

November 25, 2021

Teriflunomide Exposure in Pregnancy Form

Date: _____

Patient I.D.: _____

Country / Province: _____

Report Type:

☐ Initial

☐ Follow up

Exposure during pregnancy:
☐ Maternal

☐ Paternal

Paternal Information:

Date of Birth (DD-MMM-YYYY):

Age: _____ years

 Ethnicity: ☐ Asian ☐ Black ☐ Caucasian ☐ Hispanic ☐ Other, specify:

 Weight: _____ ☐ kgs ☐ lbs

 Height: _____ ☐ cm ☐ in

Rhesus Factor: _____

Medical History

Risk Factor	Yes	No	Risk Factor	Frequency			
				Never	Occasionally	Often	Previously /Quit
Hepatitis			Substance Abuse				
Hypertension			Alcohol				
Psychiatric Illness			Smoking				
Epilepsy							
Diabetes							
HIV							

Other Notable Health Disorders /Conditions: Please describe							

Maternal Information:

Date of Birth (DD-MMM-YYYY): _____

Age: ____ years

Ethnicity: Asian Black Caucasian Hispanic Other, specify: _____

Weight: kgs lbs _____

Height: cm in _____

Rhesus Factor: ____

Medical History

Risk Factor	Yes	No	Risk Factor	Frequency			
				Never	Occasionally	Often	Previously /Quit
Hepatitis			Substance Abuse				
Hypertension			Alcohol				
Psychiatric Illness			Smoking				
Epilepsy							
Diabetes							
HIV							

Other Notable Health Disorders /Conditions							
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Immunizations:

Immunization	Yes, Date (DD-MMM-YYYY):	No
Rubella		
Toxoplasmosis		
CMV		

Was a contraception method used? ☐ Yes ☐ No ☐ Unknown
 If yes, please check type of contraception:

☐ Oral contraception (type not known) ☐ Oral contraception (Progesterone)
☐ Contraceptive Implant ☐ Intra-uterine device
☐ Oral contraception (Oestrogen + Progesterone)
☐ Transdermal contraception ☐ Contraceptive injection
☐ Condom

History of ☐ normal or ☐ abnormal menstrual cycles
 History of infertility ☐ Yes ☐ No

First Day of Last Menstrual Period (LMP) (DD-MMM-YYYY): _____

Estimated Delivery Date (DD-MMM-YYYY): _____
 Specify method of calculation: _____

☐ LMP
☐ Ultrasound Date (DD-MMM-YYYY): _____
☐ Other, please specify: _____

Did you become pregnant while on teriflunomide? ☐ Yes ☐ No
 If you got pregnant while on teriflunomide, was accelerated elimination used? ☐ Yes ☐ No

Teriflunomide Dosage at conception:

Gestational Age at Last Dose:

Duration of Treatment with Product while Pregnant:							
Did you become pregnant after teriflunomide discontinuation? <input type="checkbox"/> Yes <input type="checkbox"/> No If yes, was accelerated elimination used? <input type="checkbox"/> Yes <input type="checkbox"/> No If yes, did you become pregnant within 11 days of teriflunomide discontinuation? <input type="checkbox"/> Yes <input type="checkbox"/> No If accelerated elimination was not used, did you become pregnant within 2 years of teriflunomide discontinuation? <input type="checkbox"/> Yes <input type="checkbox"/> No							
PATIENT'S MEDICAL HISTORY (include information on familial disorders, known risk factors or conditions that may affect the outcome of the pregnancy e.g. alcohol, smoking, other substance consumption, hypertension, eclampsia, diabetes including gestational, infections during pregnancy, environmental or occupational exposure that may pose a risk factor):							
PREVIOUS OBSTETRIC HISTORY - provide details on all previous pregnancies below, including abortion or stillbirth: _____ _____ Gestation Weeks at Delivery: _____ Outcome of the pregnancy including any previous maternal complications and previous fetal / neonatal abnormalities and type: _____ _____							
Family History: Is there any history of congenital abnormalities, children dying young, chromosomal abnormalities, developmental delays or hereditary diseases in paternal or maternal family? <input type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> Unknown If yes, please specify: Blood relationship between parents? <input type="checkbox"/> Yes <input type="checkbox"/> No <input type="checkbox"/> Unknown (If yes, specify degree)							
DRUG INFORMATION - please list all medications, including OTC medications, and dietary supplements taken prior to or during pregnancy							
Drug Name	Daily Dose	Route	Treatment Dates		Indication	Week of pregnancy	
			Start (DD-MMM-YYYY):	Stop (DD-MMM-YYYY):		Start	Stop

Were administered drugs discontinued due to pregnancy? ☐ Yes ☐ No
 If yes, which drugs? _____

PRENATAL TESTING:

Have any specific tests, e.g. amniocentesis, ultrasound, maternal serum AFP, chorionic villi sampling, fetal stress test, genetic screening or other been performed during the pregnancy so far?

☐ Yes ☐ No ☐ Unknown

If yes, please specify test date and results:

Test	Date: (DD-MMM-YYYY)	Results

PREGNANCY OUTCOME

Pregnancy Ongoing: ☐ Yes ☐ No

If yes, Gestational age: (weeks) _____

Number of embryos / foetus(es): _____

Last ultrasound scan date (DD-MMM-YYYY): _____

☐ Normal ☐ Abnormal, please specify: _____

Delivery Date: (DD-MMM-YYYY): _____

☐ Vaginal ☐ Forceps/ventouse ☐ Caesarean section

Status of amniotic fluid: ☐ Clear ☐ Not clear

Placenta: ☐ Normal ☐ Abnormal

Medications provided during delivery: ☐ yes, please specify _____ ☐ No

Delivery duration: _____

Maternal complications or problems related to birth: _____

Abortion

Date:

☐ Therapeutic ☐ Elective ☐ Spontaneous

Please, specify reason and any abnormalities (if known): _____

☐ Unspecified: _____

At week ____

Complication:

☐ Mother died (DD-MMM-YYYY): _____

☐ Neonate died (DD-MMM-YYYY): _____

MATERNAL PREGNANCY ASSOCIATED EVENTS:

If the mother experienced an adverse drug reaction during pregnancy, please complete a data collection form and submit as requested to the Sponsor and to the Canada Vigilance Program (<https://www.canada.ca/en/health-canada/services/drugs-health-products/medeffect-canada/adverse-reaction-reporting.html>)

Date	Drug	Adverse Event	Outcome	Form Tracking Number

First trimester Follow-up (please provide details of embryo/fetal development):

Second trimester Follow-up (please provide details of embryo/fetal development):

Third trimester Follow-up (please provide details of embryo/fetal development):

CHILD INFORMATION:

Neonate

☐ Live [Normal] ☐ Live with congenital abnormality ☐ Stillbirth at week

Please specify any abnormalities: _____

☐ Full term ☐ Premature Number of weeks _____ ☐ Post-mature Number of weeks _____

Sex: ☐ Male ☐ Female

Height: _____ cms Weight: _____ kgs

Apgar Scores: _____ 1 min _____ 5 mins _____ 10 mins

Head circumference: _____ cms

☐ Breast Fed ☐ Bottle Fed

Neonatal Illness, developmental delay or immaturity? ☐ Yes, Please specify
 _____ ☐ No

Corrective treatment Required? ☐ Yes, Please specify _____ ☐ No

Transfer to ICU or paediatric department?

☐ Yes, please provide details of location and contact information _____

☐ No

For additional information, (please provide copies of relevant documentation)

ASSESSMENT OF PREGNANCY OUTCOME

SERIOUSNESS CRITERIA

☐ Non-serious ☐ Congenital anomaly/birth defect ☐ Death of mother or neonate

☐ Involved or prolonged inpatient hospitalization ☐ Life-threatening (immediate risk of death)

☐ Other significant medical events (may jeopardise the patient or require intervention to prevent one of other criteria).

☐ Resulted in persistent or significant disability/incapacity.

REPORTER INFORMATION

Name: _____

Title: _____



Address: _____

City: _____ **Province:** _____ **Postal Code:** _____

Country: _____

Institution: _____ **Department:** _____

Phone: _____ **Fax:** _____ **E- mail:** _____

Healthcare professional: ☐ Yes ☐ No **If yes, please specify occupation:**

Did patient give consent to follow up with their Healthcare Practitioner for pregnancy outcome and at intervals of 1 week, 6, 12 and 24 months post-delivery?

Patient Name: _____

Healthcare Practitioner:

Name: _____

Address: _____

Phone: _____

Email: _____