

November 25, 2021

Teriflunomide Exposure in Pregnancy Form

Date:								
Patient I.D.:		_						
Country / Province:								
Report Type:								
Initial								
Follow up								
Exposure during p	oregnan	icy:						
☐ Maternal								
☐ Paternal								
Paternal Informat	tion:							
Age:years Ethnicity: Asiar Weight:k Height:cr Rhesus Factor: Medical History Risk Factor	gs 🗆 lb n 🗀 in		Caucasian H	ispanic 🗌	Other, specify:			
NISK Factor	163	NO	KISK Factor					
				Never	Occasionally	Often	Previously /Quit	
Hepatitis			Substance Abuse					
Hypertension			Alcohol					
Psychiatric			Smoking					
Illness								
Epilepsy								
Diabetes								
HIV								

CGPA-Member Companies' Teriflunomide Enhanced Pharmacovigilance Pregnancy Active Surveillance Program

November 25, 2021



	Other Notable Health Disorders /Conditions: Please describe								
	Maternal Information:								
	Date of Birth (DD-N Age: years	'IIVIIVI-1	(
	thnicity: Asian I	Black C	aucasi	an Hispanic O	ther, spec	ify:			
١	Neight: kgs lbs			•					
ŀ	leight: cm in		_						
F	Rhesus Factor:								
	1edical History								
Ī	Risk Factor	Yes	No	Risk Factor		Frequ	iencv		
					Never			Previously	_
					Never	Occasionally	Orten	/Quit	'
Ī	Hepatitis			Substance					
				Abuse					
L	Hypertension			Alcohol					
	Psychiatric			Smoking					
ŀ	Illness								
ŀ	Epilepsy								
-	Diabetes						+		
	HIV								



Other Notable Health Disorders /Conditions										
Immunizations:										
Immunization		Yes, Date (I	Yes, Date (DD-MMM-YYYY):			No				
Rubella										
Toxoplasmosis										
CMV										
Was a contraception If yes, please check to			No □Unk	nown						
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 Me	First Day of Last Menstrual Period (LMP) (DD-MMM-YYYY):									
Estimated Delivery [Date (DD-N	/IMM-YYYY):								
Specify method of ca	alculation:									
 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 Dosaş	ge at conc	eption:								
Gestational Age at L	ast Dose:									



Duration of Treatment with Product while Pregnant:							
Did you become pregnant after teriflunomide discontinuation? Yes No If yes, was accelerated elimination used? Yes No If yes, did you become pregnant within 11 days of teriflunomide discontinuation? Yes No If accelerated elimination was not used, did you become pregnant within 2 years of teriflunomide discontinuation? Yes No							
risk factors or other substance	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 OBS		•		•	. •		
Outcome of th	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? Yes No Unknown							
If yes, please specify: Blood relationship between parents? Yes No Unknown (If yes, specify degree)							
DRUG INFORMATION - please list all medications, including OTC medications, and dietary supplements taken prior to or during pregnancy							
			Treatme	nt Dates		Week of p	regnancy
Drug Name	Daily Dose	Route	Start (DD-MMM- YYYY):	Stop (DD-MMM- YYYY):	Indication St	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	l results:							
Test	Date: (DD-MMM-YYY)	Result	s					
	•	<u>.</u>			<u>.</u>			
PREGNANCY OUTCOME								
Pregnancy Ongoing: Yes No								
If yes, Gestational age: (weeks) Number of embryos / foetus(es):								
1	48.4.10000							
Last ultrasound scan date (DD-MN Normal Abnormal, please sp	,							
Delivery Date: (DD-MMM-YYYY):								
□Vaginal □ Forceps/ventouse □ Caesarean section								
Status of amniotic fluid: Clear	Status of amniotic fluid: Clear Not clear							
Placenta: Normal Abnormal								
Medications provided during delivery: yes, please specify No								

CGPA-Member Companies' Teriflunomide Enhanced Pharmacovigilance Pregnancy Active Surveillance Program
November 25, 2021



Delivery duration: _		_		
Maternal complicat	tions or problems	related to birth:		
Abortion Date:				
Therapeutic Please, specify reas		ntaneous rmalities (if known):		
Unspecified:				
At week Complication:				
		Y): YY):		
collection form and	rienced an advers d submit as reque ada.ca/en/health-	se drug reaction during sted to the Sponsor an canada/services/drug	nd to the Canada	Vigilance Program
Date	Drug	Adverse Event	Outcome	Form Tracking Number
First trimester Follo	ow-up (please pro	ovide details of embryo	o/fetal developmo	ent):
Second trimester F	ollow-up (please	provide details of emb	ryo/fetal develop	oment):
Third trimester Fol	low-up (please pr	ovide details of embry	o/fetal developm	nent):
CHILD INFORMATION Neonate	DN:			

CGPA-Member Companies' Teriflunomide Enhanced Pharmacovigilance Pregnancy Active Surveillance Program
November 25, 2021



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:kgs						
Apgar Scores:1 min5 mins10 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:						

CGPA-Member Companies' Teriflunomide Enhanced Pharmacovigilance Pregnancy Active Surveillance Program

November 25, 2021



Address:			
City:	Province:	Postal Code:	_
Country:			
Institution:		Department:	_
Phone:	Fax:	E- mail:	_
Healthcare professional: Y	es 🗌 No 🛮 If yes, plea	se specify occupation:	
Did patient give consent to fo intervals of 1 week, 6, 12 and	•	olthcare Practitioner for pregnancy outcome ery?	and at
Patient Name:			
Healthcare Practitioner:			
Name:		_	
Address:		-	
Phone:		_	
Email:		_	