

Male	
1. Of the need to avoid foetal exposure	Tick
2. To not share the medicinal product with any other person.	Tick
3. That they should not donate blood during treatment (including during dose interruptions) and for at least 7 days following discontinuation of pomalidomide.	Tick
4. That they should return the unused capsules to the pharmacist at the end of treatment	Tick
5. Of the effective contraceptive measures he or his female partner can use.	Tick
6. That pomalidomide is found in semen, so there is a need to use condoms if the sexual partner is pregnant or is a woman of childbearing potential not on effective contraception (even if the man has had a vasectomy) throughout treatment duration, during dose interruption and for at least 7 days after cessation of treatment.	Tick
7. That if his partner becomes pregnant, he should inform his treating prescriber	Tick

**Can you confirm your patient:**

## Pregnancy Prevention

## Prescriber Confirmation

[illegible]

**Patient: please read thoroughly and initial the adjacent box if you agree with the statement**

I understand that severe birth defects may occur with the use of pomalidomide. I have been warned by my prescriber that any unborn baby has a high risk of birth defects and could even die if a woman is pregnant or becomes pregnant while taking pomalidomide.	Patient Initials
I have been told by my prescriber that I must NEVER have unprotected sexual contact with women who are pregnant or may become pregnant, whilst I am taking pomalidomide and for at least 7 days after stopping treatment.	Patient Initials
I understand that pomalidomide passes into human semen. If my partner is pregnant or able to become pregnant, and she doesn't use effective contraception, I must use condoms throughout the duration of my treatment, during dose interruptions and for at least 7 days after I stop pomalidomide even if I have had a vasectomy.	Patient Initials
I understand that if my partner does become pregnant whilst I am taking pomalidomide or within 7 days after I have stopped taking pomalidomide I should inform my prescriber immediately and my partner should be referred to a physician specialised or experienced in teratology for evaluation and advice.	Patient Initials
I understand that pomalidomide will be prescribed ONLY for me. I must not share it with ANYONE	Patient Initials
I have read and understand the Pomalidomide Patient Guide and understand the contents, including the information about other possible important health problems (side effects) associated with the use of pomalidomide.	Patient Initials
I know that I cannot donate blood while taking pomalidomide (including dose interruptions) and for at least 7 days after stopping treatment.	Patient Initials
I know that I cannot donate semen or sperm while taking pomalidomide, during dose interruptions and for at least 7 days after discontinuation of pomalidomide.	Patient Initials
I understand that I must return any unused pomalidomide capsules to my pharmacy at the end of my treatment	Patient Initials
I have been informed about the effective contraceptive methods that my female partner can use.	Patient Initials
I have been informed about the thromboembolic risk and possible requirement to take thromboprophylaxis during treatment with pomalidomide.	Patient Initials
I understand that my prescriber will send or may provide me with a completed 'Prescription Authorisation Form' with each pomalidomide prescription for the pharmacy.	Patient Initials
I understand that the 'Prescription Authorisation Form' contains non-identifiable information about me, which will ensure pomalidomide is dispensed safely. This information may be used by the Marketing Authorisation Holder and the distributor for the product I receive and the Health Products Regulatory Authority (HPRA) to evaluate the safe use of pomalidomide.	Patient Initials

**Patient Confirmation**

I confirm that I understand and will comply with the requirements of the POMALIDOMIDE Pregnancy Prevention Programme, and I agree that my prescriber can initiate my treatment with pomalidomide.

Your personal data is used solely for the purpose of entering you into the Pomalidomide Pregnancy Prevention Programme and is processed by the marketing authorisation holder (MAH) of the pomalidomide product you receive, its third party service providers and any worldwide Affiliates the MAH may have, for the purposes of compliance with the Risk Management Plan legal obligations and for storage purposes. Third party service providers include, for example, the distributor of the pomalidomide product you receive.

Your pharmacist can confirm the details of the MAH for the pomalidomide product you are given and this will also be mentioned on the packaging and package leaflet. Queries on how your personal data will be processed can be directed to the MAH in question, by consulting their publicly available information (e.g. on their website) which details how they process your personal data and provides a contact point for any queries in relation to their use of your personal data.

Patient Signature		Date	DD	MM	YYYY
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**Statement of the interpreter (Where Appropriate)**

I have interpreted the information above to the patient/parent/carer to the best of my ability and in a way in which I believe she/he/they can understand. She/he/they agree to follow the necessary precautions to prevent an unborn child being exposed to pomalidomide.

<b>Signed:</b>		<b>Name: (Print)</b>		<b>Date</b>	DD	MM	YYYY
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**Important Safety Information:**

This is risk minimisation material and is provided as a collaborative project between Accord Healthcare Ireland Ltd., Clonmel Healthcare Ltd., Viatris Limited., AS Grindeks, Teva Pharmaceuticals Ireland and Rowex Ltd. For further information, please refer to the Summary of Product Characteristics (SmPC) for the respective medicinal product from the relevant Marketing Authorisation Holder available at [www.hpra.ie](http://www.hpra.ie)

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Approved by HPRA: 01-DEC-2025