



EUROPEAN MEDICINES AGENCY
SCIENCE MEDICINES HEALTH

24 January 2018
EMA/44742/2018
Human Medicines Evaluation Division

List of nationally authorised medicinal products

Active substance: mifepristone / misoprostol

Procedure no.: PSUSA/00010378/201705



Product Name (in authorisation country)	MRP / DCP Authorisation number	National Authorisation Number	MAH of product in the member state	Member State where product is authorised
Medabon Combipack of Mifepristone 200 mg tablet and Misoprostol 4 x 0.2 mg vaginal tablets	SE/H/0752/001	PL 31750/0042	SUN PHARMACEUTICAL INDUSTRIES EUROPE B.V.	UK
Medabon Pachet combinat de Mifepriston 200 mg comprimate și Misoprostol 4 x 0,2 mg comprimate vaginale	SE/H/0752/001	5071/2012/01	SUN PHARMACEUTICAL INDUSTRIES EUROPE B.V.	RO
Sunmedabon, Combinatieverpakking mifepriston 200 mg tablet en misoprostol 4 x 0,2 mg vaginale tabletten	SE/H/0752/001	RVG 106099	SUN PHARMACEUTICAL INDUSTRIES EUROPE B.V.	NL
Medabon Kombinationsförpackning av mifepriston 200 mg tablett och misoprostol 4 x 0,2 mg vaginaltabletter	SE/H/0752/001	43107	SUN PHARMACEUTICAL INDUSTRIES EUROPE B.V.	SE