REBLOZYL® (luspatercept) Prescriber's Checklist

Important information for healthcare providers prescribing REBLOZYL for women of)f
childbearing potential	

The Prescriber's Checklist is to be used before initiating treatment, at each administration, and then at regular intervals when performing follow-up.

For more information about REBLOZYL or to obtain further copies of this document, please contact AM Mangion Ltd on 00 n356 2397 6333

Please see the REBLOZYL SmPC for complete prescribing information.

REBLOZYL® (luspatercept) Prescriber's Checklist

REBLOZYL Prescriber's Checklist for Women of Childbearing Potential

Patient Identification	Prescriber Details
Name:	Name:
	Signature:
	Date:

Prior to Initiation of Treatment			
Treatment with REBLOZYL is contraindicated in pregnancy and in women of childbearing potential not using at least one			
highly effective method of contraception			
- There are no data from the use of REBLOZYL in pregnant women. Studies in animals have shown reproductive			
toxicity and embryo-foetal toxicity. Clinical implications are potential foetal loss and teratogenicity.			
□ Provide counselling before treatment initiation regarding the potential teratogenic risk of REBLOZYL and required			
actions to minimise this risk.			
☐ Inform women of childbearing potential of the necessity for at least one highly effective method of contraception			
while on treatment and for 3 months after discontinuation.			
A pregnancy test must be carried out and a negative result must be verified in women of childbearing potential			
before starting treatment.			
☐ Provide the Patient Card to a women of childbearing potential.			
Duration of Treatment			
□ Provide regularly counselling regarding the potential teratogenic risk of REBLOZYL and required actions to minimise)		
this risk			
Remind women of childbearing potential that they must use at least one highly effective method of contraception			
during treatment with REBLOZYL.			
During treatment with REBLOZYL, women must not become pregnant. If a woman becomes pregnant or wants to			
become pregnant, REBLOZYL should be discontinued.			
During treatment with REBLOZYL, pregnancy tests must be repeated at suitable intervals and medically verified as			
negative.			
Discontinuation of Treatment			
□ Counsel women of childbearing potential that at least one highly effective method of contraception should be			
maintained for at least 3 months following discontinuation of treatment with REBLOZYL.			
☐ Provide counselling in the event of pregnancy and evaluation of the outcome of any pregnancy.			
□ Not applicable (this patient did not become pregnant while on treatment or within 3 months of			
discontinuation of REBLOZYL.)			

Should a pregnancy occur during treatment or within 3 months following discontinuation of treatment with REBLOZYL, remind the patient that it should be reported to the prescriber, Medicines Authority and to BMS byAM Mangion Ltd on Tel 00 356 23976333 and email - pv@ammangion.com, irrespective of adverse outcomes observed.

This Prescriber checklist fulfils the conditions of the marketing authorisation and has been approved by the competent authority, version number 2007-MT-2500001 and date of approval by the NCA "<month> <year>.

REPORTING OF ADVERSE REACTIONS

Suspected adverse reactions and medication errors should be reported at ADR Reporting, The Medicines Authority, Post-Licensing Directorate, Sir Temi Zammit Buildings, Malta Life Sciences Park, San Gwann SGN 3000, Malta

Website: www.medicinesauthority.gov.mt

e-mail: postlicensing.medicinesauthority@gov.mt

OR

ADR Reporting: www.medicinesauthority.gov.mt/adrportal

AND

AM Mangion Ltd Mangion House New Street off Valletta Road Luqa LQA6000, Malta

Email: pv@ammangion.com

Tel - 00 356 23976333