NOVARTIS FINLAND OY	FI
Femar, filmovertrukne tabletter	FR/H/0110/001	18420	NOVARTIS HEALTHCARE A/S	DK
Femar 2,5 mg filmuhúðuð tafla.	FR/H/0110/001	960198	NOVARTIS HEALTHCARE A/S	IS
Femar 2,5 mg filmuhúðuð tafla.	FR/H/0110/001	960198	NOVARTIS HEALTHCARE A/S	IS
Femar 2,5 mg filmuhúðuð tafla.	FR/H/0110/001	960198	NOVARTIS HEALTHCARE A/S	IS
Femar 2,5 mg filmuhúðuð tafla.	FR/H/0110/001	960198	NOVARTIS HEALTHCARE A/S	IS
Femar 2,5 mg filmuhúðuð tafla.	FR/H/0110/001	960198	NOVARTIS HEALTHCARE A/S	IS
Femara 2,5 mg filmom obložene tablete	FR/H/0110/001	HR-H-020668711-02	NOVARTIS HRVATSKA D.O.O.	HR
Femara 2,5 mg filmom obložene tablete	FR/H/0110/001	HR-H-020668711-03	NOVARTIS HRVATSKA D.O.O.	HR
Femara 2,5 mg filmom obložene tablete	FR/H/0110/001	HR-H-020668711-04	NOVARTIS HRVATSKA D.O.O.	HR

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Femara 2,5 mg filmom obložene tablete	FR/H/0110/001	HR-H-020668711-05	NOVARTIS HRVATSKA D.O.O.	HR
Femara 2,5 mg filmom obložene tablete	FR/H/0110/001	HR-H-020668711-01	NOVARTIS HRVATSKA D.O.O.	HR
Femara 2,5 mg filmtabletta	FR/H/0110/001	OGYI-T-5712/02	NOVARTIS HUNGÁRIA KFT.	HU
Femara® 2,5 mg, επικαλυμμένο με λεπτό υμένιο δισκίο	FR/H/0110/001	18468	NOVARTIS IRELAND LIMITED	CY
Femara 2.5 mg film-coated tablets	FR/H/0110/001	PA0896/012/001	NOVARTIS IRELAND LIMITED	IE
Femar 2,5 mg filmdrasjerte tabletter	FR/H/0110/001	96-1423	NOVARTIS NORGE AS	NO
Femara 2,5 mg - Filmtabletten	FR/H/0110/001	1-21781	NOVARTIS PHARMA GMBH	AT
Femara® 2,5 mg Filmtabletten	FR/H/0110/001	38895.00.00	NOVARTIS PHARMA GMBH	DE
FEMARA 2,5 mg Filmtabletten	FR/H/0110/001	BE182926	NOVARTIS PHARMA N.V.	BE
Femara 2,5 mg comprimés pelliculés	FR/H/0110/001	BE182926	NOVARTIS PHARMA N.V.	BE
Femara 2,5 mg filmomhulde tabletten	FR/H/0110/001	BE182926	NOVARTIS PHARMA N.V.	BE
Femara 2,5 mg comprimés pelliculés	FR/H/0110/001	2008079869	NOVARTIS PHARMA N.V.	LU
FEMARA 2,5 mg Filmtabletten	FR/H/0110/001	2008079869	NOVARTIS PHARMA N.V.	LU
FEMARA 2,5 mg, comprimé pelliculé	FR/H/0110/001	FR/H/0110/001	NOVARTIS PHARMA S.A.S.	FR
FEMARA 2,5 mg, comprimé pelliculé	FR/H/0110/001	FR/H/0110/001	NOVARTIS PHARMA S.A.S.	FR
FEMARA 2,5 mg, comprimé pelliculé	FR/H/0110/001	FR/H/0110/001	NOVARTIS PHARMA S.A.S.	FR
FEMARA 2,5 mg, comprimé pelliculé	FR/H/0110/001	34009 341 474 2 5	NOVARTIS PHARMA S.A.S.	FR
FEMARA 2,5 mg, comprimé pelliculé	FR/H/0110/001	34009 341 475 9 3	NOVARTIS PHARMA S.A.S.	FR
Femar 2,5 mg	FR/H/0110/001	13261	NOVARTIS SVERIGE AB	SE

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filmdragerad tablett				
Femara, 2,5 mg õhukese polümeerikattega tabletid	FR/H/0110/001	246799	SIA NOVARTIS BALTICS	EE
Femara 2,5 mg plēvele dengtos tabletēs	FR/H/0110/001	LT/1/98/0128/002	SIA NOVARTIS BALTICS	LT
Femara 2,5 mg plēvele dengtos tabletēs	FR/H/0110/001	LT/1/98/0128/003	SIA NOVARTIS BALTICS	LT
Femara 2,5 mg plēvele dengtos tabletēs	FR/H/0110/001	LT/1/98/0128/004	SIA NOVARTIS BALTICS	LT
Femara 2,5 mg plēvele dengtos tabletēs	FR/H/0110/001	LT/1/98/0128/005	SIA NOVARTIS BALTICS	LT
Femara 2,5 mg plēvele dengtos tabletēs	FR/H/0110/001	LT/1/98/0128/001	SIA NOVARTIS BALTICS	LT
Letrozole 2.5mg film-coated tablets	not available	PL 36687/0308	TORRENT PHARMA (UK) LTD.	XI