



Odevixibat Reimbursement Application Form

For MIMP Use Unity						
Case Reference:		Date Received:				
	1					
Date of Application						
	•	Part 1:	Patient Do	etails		
Name of Patient						
Date of Birth						
Address						
GMS / DPS / PPS Number		GMS	DPS	PPSN		
(Please tick and insert number)	Num	oer:				
	P	art 2: P	rescriber l	Details		
Name of Consultant	Pa	art 2: P	rescriber l	Details		
Name of Consultant Medical Council Number	Pa	art 2: P	rescriber l	Details		
	Pa Hosp		rescriber l	Details		
Medical Council Number		ital:	rescriber I	Details		
Medical Council Number	Hosp	ital:	rescriber I	Details		
Medical Council Number	Hosp	ital:	rescriber I	Details		

Please refer to the HSE Managed Access Protocol for Odevixibat (Bylvay®) when completing Part 3 of this application form

Part 3: Reimbursement criteria - Initiation						
Part 3(a): Patient age, weight and dosage						
To enable a positive recommendation, information relating to the patient's age, weight and dosage must be provided.						
1. This patient	1. This patient is aged six months or older at the time of application: Yes No]
a) Please pi. Patier	a) Please provide: i. Patient's weight in kilograms (kg):					
 3. Please indicate if this patient is currently accessing treatment with Bylvay® via compassionate use / early access programme / ongoing clinical trial: Yes No No If Yes, please complete (a) below. If No, please complete (b) on the next page. a) Please provide details of the patient's current dose and the number of Bylvay® capsules needed to achieve that dose based on the patient's body weight. 						
Patient's body weight						
(kg)	e.g. 40 mcg/kg/day	(mcg)	Body weig < 19.5 kg	ht	Body weig > 19.5 kg	jht
			Strength (mcg)	Number	Strength (mcg)	Number
			600		1200	

If No, please complete (b) below.

b) Please indicate the number of Bylvay[®] capsules needed to achieve the recommended initiation dose of 40 mcg/kg/day based on the patient's body weight.

Please refer to Tables 1 and 2 in section 4.2 of the SmPC.

Patient's body weight (kg)	Recommended initiation dose	Total daily dose (mcg)	Bylvay [®] capsules administered per day			
	(mcg/kg/day)		Body weig < 19.5 kg	ght	Body weig > 19.5 kg	ght
			Strength (mcg)	Number	Strength (mcg)	Number
40		200		400		
			600		1200	

	Part 3(b): Patient diagnosis
	o enable a positive recommendation, evidence relating to the patient's diagnosis must be ovided.
4.	Patient has a confirmed diagnosis of progressive familial intrahepatic cholestasis (PFIC): Yes No
5.	 Please indicate the following: a) Age at presentation with jaundice: week(s) old (0 / zero indicates 'at birth') b) Age at presentation with abnormal liver function tests: week(s) old (0 / zero indicates 'at birth') c) Presence of any extrahepatic manifestations: Yes No If Yes, (presence of extrahepatic manifestations), please provide further information in the space below:

6.	Please provide a copy of the genetic testing report confirming the diagnosis of PFIC:
	Enclosed
7.	Is the subtype of PFIC known? Yes No
	a) If Yes, please provide the subtype of PFIC (e.g. PFIC1, PFIC2, PFIC3):
	PFIC
0	Does the nations have nathologic variations of the ARCR11 gape that prodict complete absence or
О.	Does the patient have pathologic variations of the ABCB11 gene that predict complete absence or lack of function of the bile salt export pump (BSEP) protein (i.e. patients with BSEP3 subtype of
	PFIC2)? Yes No Not known
	a) If Not known , please provide further information in the space below:
Г	
9.	Please provide copies of the following investigations performed to support the diagnosis of PFIC:
	Liver biopsy Enclosed
	Liver function tests (to include serum bile acid level) Enclosed
	Liver ultrasound Enclosed
	a) If No liver biopsy investigation has been performed, please outline the reason(s) in the space
	below:

Part 3(c): Patient clinical status				
To enable a positive recommendation, the status of the patient in relation to the contraindications for treatment and exclusion criteria must be satisfied.				
 10. Does the patient have any contraindications for treatment outlined in the SmPC for Bylvay[®], i.e. hypersensitivity to the active substance or any of the excipients listed in section 6.1 of the SmPC? Yes No				
 11. Does the patient have the following exclusion criteria outlined in section 2.4.1 of the Managed Access Protocol? a) Conditions, medications or surgical procedures that impair either gastrointestinal motility, or enterohepatic circulation of bile acids, including bile salt transport to biliary canaliculi which have the potential to reduce the efficacy of odevixibat: Yes No b) Severe hepatic impairment (Child-Pugh C): Yes No C) Past medical history or ongoing presence of other types of liver disease: Yes No No Decompensated liver disease, coagulopathy, history or presence of clinically significant ascites, variceal haemorrhage, and/or encephalopathy: Yes No 				
Part 3/d)· Patient's m	nedical trea	atment	
Part 3(d): Patient's medical treatment To enable a positive recommendation, information relating to the patient's current medical treatment must be provided. 12. Please provide details of the patient's current medications in the table below:				
Medication	Strength	Dose	Indication	

Space for additional medications if required:		
Part 4: Patient Baseline	e Parameters	
To enable a positive recommendation, evidence relating		s
must be provided, and taken within one month of the da	ate of application.	
Serum bile acid level	Enclosed	
Liver function tests	Enclosed	
Pruritus assessment (e.g. evaluation report / measurement scale of the severity	Enclosed	
(e.g. evaluation report? measurement scale of the severity	or pruntus)	
Additional space for supporting	information if required	

Completed forms should be returned to: Scan the completed form and return via secure email (e.g. HSE email or healthmail) to: mmp@hse.ie

Authorisation of Request		
Signature of		
Prescribing		
Consultant		
Institution		

Data Protection Notice

- The information on this form will be used by the Health Service Executive (HSE) to assess
 the suitability of the items listed to be provided under Section 20 of the Health (Pricing and
 Supply of Medical Goods) Act 2013.
- Details of prescription items dispensed to the named person may be notified to the HSE by the dispensing pharmacist to ensure that the named person receives the items required.
- The named person may access information relating to themselves only, on prescription claims processed in their name by the HSE.
- We may share information with the Department of Health, healthcare practitioners and other healthcare bodies.
- We may also disclose information to other parties if the law requires us to do so.
- The PCRS privacy statement can be located at www.pcrs.ie.