Package leaflet: Information for the patient

Pomalidomide Grindeks 1 mg hard capsules Pomalidomide Grindeks 2 mg hard capsules Pomalidomide Grindeks 3 mg hard capsules Pomalidomide Grindeks 4 mg hard capsules pomalidomide

Pomalidomide Grindeks is expected to cause severe birth defects and may lead to the death of an unbornbaby.

- Do not take this medicine if you are pregnant or could become pregnant.
- You must follow the contraception advice described in this leaflet.

Read all of this leaflet carefully before you start taking this medicine because it contains important information for you.

- Keep this leaflet. You may need to read it again.
- If you have any further questions, ask your doctor or pharmacist.
- This medicine has been prescribed for you only. Do not pass it on to others. It may harm them, even if their signs of illness are the same as yours.
- If you get any side effects, talk to your doctor or pharmacist.
- This includes any possible side effects not listed in this leaflet. See section 4.

What is in this leaflet

- 1. What Pomalidomide Grindeks is and what it is used for
- 2. What you need to know before you take Pomalidomide Grindeks
- 3. How to take Pomalidomide Grindeks
- 4. Possible side effects
- 5. How to store Pomalidomide Grindeks
- 6. Contents of the pack and other information

1. What Pomalidomide Grindeks is and what it is used for

What Pomalidomide Grindeks is

Pomalidomide Grindeks contains the active substance 'pomalidomide'. This medicine is related to thalidomide and belongs to a group of medicines which affect the immune system (the body's natural defences).

What Pomalidomide Grindeks is used for

Pomalidomide Grindeks is used to treat adults with a type of cancer called 'multiple myeloma'.

Pomalidomide Grindeks is either used with:

• **two other medicines** - called 'bortezomib' (a type of chemotherapy medicine) and 'dexamethasone' (an anti-inflammatory medicine) in people who have had at least one other treatment - including lenalidomide.

Or

• **one other medicine** - called 'dexamethasone' in people whose myeloma has become worse, despite having at least two other treatments - including lenalidomide and bortezomib.

What is multiple myeloma

Multiple myeloma is a type of cancer which affects a certain type of white blood cell (called the 'plasma cell'). These cells grow out of control and accumulate in the bone marrow. This results indamage to the bones and kidneys.

Multiple myeloma generally cannot be cured. However, treatment can reduce the signs and symptoms of the disease or make them disappear for a period of time. When this happens, it is called 'response'.

How Pomalidomide Grindeks works

Pomalidomide Grindeks works in a number of different ways:

- by stopping the myeloma cells developing
- by stimulating the immune system to attack the cancer cells
- by stopping the formation of blood vessels supplying the cancer cells.

The benefit of using Pomalidomide Grindeks with bortezomib and dexamethasone

When pomalidomide is used with bortezomib and dexamethasone, in people who have had at least one other treatment, it can stop multiple myeloma getting worse.

On average, pomalidomide when used with bortezomib and dexamethasone stopped multiple myeloma from coming back for up to 11 months - compared with 7 months for those patients who only used bortezomib and dexamethasone.

The benefit of using Pomalidomide Grindeks with dexamethasone

When Pomalidomide Grindeks is used with dexamethasone, in people who have had at least two other treatments, it can stop multiple myeloma getting worse.

On average, Pomalidomide Grindeks when used with dexamethasone stopped multiple myeloma from coming back for up to 4 months - compared with 2 months for those patients who used only dexamethasone.

2. What you need to know before you take Pomalidomide Grindeks

Do not take Pomalidomide Grindeks:

- if you are pregnant or think you may be pregnant or are planning to become pregnant this is because **Pomalidomide Grindeks is expected to be harmful to an unborn child**. (Men and women taking this medicine must read the section "Pregnancy, contraception and breast-feeding information for women and men" below).
- if you are able to become pregnant, unless you follow all the necessary measures to prevent you from becoming pregnant (see "Pregnancy, contraception and breast-feeding information for women and men"). If you are able to become pregnant, your doctor will record with each prescription that the necessary measures have been taken and will provide you with this confirmation.
- if you are allergic to pomalidomide or any of the other ingredients of this medicine (listed in section 6). If you think you may be allergic, ask your doctor for advice.

If you are uncertain whether any of the conditions above apply to you, talk to your doctor, pharmacist or nurse before taking Pomalidomide Grindeks.

Warnings and precautions

Talk to your doctor, pharmacist or nurse before taking Pomalidomide Grindeks if:

- you have ever had blood clots in the past. During the treatment with Pomalidomide Grindeks you have an increased risk of getting blood clots in your veins and arteries. Your doctor may recommend you take additional treatments (e.g. warfarin) or lower the dose of Pomalidomide Grindeks to reduce the chance that you get blood clots.
- you have ever had an allergic reaction such as rash, itching, swelling, feeling dizzy or trouble breathing while taking related medicines called 'thalidomide' or 'lenalidomide'.
- you have had a heart attack, have heart failure, have difficulty breathing, or if you smoke, have high blood pressure or high cholesterol levels.
- you have a high total amount of tumour throughout the body, including your bone marrow. This could lead to a condition where the tumours break down and cause unusual levels of chemicals in the blood which can lead to kidney failure. You may also experience an uneven heartbeat. This condition is called tumour lysis syndrome.
- you have or have had neuropathy (nerve damage causing tingling or pain in your hands or feet).
- you have or have ever had hepatitis B infection. Treatment with Pomalidomide Grindeks may cause the hepatitis B virus to become active again in patients who carry the virus, resulting in a recurrence of the infection. Your doctor should check whether you have ever had hepatitis B infection.

• you experience or have experienced in the past a combination of any of the following symptoms: rash on face or extended rash, red skin, high fever, flu-like symptoms, enlarged lymph nodes (signs of severe skin reaction called Drug Reaction with Eosinophilia and Systemic Symptoms (DRESS) or drug hypersensitivity syndrome, Toxic Epidermal Necrolysis(TEN) or Stevens-Johnson Syndrome (SJS). See also section 4 "Possible side effects".

It is important to note that patients with multiple myeloma treated with pomalidomide may develop additional types of cancer, therefore your doctor should carefully evaluate the benefit and risk whenyou are prescribed this medicine.

At any time during or after your treatment, tell your doctor or nurse immediately if you: experience blurred, loss of or double vision, difficulty speaking, weakness in an arm or a leg, a change in the way you walk or problems with your balance, persistent numbness, decreased sensation or loss of sensation, memory loss or confusion. These may all be symptoms of a serious and potentially fatal brain condition known as progressive multifocal leukoencephalopathy (PML). If you had these symptoms prior to treatment with Pomalidomide Grindeks, tell your doctor about any change in these symptoms.

At the end of the treatment, you should return all unused capsules to the pharmacist.

Pregnancy, contraception and breast-feeding – information for women and men

The following must be followed as stated in the Pomalidomide Grindeks Pregnancy Prevention Programme. Women and men taking Pomalidomide Grindeks must not become pregnant or father a child. This is because pomalidomide is expected to harm the unborn baby. You and your partner should use effective methods of contraception while taking this medicine.

Women

Do not take Pomalidomide Grindeks if you are pregnant, think you may be pregnant or are planning to become pregnant. This is because this medicine is expected to harm the unborn baby. Before starting the treatment, you should tell your doctor if you are able to become pregnant, even if you think this is unlikely.

If you are able to become pregnant:

- you must use effective methods of contraception for at least 4 weeks before starting treatment, for the whole time you are taking treatment, and until at least 4 weeks after stopping treatment. Talk to your doctor about the best method of contraception for you.
- each time your doctor writes a prescription for you, he will ensure you understand the necessary measures that have to be taken to prevent pregnancy.
- your doctor will arrange pregnancy tests before treatment, at least every 4 weeks during treatment, and at least 4 weeks after the treatment has finished.

If you become pregnant despite the prevention measures:

• you must stop the treatment immediately and talk to your doctor straight away.

Breast-feeding

It is not known if Pomalidomide Grindeks passes into human breast milk. Tell your doctor if you are breast-feeding orintend to breast-feed. Your doctor will advise if you should stop or continue breast-feeding.

Men

Pomalidomide Grindeks passes into human semen.

- If your partner is pregnant or able to become pregnant, you must use condoms for the whole time you are taking treatment and for 7 days after the end of treatment.
- If your partner becomes pregnant while you are taking Pomalidomide Grindeks, tell your doctor straight away. Your partner should also tell her doctor straight away.

You should not donate semen or sperm during treatment and for 7 days after the end of treatment.

Blood donation and blood tests

You should not donate blood during treatment and for 7 days after the end of treatment.

Before and during the treatment with Pomalidomide Grindeks you will have regular blood tests. This is because your medicine may cause a fall in the number of blood cells that help fight infection (white cells) and in the number of cells that help to stop bleeding (platelets).

Your doctor should ask you to have a blood test:

- before treatment
- every week for the first 8 weeks of treatment
- at least every month after that for as long as you are taking Pomalidomide Grindeks.

As a result of these tests, your doctor may change your dose of Pomalidomide Grindeks or stop your treatment. The doctor may also change the dose or stop the medicine because of your general health.

Children and adolescents

Pomalidomide Grindeks is not recommended for use in children and young people under 18 years.

Other medicines and Pomalidomide Grindeks

Tell your doctor, pharmacist or nurse if you are taking, have recently taken or might take any other medicines. This is because Pomalidomide Grindeks can affect the way some other medicines work. Also, some other medicines can affect the way Pomalidomide Grindeks works.

In particular, tell your doctor, pharmacist or nurse before taking Pomalidomide Grindeks if you are taking any of the following medicines:

- some antifungals such as ketaconazole
- some antibiotics (for example ciprofloxacin, enoxacin)
- certain antidepressants such as fluvoxamine.

Driving and using machines

Some people feel tired, dizzy, faint, confused or less alert when taking Pomalidomide Grindeks. If this happens to you,do not drive or operate tools or machinery.

Pomalidomide Grindeks contains sodium

This medicine contains less than 1 mmol sodium (23 mg) per capsule, therefore it is considered essentially 'sodium-free'.

Pomalidomide Grindeks contains azo colouring agents

The capsules contain the azo colouring agents Brilliant Black PN (all strengths), Azorubine – Carmoisine (all strengths) and Sunset yellow FCF (2 mg capsules only). These colouring agents may cause allergic reactions.

3. How to take Pomalidomide Grindeks

Pomalidomide Grindeks must be given to you by a doctor with experience in treating multiple myeloma.

Always take your medicines exactly as your doctor has told you. Check with your doctor, pharmacist or nurse if you are not sure.

When to take Pomalidomide Grindeks with other medicines

Pomalidomide Grindeks with bortezomib and dexamethasone

- See the leaflets that come with bortezomib and dexamethasone for further information on their use and effects.
- Pomalidomide Grindeks, bortezomib and dexamethasone are taken in 'treatment cycles'. Each cycle lasts 21 days (3 weeks).

- Look at the chart below to see what to take on each day of the 3-week cycle.
 - o Each day, look down the chart and find the correct day to see which medicines to take.
 - O Some days, you take all 3 medicines, some days just 2 or 1 medicines, and some days none at all.

PMD: Pomalidomide Grindeks; BOR: Bortezomib; DEX: Dexamethasone
Cycle 1 to 8

Cycle 9 and onwards

	Medicine name		
Day	PMD	BOR	DEX
1	V	1	V
3 4	V		V
3			
	V	√	
5	V		
6	V		
7	V		
8	V		
9	V		
10	V		
11	V		
12	V		
13			
14			
15			
16			
17			
18	_		
19	_		
20			
21			

	Medicine name		
Day	PMD	BOR	DEX
1	√	1	V
2	$\sqrt{}$		$\sqrt{}$
3	$\sqrt{}$		
4	$\sqrt{}$		
5	$\sqrt{}$		
6	$\sqrt{}$		
7	$\sqrt{}$		
8	$\sqrt{}$	$\sqrt{}$	$\sqrt{}$
9	$\sqrt{}$		$\sqrt{}$
10	$\sqrt{}$		
11	$\sqrt{}$		
12	$\sqrt{}$		
13	$\sqrt{}$		
14			
15			
16			
17			
18			
19			
20			
21			

After completing each 3-week cycle, start a new one.

Pomalidomide Grindeks with dexamethasone only

- See the leaflet that comes with dexamethasone for further information on its use and effects.
- Pomalidomide Grindeks and dexamethasone are taken in 'treatment cycles'. Each cycle lasts 28 days (4 weeks).
- Look at the chart below to see what to take on each day of the 4-week cycle:
 - o Each day, look down the chart and find the correct day to see which medicines to take.
 - O Some days, you take both medicines, some days just 1 medicine, and some days none at all.

PMD: Pomalidomide Grindeks; DEX: Dexamethasone

	Medicine name	
Day	PMD	DEX
1	V	$\sqrt{}$
2	V	
3	V	
4	V	
5	V	
6	V	
7	V	
8	V	V
9	V	

10	V	
11	V	
12	V	
13	V	
14	V	
15	V	V
16	V	
17	V	
18	V	
19	V	
20	V	
21	V	
22		V
23		
24		
25		
26		
27		
28		

After completing each 4-week cycle, start a new one.

How much Pomalidomide Grindeks to take with other medicines

Pomalidomide Grindeks with bortezomib and dexamethasone

- The recommended starting dose of Pomalidomide Grindeks is 4 mg per day.
- The recommended starting dose of bortezomib will be worked out by your doctor and based on your height and weight (1.3 mg/m² body surface area).
- The recommended starting dose of dexamethasone is 20 mg per day. However, if you are over 75, the recommended starting dose is 10 mg per day.

Pomalidomide Grindeks with dexamethasone only

- The recommended dose of Pomalidomide Grindeks is 4 mg per day.
- The recommended starting dose of dexamethasone is 40 mg per day. However, if you are over 75, the recommended starting dose is 20 mg per day.

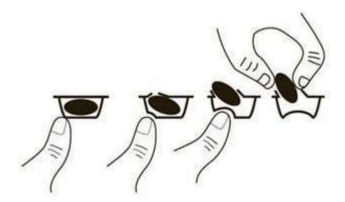
Your doctor may need to reduce the dose of Pomalidomide Grindeks, bortezomib or dexamethasone or stop one or more of these medicines based on the results of your blood tests, your general condition, other medicines you may be taking (e.g. ciprofloxacin, enoxacin and fluvoxamine) and if you experience side effects (especially rash or swelling) from treatment.

If you suffer from liver or kidney problems your doctor will check your condition very carefully whilst you are receiving this medicine.

How to take Pomalidomide Grindeks

- Do not break, open or chew the capsules. If powder from a broken Pomalidomide Grindeks capsule makes contact with the skin, wash the skin immediately and thoroughly with soap and water.
- Healthcare professionals, caregivers and family members should wear disposable gloves when handling the blister or capsule. Gloves should then be removed carefully to prevent skin exposure, placed in a sealable plastic polyethylene bag and disposed of in accordance with local requirements. Hands should then be washed thoroughly with soap and water. Women who are pregnant or suspect they may be pregnant should not handle the blister or capsule.
- Swallow the capsules whole, preferably with water.
- You can take the capsules either with or without food.
- Take Pomalidomide Grindeks at about the same time each day.

To remove the capsule from the blister, press only one end of the capsule out to push it through the foil. Do not apply pressure on the centre of the capsule as this can cause it to break.



Your doctor will advise you of how and when to take Pomalidomide Grindeks if you have kidney problems and are receiving dialysis treatment.

Duration of the treatment with Pomalidomide Grindeks

You should continue the cycles of treatment until your doctor tells you to stop.

If you take more Pomalidomide Grindeks than you should

If you take more Pomalidomide Grindeks than you should, talk to a doctor or go to a hospital straight away. Take the medicine pack with you.

If you forget to take Pomalidomide Grindeks

If you forget to take Pomalidomide Grindeks on a day when you should, take your next capsule as normal the next day. Do not increase the number of capsules you take to make up for not taking Pomalidomide Grindeks the previous day.

If you have any further questions on the use of this medicine, ask your doctor or pharmacist.

4. Possible side effects

Like all medicines, this medicine can cause side effects, although not everybody gets them.

Serious side effects

Stop taking Pomalidomide Grindeks and see a doctor straight away if you notice any of the following serious side effects – you may need urgent medical treatment:

- Fever, chills, sore throat, cough, mouth ulcers or any other signs of infection (due to less white blood cells, which fight infection).
- Bleeding or bruising without a cause, including nosebleeds and bleeding from the bowels or stomach (due to effects on blood cells called 'platelets').
- Rapid breathing, rapid pulse, fever and chills, passing very little to no urine, nausea and vomiting, confusion, unconsciousness (due to infection of blood called sepsis or septic shock).
- Severe, persistent or bloody diarrhoea (possibly with stomach pain or fever) caused by bacteria called *Clostridium difficile*.
- Chest pain, or leg pain and swelling, especially in your lower leg or calves (caused by blood clots).
- Shortness of breath (from serious chest infection, inflammation of the lung, heart failure or blood clot).
- Swelling of face, lips, tongue and throat, which may cause difficulty breathing (due to serious types of allergic reaction called angioedema and anaphylactic reaction).
- Certain types of skin cancer (squamous cell carcinoma and basal cell carcinoma), which can cause changes in the appearance of your skin or growths on your skin. If you notice any changes to your

- skin whilst taking Pomalidomide Grindeks, tell your doctor as soon as possible.
- Recurrence of hepatitis B infection, which can cause yellowing of the skin and eyes, dark, brown-coloured urine, right-sided abdominal pain, fever and feeling nauseous or being sick. Tell your doctor straightaway if you notice any of these symptoms.
- Widespread rash, high body temperature, enlarged lymph nodes and other body organs involvement (Drug Reaction with Eosinophilia and Systemic Symptoms which is also known as DRESS or drug hypersensitivity syndrome, Toxic Epidermal Necrolysis or Stevens-Johnson Syndrome). Stop using pomalidomide if you develop these symptoms and contact your doctor or seek medical attention immediately. See also section 2.

Stop taking Pomalidomide Grindeks and see a doctor straight away if you notice any of the serious side effects listed above – you may need urgent medical treatment.

Other side effects

Very common (may affect more than 1 in 10 people):

- Shortness of breath (dyspnoea).
- Infections of the lungs (pneumonia and bronchitis).
- Infections of the nose, sinuses, and throat, caused by bacteria or viruses.
- Flu-like symptoms (influenza).
- Low red blood cells, which may cause anaemia leading to tiredness and weakness.
- Low blood levels of potassium (hypokalaemia), which may cause weakness, muscle cramps, muscle aches, palpitations, tingling or numbness, dyspnoea, mood changes.
- High blood levels of sugar.
- A fast and irregular heartbeat (atrial fibrillation).
- Loss of appetite.
- Constipation, diarrhoea, or nausea.
- Being sick (vomiting).
- Abdominal pain.
- Lack of energy.
- Difficulty in falling asleep or staying asleep.
- Dizziness, tremor.
- Muscle spasm, muscle weakness.
- Bone pain, back pain.
- Numbness, tingling or burning sensation to the skin, pains in hands or feet (peripheral sensory neuropathy).
- Swelling of the body, including swelling of the arms or legs.
- Rashes.
- Urinary tract infection, which may cause a burning sensation when passing urine, or a need to pass urine more often.

Common (may affect up to 1 in 10 people):

- Fall.
- Bleeding within the skull.
- Decreased ability to move or feel (sensation) in your hands, arms, feet and legs because of nerve damage (peripheral sensorimotor neuropathy).
- Numbness, itching, and a feeling of pins and needles on your skin (paraesthesia).
- A spinning feeling in your head, making it difficult to stand up and move normally.
- Swelling caused by fluid.
- Hives (urticaria).
- Itchy skin.
- Shingles.
- Heart attack (chest pain spreading to the arms, neck, jaw, feeling sweaty and breathless, feeling sick or vomiting).
- Chest pain, chest infection.
- Increased blood pressure.

- A fall in the number of red and white blood cells and platelets at the same time (pancytopenia), which will make you more prone to bleeding and bruising. You may feel tired and weak, and short of breath and you are also more likely to get infections.
- Decreased number of lymphocytes (one type of white blood cells) often caused by infection (lymphopenia).
- Low blood levels of magnesium (hypomagnesaemia), which may cause tiredness, generalised weakness, muscle cramps, irritability and may result in low blood levels of calcium (hypocalcaemia), which may cause numbness and, or tingling of hands, feet, or lips, muscle cramps, muscle weakness, light-headedness, confusion.
- Low blood level of phosphate (hypophosphataemia), which may cause muscle weakness and irritability or confusion.
- High blood level of calcium (hypercalcaemia), which may cause slowing reflexes and skeletal muscle weaknesses.
- High blood levels of potassium, which may cause abnormal heart rhythm.
- Low blood levels of sodium, which may cause tiredness and confusion, muscle twitching, fits (epileptic seizures) or coma.
- High blood levels of uric acid, which may cause a form of arthritis called gout.
- Low blood pressure, which may cause dizziness or fainting.
- Sore or dry mouth.
- Changes in the way things taste.
- Swollen abdomen.
- Feeling confused.
- Feeling down (depressed mood).
- Loss of consciousness, fainting.
- Clouding of your eye (cataract).
- Damage to the kidney.
- Inability to pass urine.
- Abnormal liver test.
- Pain in the pelvis.
- Weight loss.

Uncommon (may affect up to 1 in 100 people):

- Stroke.
- Inflammation of the liver (hepatitis) which can cause itchy skin, yellowing of the skin and the whites of the eyes (jaundice), pale coloured stools, dark coloured urine and abdominal pain.
- The breakdown of cancer cells resulting in the release of toxic compounds into the bloodstream (tumour lysis syndrome). This can result in kidney problems.
- Underactive thyroid gland, which may cause symptoms such as tiredness, lethargy, muscle weakness, slow heart rate, weight gain.

Not known (frequency cannot be estimated from the available data):

• Rejection of solid organ transplant (such as heart or liver).

Reporting of side effects

If you get any side effects, talk to your doctor, pharmacist, or nurse. This includes any possible side effects not listed in this leaflet. You can also report side effects directly:

Website: www.hpra.ie

HPRA Pharmacovigilance

By reporting side effects you can help provide more information on the safety of this medicine.

5. How to store Pomalidomide Grindeks

Keep this medicine out of the sight and reach of children.

Do not use this medicine after the expiry date which is stated on the blister and carton after EXP. The expiry date refers to the last day of that month.

This medicine does not require any special storage conditions.

Do not use Pomalidomide Grindeks if you notice any damage or signs of tampering to medicine packaging.

Do not throw away any medicines via wastewater or household waste. Any unused medicine should be returned to the pharmacist at the end of treatment. These measures will help protect the environment.

6. Contents of the pack and other information

What Pomalidomide Grindeks contains

Pomalidomide Grindeks 1 mg hard capsules:

- The active substance is pomalidomide. Each capsule contains 1 mg pomalidomide.
- The other ingredients are: pregelatinised starch; maltodextrin; crospovidone; silica, colloidal anhydrous; sodium stearyl fumarate.
- The capsule shells contain: gelatine; titanium dioxide (E171); colourants (yellow iron oxide (E172); black iron oxide (E172); brilliant black PN (E151); patent blue V (E131); azorubine carmoisine (E122); brilliant blue FCF (E133)) and white printing ink (shellac; titanium dioxide (E171); sodium hydroxide; propylene glycol (E1520) and povidone (E1201)).

Pomalidomide Grindeks 2 mg hard capsules:

- The active substance is pomalidomide. Each capsule contains 2 mg pomalidomide.
- The other ingredients are: pregelatinised starch; maltodextrin; crospovidone; silica, colloidal anhydrous; sodium stearyl fumarate.
- The capsule shells contain: gelatine; titanium dioxide (E171); colourants (sunset yellow FCF (E110); brilliant black PN (E151); patent blue V (E131); azorubine carmoisine (E122)) and white printing ink (shellac; titanium dioxide (E171); sodium hydroxide; propylene glycol (E1520) and povidone (E1201)).

Pomalidomide Grindeks 3 mg hard capsules:

- The active substance is pomalidomide. Each capsule contains 3 mg pomalidomide.
- The other ingredients are: pregelatinised starch; maltodextrin; crospovidone; silica, colloidal anhydrous; sodium stearyl fumarate.
- The capsule shells contain: gelatine; titanium dioxide (E171); colourants (brilliant black PN (E151); patent blue V (E131); azorubine carmoisine (E122); brilliant blue FCF (E133); erythrosine (E127)) and white printing ink (shellac; titanium dioxide (E171); sodium hydroxide; propylene glycol (E1520) and povidone (E1201)).

Pomalidomide Grindeks 4 mg hard capsules:

- The active substance is pomalidomide. Each capsule contains 4 mg pomalidomide.
- The other ingredients are: pregelatinised starch; maltodextrin; crospovidone; silica, colloidal anhydrous; sodium stearyl fumarate.
- The capsule shells contain: gelatine; titanium dioxide (E171); colourants (brilliant blue FCF (E133); brilliant black PN (E151); patent blue V (E131); azorubine carmoisine (E122); erythrosine (E127)) and white printing ink (shellac, titanium dioxide (E171); sodium hydroxide; propylene glycol (E1520) and povidone (E1201)).

What Pomalidomide Grindeks looks like and contents of the pack

Pomalidomide Grindeks 1 mg hard capsules are size 4 (approximately 14 mm × 5 mm), hard gelatine capsules with light grey body imprinted with P1 in white ink and dark blue opaque cap.

Pomalidomide Grindeks 2 mg hard capsules are size 3 (approximately 16 mm × 6 mm), hard gelatine

capsules with orange opaque body imprinted with P2 in white ink and dark blue opaque cap.

Pomalidomide Grindeks 3 mg hard capsules are size 2 (approximately 18 mm × 6 mm), hard gelatine capsules with light blue body imprinted with P3 in white ink and dark blue opaque cap.

Pomalidomide Grindeks 4 mg hard capsules are size 1 (approximately 19 mm \times 7 mm), hard gelatine capsules with blue opaque body imprinted with P4 in white ink and dark blue opaque cap.

The capsules are provided in blister packs of 14 or 21 capsules (2 or 3 blisters per pack, with 7 capsules in each blister) or perforated unit dose blister packs of 14 x 1 or 21 x 1 capsules.

Not all pack sizes may be marketed.

Marketing Authorisation Holder and manufacturer

AS GRINDEKS Krustpils iela 53, Rīga, LV-1057, Latvia

This medicine is authorised in the Member States of the European Economic Area under the following names:

Austria Pomalidomid Grindeks 1 mg, 2 mg, 3 mg, 4 mg Hartkapseln Belgium Pomalidomide Grindeks 1 mg, 2 mg, 3 mg, 4 mg gélules

Bulgaria Помалидомид Гриндекс 1 mg, 2 mg, 3 mg, 4 mg твърди капсули

Pomalidomide Grindeks 1 mg, 2 mg, 3 mg, 4 mg hard capsules

Czech RepublicPomalidomide GrindeksDenmarkPomalidomid GrindeksEstoniaPomalidomide Grindeks

Finland Pomalidomide Grindeks 1 mg, 2 mg, 3 mg, 4 mg kovat kapselit

France POMALIDOMIDE GRINDEKS 1 mg, gélule POMALIDOMIDE GRINDEKS 2 mg, gélule

POMALIDOMIDE GRINDEKS 2 mg, gélule POMALIDOMIDE GRINDEKS 3 mg, gélule POMALIDOMIDE GRINDEKS 4 mg, gélule

Germany Pomalidomid Grindeks 1 mg, 2 mg, 3 mg, 4 mg Hartkapseln

Greece Pomalidomide/Grindeks

Hungary Pomalidomide Grindeks 1 mg, 2 mg, 3 mg, 4 mg kemény kapszula Ireland Pomalidomide Grindeks 1 mg, 2 mg, 3 mg, 4 mg hard capsule

Italy Pomalidomide Grindeks

Latvia Pomalidomide Grindeks 1 mg, 2 mg, 3 mg, 4 mg cietās kapsulas Lithuania Pomalidomide Grindeks 1 mg, 2 mg, 3 mg, 4 mg kietosios kapsulės

The Netherlands Pomalidomide Grindeks 1 mg harde capsules

Pomalidomide Grindeks 2 mg harde capsules Pomalidomide Grindeks 3 mg harde capsules Pomalidomide Grindeks 4 mg harde capsules

Norway Pomalidomide Grindeks Poland Pomalidomide Grindeks

Portugal Pomalidomida Grindeks 1 mg, 2 mg, 3 mg, 4 mg cápsula

Romania Pomalidomidă Grindeks 1 mg capsule

Pomalidomidă Grindeks 2 mg capsule Pomalidomidă Grindeks 3 mg capsule Pomalidomidă Grindeks 4 mg capsule

Slovenia Pomalidomid Grindeks 1 mg, 2 mg, 3 mg, 4 mg trde kapsule

Slovakia Pomalidomid Grindeks 1 mg tvrdé kapsuly

Pomalidomid Grindeks 2 mg tvrdé kapsuly

Pomalidomid Grindeks 3 mg tvrdé kapsuly

Pomalidomid Grindeks 4 mg tvrdé kapsuly Pomalidomide Grindeks 1 mg, 2 mg, 3 mg, 4 mg cápsula dura Spain Pomalidomide Grindeks 1 mg, 2 mg, 3 mg, 4 mg hårda kapslar Sweden

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