Adomp				nrollmen					CO 5000	
1 Patient Info	Access thi ormation (* indic			idempasREMS.coi	n, or fax thi	s form to the A	dempas	s REMS at 1-855-6	662-5200	
First Name*:	mation ( mai	cates require	Middle	Last Name*:			В	Birthdate*		Gender*: ☐ Male
			Initial:					MM/DD/YYYY):		Female
Address Line 1*:							A	Address Line 2:		
City*:						State*:		Zip code*:		
Preferred Phone*:			Can we leave a message on this phone?   Yes   N			□ No	Preferred Time to Contact:   Day   Evening			
Cell/Alternate Ph	one:				Email:					
Alternate Contac	Contact Name: Phone:				Relationship:					
*Adempas Samp	as Sample Dispense le should only be dis	pensed as a 30			Adempas San	ple Dispensed* /	Date:		_	
	of Medical Nece			· · · · · · · · · · · · · · · · · · ·						
The following of Diagnosis*:	does not suggest a	approved use	s or indica	tions.				Therapy Statu	ıs:	
Pulmonary arteri	al hypertension		nboembolic pulmonary hypertension			HER (please speci	fy)	☐ Initial therapy (monotherapy or in combination)		
☐ I27.0 I27.24   ☐ I27.21 ☐ Inope			ble					☐ Add-on therapy ☐ Transition from other therapy		
			ent/Recurrent							
3 Female Patient Agreement										
during Adempas treatment and for one month after stopping Adempas treatment, my medical options in the event of unprotected sexual intercourse or known or suspected contraception failure, and to immediately contact my prescriber if I miss a menstrual period or suspect that I am pregnant. Before each prescription, I will receive counseling by the pharmacy or the prescriber who dispenses Adempas on the risk of serious birth defects, the need to use effective contraception during Adempas treatment and for one month after stopping Adempas treatment, to get monthly pregnancy tests, and to report a pregnancy immediately. Ensure that I have completed pregnancy testing before I started Adempas, monthly before each refill, and for one month after stopping Adempas. I understand that I may be contacted by Bayer and/or its agents and contractors to obtain information about my pregnancy. I will communicate with the pharmacy to confirm completion of pregnancy testing.  For Pre-Pubertal Females: I have been counseled on the risks of Adempas, including the risk of serious birth defects, and that I have read the Guide for Female Patients. I understand that I must immediately contact my healthcare provider if I get my menstrual period.  For Post-Menopausal Females: I have received and read the Guide for Female Patients and that I will inform my prescriber if there is a change in my reproductive status.  For Females with other medical reasons for permanent, irreversible infertility: I have received and read the Guide for Female Patients and that I will inform my prescriber if there is a change in my reproductive status.										
REQUIRED FOR ALL FEMALE Patient or Parent/Guardian Signature:  Date:  Date:										
4 Prescriber	Information (* i	ndicates rec	uired fiel	d)						
First Name*:			Last Name	*:				1	NPI*:	
Practice/Facility	Name (where you see	e this patient):						'		
Address Line 1*:					A	Idress Line 2:				
City:		(	State:	Zip code:	Р	ione*:			State Licer	nse #:
5 Prescriber	Authorization									
For female patients, please indicate the patient's current reproductive status below.    Female of NON-Reproductive Potential										uctive Potential ed prior to No
Prescriber Signature*:						<u> </u>	TO duditority.			
REQUIRED										
Definitions:  Females of Reproductive Potential, I will:  • counsel Females of Reproductive Potential (FRP) on Adempas risks, including serious birth defects; and review the <i>Guide for Female Patients</i> with the patient.  • counsel each FRP to immediately contact her prescriber if she misses a menstrual period or suspects pregnancy.										

- For the purposes of this REMS, puberty includes those girls who are at least Tanner Stage 3 and have not yet had a menses (premenarchal).
- Females of Non-Reproductive Potential, I will:
- Pre-Pubertal Females: Females who are at Tanner Stages 1 and 2 are not considered to be of reproductive potential.
- Post-Menopausal Females: Females who have passed through menopause. Menopause is defined as 12 months of spontaneous amenorrhea (not amenorrhea induced by a medical condition or medical therapy) or post-surgical from bilateral oophorectomy.
- Females with other medical reasons for permanent, irreversible infertility.

Prescriber Obligations under the Adempas REMS

For All Females, I will:

- determine the reproductive potential status of all female patients using the definitions provided in the Prescriber and Pharmacy Guide.
- advise all females that Adempas is only available through a restricted distribution program called the Adempas REMS.
- enroll all female patients into the Adempas REMS by completing the *Patient Enrollment and Consent Form* and submitting it to the REMS

- order and review pregnancy tests for FRPs before the start of treatment, monthly during treatment, and for one month after stopping treatment.
- counsel each FRP to use effective contraception during Adempas treatment and for one month after stopping treatment and discuss her medical options in the event of unprotected sexual intercourse or known or suspected contraceptive failure using the *Guide for Female Patients*.
- counsel each FRP during treatment if she is not complying with the required testing or if she is not using effective contraception, and to immediately contact her prescriber if she misses a menstrual period or suspects that she is pregnant.

For Pre-Pubertal Females, I will:

- counsel the Pre-Pubertal Female (PPF) patient on the Adempas risks, including serious birth defects and to immediately contact her prescriber if she begins to menstruate
- Review the Guide for Female Patients with the patient.
- for PPF, regularly assess the reproductive status of each pre-pubertal female during their treatment with Adempas.

Submit this form online at www.adempasREMS.com or fax this form to 1-855-662-5200

Phone: 1-855-4ADEMPAS (1-855-423-3672) Fax: 1-855-662-5200

