Introduction

This form must be completed and signed by each male patient prior to the initiation of their therapy with Myrin[®] 50 mg and 100 mg tablets (Thalidomide).

The aim of this treatment initiation form is to ensure that patients are fully informed and understand the risk of malformations and even death for the unborn baby associated with the intake of thalidomide.

WARNING: Severe-life threatening birth defects! If Myrin[®] (Thalidomide) is taken during pregnancy it can cause severe birth defects or death to an unborn baby.

Patient declaration

I, the undersigned, Mr.______ born the __/__/___ confirm that:

I will use a condom during intercourse with a woman of childbearing potential	
My female partner is using an effective method of pregnancy prevention	
My female partner is of non-childbearing potential	
Or I am committed to absolute abstinence	
I was informed by my physician about the risks of the treatment with thalidomide and I	
agree with the following statements:	
 I understand that severe birth defects can occur after exposure to Myrin[®] (Thalidomide). 	
 I have been warned by my doctor that any unborn baby has a high risk of birth 	
defects and could also die if a woman is pregnant or become pregnant while taking Myrin [®] (Thalidomide)	
• I was talked by my doctor to never have unprotected sexual contact with women	
who are pregnant or may become pregnant while I am taking Myrin [®] (Thalidomide) and for 1 week after I stopped	
I must inform my doctor if I think that my partner may be pregnant	
 I understand that Myrin[®] (Thalidomide) was prescribed only for me, I will not share 	
it with anyone else	
I will not drink any alcohol while taking Myrin [®]	
 I have read the Myrin[®] (Thalidomide) patient information and understood its content 	
 I understand that I cannot donate semen and blood while taking Myrin[®] 	
(thalidomide) up to 12 weeks after stopping therapy	
 I understand that I have to return any unused Myrin[®] tablets (thalidomide) to the phone and at the and of my treatment. 	
pharmacy at the end of my treatment	

Reporting of any other adverse events

In the instance of any adverse event suspected to be related to the treatment with Myrin[®] (Thalidomide) you must also report it to the Malta Medicines Authority by post or e-mail: ADR reporting/ Sir Temi Zammit Building, Malta Life Sciences Park, San Gwann or on www.medicinesauthority.gov.mt/adrportal using the adverse drug reaction form of the Medicines Authority www.maltamedicinesauthority gov.mt

OR report it to Cherubino Ltd on 21343270 or elaine@cherubino.com.mt

Informed consent form for men treated with Myrin[®] 50 mg and 100 mg tablets (Thalidomide)

Patient Signature

Patient name (first and second name)	Place and Date:
Patient Signature	

Prescriber signature

Prescriber name (first and second name)	Place and Date:
Prescriber Signature	