ull Bristol Myers Squibb™

Pregnancy Surveillance Form Part I (Antepartum Information)

PATIENT IDENTIFIER: (FOR STUDIES, MUST INCLUDE PROTOCOL, SITE & SUBJECT NUMBERS)		Case # (BMS only)					Loca	LOCAL COUNTRY NUMBER: (BMS ONLY)			
BMS receipt date (BMS use only)	Click here to enter	a date.				/WPS reci 3MS use o		ATE	Click	here	to enter a date.
	C SPONTAN OR	EOUS	S TUDY				Сс	UNTRY*			
REPORT TYPE:				W-UP REPORT			*If UK, was Country of Incidence, Specify if Northern Ireland below? Yes No			nce, Specify if	
EVENT: PREGNANCY											
EXPOSURE TYPE:		AL DRUG EXP	OSURE OR		P	ATERNAL DI	RUG EX	(POSURE			
for <u>Paternal Drug Exposu</u>	IRE ONLY: WAS PREGN	ANT PARTNE	R INFORMED (ONSENT FO	ORM SI	IGNED?		Γ	No		Yes
IF NO, DID THE MALE SUBJECT	PROVIDE ALL OF THE P	REGNANCY S	URVEILLANCE	INFORMATI	on be	ELOW?		Г	No		Yes
Report type:		TIVE REPORT	- OR		R	ETROSPECT	IVE RE	PORT			
WERE THERE ANY ADDITIONAL	.MATERNAL/PATERNAL	ADVERSE EV	ENTS?			🗖 No		Yes			
IF YES, REPORT THE ADVERSE	EVENTS APPROPRIATEL	Y (FOR STUD	DIES, REFER T	O STUDY-SF	PECIFI	C INSTRUCT	TIONS)				
MATERNAL INFORMATION	Age at	HEIGHT:	WEIG	HT: RA	CE:						
DATE OF BIRTH:	CONCEPTION:				WHIT	ΓE			Black		Asian
		inche	es 🔽	lb	AME	rican Indian oi	OR ALASKAN NATIVE				
		C cm		kg NATIVE HAWAIIAN OF			r Other Pacific Islander				
Click here to enter a date.				Aboriginal			Torres Strait Islander			Torres Strait Islander	
					Отн	ER RACE:					
NUMBER OF PREGNANCIES INC	LUDING THIS ONE		NUMBER	OF BIRTHS			Nume	BER OF LIVI	NG CHI	LDREN	
		Approximate	(LICK)	nere to enter a	a date.	DATE PREGNA		Click here t	o enter	a date.	,
ONSET DATE LAST MENSTRUAL Click here to enter a date.		nere to enter a	a date.	WAS CONFIRM		SE	RUM				
ESTIMATED GESTATIONAL AGE WHEN PREGNANCY DIAGNOSED: WEEKS			VEEKS	D	ETERMII	NED BY:		Fetal ultras	OUND	Γ	DATE FROM LMP
CONTRACEPTION AT TIME OF CONCEPTION	N: 🔽 N	o Г	YES	UNKNOWN			(IF YES,	SPECIFY)			
RELEVANT MATERNAL MEDICAL HISTORY/RISK FACTORS			Date	OF O	NSET	IF APPLICABLE SPECIFY PERTINENT DETAILS			PERTINENT		
				Click here	to ent	er a date.					
				Click here	to ent	er a date.					
				Click here	to ent	er a date.					
				Click here	to ent	er a date.	,				
PATERNAL INFORMATION:	Age	YEAR	RS			Date of bi	RTH:	Click her	re to en	ter a da	ate.
RELEVANT PATERNAL MEDICAL HISTORY / RISK FACTORS			DATE OF ONSET		PERTINENT						
			Click here to enter a date.								
				Click here to enter a date.							
			Click here	e to enter a date.							
				Click here to enter a date.							

H Bristol Myers Squibb

Pregnancy Surveillance Form Part I (Antepartum Information)

PATIENT IDENTIFIER: (FOR STUDIES, MUST INCLUDE PROTOCOL, SITE & SUBJECT NUMBERS)		Case # (BMS only)				LOCAL COUNTRY NUMBER: (BMS ONLY)			
MEDICATION NAME AND INDICATION	PREGNA Relate Medica	ED TO	Dose and UNITS	Freq	Route	OF D	DD(S) RUG JRE ***	Oncology drugs ONLY	START AND STOP DATES
1.								Cycle #:	Click here to enter a date.
INDICATION							_		
MATERNAL OR PATERNAL	Not	RELATED						CUMULATIVE DOSE WITH UNITS	Click here to enter a date.
Non-study or Study	REL/	ATED							OR ONGOING
2.								Cycle #:	Click here to enter a date.
							_		
MATERNAL OR PATERNAL	Not	RELATED				ļ		CUMULATIVE DOSE WITH UNITS	Click here to enter a date.
Non-study or Study	REL/	ATED							OR ONGOING
3.								CYCLE #:	Click here to enter a date.
							_		
MATERNAL OR PATERNAL	Not	RELATED						CUMULATIVE DOSE WITH UNITS	Click here to enter a date.
Non-study or Study	REL/	ATED							OR ONGOING
4.								Cycle #:	Click here to enter a date.
							_		
MATERNAL OR PATERNAL	Not	RELATED						CUMULATIVE DOSE WITH UNITS	Click here to enter a date.
Non-study or Study	RELA	ATED							OR ONGOING
5.								Cycle #:	Click here to enter a date.
							_		
MATERNAL OR PATERNAL	Not	RELATED						CUMULATIVE DOSE WITH UNITS	Click here to enter a date.
Non-study or Study	RELA	ATED							OR ONGOING
6.								Cycle #:	Click here to enter a date.
							_		
MATERNAL OR PATERNAL	Not	RELATED						CUMULATIVE DOSE WITH UNITS	Click here to enter a date.
Non-study or Study	REL4	ATED							OR ONGOING
7.								Cycle #:	Click here to enter a date.
INDICATION							_		
MATERNAL OR PATERNAL	Not	RELATED						CUMULATIVE DOSE WITH UNITS	Click here to enter a date.
Non-study or Study	RELA	ATED							OR ONGOING
* MANDATORY FOR ALL STUDIES									
**Route:									
1 = ORAL 2 = IN ***PERIOD(S) OF DRUG EXPOSURE: (INCLUD				= Subcuta	NEOUS			4 = OTHER	
$0 = PRIOR TO CONCEPTION \qquad 1 = 19$	E ALL TH			= 2ND TRIA	AESTER				

5 = UNKNOWN

4 = LABOR & DELIVERY

3 = 3rd trimester

ull Bristol Myers Squibb™

Pregnancy Surveillance Form Part I (Antepartum Information)

PATIENT IDENTIFIER: (FOR STUDIES, MUST INCLUDE PROTOCOL, SITE & SUBJECT NUMBERS)	CASE # (BMS ONLY)			LOCAL COUNTRY NUMBER: (BMS ONLY)				
	Base-	Dire	Test res	Test results		NORMAL RANGE		
PRENATAL DIAGNOSTIC TESTING	LINE	Date	UNIT		Lov	w Нідн		
		Click here to enter a date.						
		Click here to enter a date.						
		Click here to enter a date.						
		Click here to enter a date.						
		Click here to enter a date.						
		Click here to enter a date.						
		Click here to enter a date.						
DESCRIBE RESULTS IN DETAIL, IF APPLICABLE:								
REPORTER INFORMATION: BMS STUDY INVESTIGATOR NON-BMS STUDY SPONSOR OTHER*								
*QUALIFICATION: (COMPLETE ONLY IF "OTHER" IS	CHECKED)							
Physician Pharmacist Nurse/Nurse practitioner Other health professional								
CONSUMER ATTORNEY OTHER NON-HEALTH PROFESSIONAL								
Person completing the form (if different from Investigator/Sponsor): Date:								
						lick here to enter a date.		
Signature								
INSTITUTION / ORGANIZATION:								
STREET ADDRESS:				CITY:				
STATE/PROVINCE:								
DST CODE: COUNTRY: PHONE NUMBER:								
Email address								
INVESTIGATOR/SPONSOR/OTHER:								
LAST NAME								
	First N	AME				MIDDLE INITIAL		
Signature:				DATE:	Clic	k here to enter a date.		

Ull Bristol Myers Squibb™

Pregnancy Surveillance Form Part II (Pregnancy Outcome)

PATIENT IDENTIFIER: (FOR STUDIES, MUST INCLUDE PROTOCOL, SITE & SUBJECT NUMBERS)	CASE # (BMS ONLY)		LOCAL COUNTRY NUMBER: (BMS ONLY)				
PREGNANCY OUTCOME:	LA	BOR/DELIVERY COMPLICATIONS	IF YES, SPECIFY				
Single gestation Multiple gestation (# of) COMPLETE AN OUTCOME FORM FOR EACH FETUS/INFANT Did Obstetrical complications on Maternal/Paternal Medical Conditions occur during this pregnancy? Date pregnancy ended: Gestational age at outcome Weeks Unknown Unknown If yes, specify:							
*For any complications noted above, report the adverse eve	ENT APPROPRIATELY (FOR STUDIES, RI	EFER TO STUDY-SPECIFIC INSTRUCTIO	אר)				
GENDER: BIRTH WEIGHT: MALE FEMALE J / J UNKNOWN	BIRTH LENGTH: grams Birth LENGTH:		1 Min. 5 Min.				
LIVE BIRTH NORMAL (PROCEED TO PART III)							
LIVE BIRTH ABNORMAL	NEONATAL DEATH (IF	F ANY ARE CHECKED, COMPLETE SEC	TIONS BELOW)				
PRE-TERM TERM SMALL FOR GESTATIONAL AGE INTRAUTERINE GROWTH RETARDATION	Post term	FAMILY HISTORY OF CONGENIT	ral abnormalities/birth defects:				
DRUG WITHDRAWAL SYNDROME IN THE NEONA MALFORMATION (SPECIFY BELOW) POST-NATAL/NEONATAL COMPLICATIONS (E.G.		PRIOR PREGNANCIES WITH COL DEFECTS: IF YES, SPECIFY #/TYPE :	NGENITAL ABNORMALITIES/BIRTH				
INFECTION, RESPIRATORY DISTRESS) (SPECIF	Y):	PRIOR STILLBIRTHS:	No Yes				
ECTOPIC MISCARRIAGE/SPONTANEOUS	SABORTION STILLBIRTH	PRIOR SPONTANEOUS ABORTIC	DNS: NO YES				
AUTOPSY/PATHOLOGY REPORT NO	Yes Unknown	SPECIFY ANY PRIOR PREGNANC	CY COMPLICATIONS:				
CAUSE:	DATE: Click here to enter a date.						
PLACENTAL ABNORMALITIES NO	Yes Unknown						
IF YES, SPECIFY: PATHOLOGY REPORT AVAILABLE NO	Yes Unknown						
DESCRIBE ANY CONGENITAL MALFORMATIONS/ABNORMALITIE	S, STRUCTURAL DEFECTS AND OT	THER FETAL/NEONATAL COMPLIC	ATIONS:				
CAUSALITY (MANDATORY FOR STUDIES)							
IN THE INVESTIGATOR'S OPINION, WAS THE DEFECT/MEDICAL IF RELATED, PLEASE COMMENT ON SPECIFIC EVENT(S) AND N IF NOT RELATED, INDICATE WHAT THE DEFECT/MEDICAL PROP	IEDICATION(S) BELOW:	DN UNDER STUDY? :	NOT RELATED RELATED				

ulli Bristol Myers Squibb[™]

Pregnancy Surveillance Form Part III (Infant Follow-up)

PATIENT IDENTIFIER: (FOR STUDIES, MUST INCLUDE PROTOCOL, SITE & SUBJECT NUMBERS)	CASE # (B/	MS ONLY) LOCAL COUNTRY N			try Numbe	NUMBER: (BMS ONLY)	
CURRENT INFANT AGE:	10	AGE UNITS:		YS 🗖	WEEKS	Монтня	
NO PROBLEMS MEDICAL PROBLEMS NOTED (SPECIFY AND DESCRIBE FINDINGS AND/OR PLANNED EVALUATIONS;							
E.G. DIAGNOSTIC TESTING, CONSULTATIONS, ETC)							
CAUSALITY (MANDATORY FOR ALL STUDIES): IN THI	E INVESTIGATO	DR'S OPINION WERE ANY PRO	BLEMS NOTE	D ABOVE RELA	TED TO TH	Ē	
MEDICATION UNDER STUDY?	NOT RE			(PLEASE SF			
Maternal breastfeeding: No Yes How long:							
MATERNAL DRUGS TAKEN WHILE BREASTFEEDING:	MATERNAL DRUGS TAKEN WHILE BREASTFEEDING: NO Yes (IF yes, specify)						
REPORTER INFORMATION: BMS STUDY INVESTIGATOR NON-BMS STUDY SPONSOR OTHER*							
*QUALIFICATION: (COMPLETE ONLY IF "OTHER" IS CHECKED)							
PHYSICIAN PHARMACIST NURSE/NURSE PRACTITIONER OTHER HEALTH PROFESSIONAL							
CONSUMER ATTORNEY OTHER NON-HEALTH PROFESSIONAL							
Person completing the form (if different from Investigator/Sponsor): Date:						DATE:	
PRINTED NAME				Click	here to enter a date.		
Signature							
Institution/organization:							
			-	CITY:		STATE/PROVINCE:	
STREET ADDRESS:							
Post code:	COUNTRY:			PHONE NUMB	ER:		
Email address							
II INVESTIGATOR/SPONSOR/OTHER:							
LAST NAME							
2	FIRST NAME	Middle I					
Signature:		,				DATE:	
					Click	here to enter a date.	

Pregnancy Surveillance Form - Quick Reference Guide

The Pregnancy Surveillance Form will be completed for all prospective (confirmed pregnancy, prior to delivery or confirmation of congenital anomaly) and retrospective (when congenital anomaly/malformation is confirmed or after delivery has occurred) reports of pregnancy and pregnancy outcomes (live births: normal or abnormal, fetal death, neonatal death etc.) It functions as a data collection and query tool to report pregnancies and related pregnancy information. AE/SAEs for all subjects/patients reported in association with the pregnancy (obstetric complications, maternal medical complications, etc.) are to be reported separately on the clinical or non-interventional SAE form or spontaneous AE/SAE form.

Pregnancy Surveillance Form Part I	Pregnancy Surveillance Form Part II	Pregnancy Surveillance Form Part III
When a pregnancy is confirmed	When the pregnancy outcome is known	When the infant outcome is known.

Site Monitor: When a pregnancy is confirmed, collaborate with the site manager or clinical scientist to ensure that the Investigator has notified the IRB/IEC or Health Authority (if required by local law).

- Ensure that documentation of pregnancy notifications sent by the Investigator to the IRB/IEC are filed in the On-site Investigator File (OSIF) and R&D Study File.
- In countries where notification of the IRB/IEC is handled by the sponsor, the site manager is responsible for ensuring that the documentation of all pregnancy notifications sent to the IRB/IEC are filed within the R&D Study File.
- Note: for Paternal Drug Exposure in Interventional Study Reports: If pregnant partner informed consent is not signed, Part I, Part II and Part III information needs to come from the male subject, and not from the female partner herself.

All Pages Header Information

- For studies the "Patient Identifier" is the same as that used throughout the CRF, and populated with the protocol, site and subject numbers i.e. CV131-345-234-1134
- For spontaneous reports, enter local country number (if applicable) at the top left and/or enter a patient identifier (i.e. initials) if available or leave blank
- Parts I, II and III will be completed with all appropriate identifying header information on each page Part I - Page 1

Complete all questions for "PREGNANCY" as the only adverse event; other SAEs reported in association with the pregnancy (obstetric complications, maternal medical complications etc.) are reported separately either on the clinical/non-interventional study SAE form or the Spontaneous AE/SAE forms.

Part I - Page 2: Medication:

- Include each medication reported as a separate entry.
- Indicate if the drug was associated with maternal or paternal exposure.
- Indicate if the drug was identified as a non study medication or study medication by the investigator or reporter. Study medications include the medications under study (for non-interventional studies), the Investigational Medicinal Product (IMP), comparator medications and background therapy identified in the protocol.

"Pregnancy Related to Medication" Column: Check whether or not the pregnancy was related to the medication. Dosing Information: For route and period(s) of drug exposure, use the codes indicated at the bottom of the page. For period(s) of drug exposure, include all that apply.

Part I - Page 3: Prenatal Diagnostic Testing: Indicate if the results are baseline by checking under "baseline"; otherwise leave this box blank when providing the relevant details. Specify the test results (including any relevant units or other data), use the space below this section to describe results in more detail if needed.

Part II - Pregnancy Outcome: Complete delivery and outcome data as requested at the top of the page. If the outcome involved multiple gestations, please complete a separate outcome form for each fetus/infant. If the pregnancy/outcome involved labor or delivery complications, obstetric complications, or maternal medical conditions, briefly specify them. NOTE: If any complications reported above meet the definition of an SAE (or an AE for non-study patients) they should be reported separately on either the clinical or non-interventional SAE form or the spontaneous AE/SAE form. If the outcome is "live birth- normal" check this box, and proceed to the next page or any adverse outcome (live birth abnormal, fetal or neonatal death) complete all requested information to the fullest extent

For any adverse outcome (live birth abnormal, fetal or neonatal death) complete all requested information to the fullest extent possible. A detailed causality assessment by the investigator is required for any reports from trials and must be provided as noted at the bottom of this page.