ANNEX I SUMMARY OF PRODUCT CHARACTERISTICS

This medicinal product is subject to additional monitoring. This will allow quick identification of new safety information. Healthcare professionals are asked to report any suspected adverse reactions. See section 4.8 for how to report adverse reactions.

1. NAME OF THE MEDICINAL PRODUCT

Zokinvy 50 mg hard capsules

Zokinvy 75 mg hard capsules

2. QUALITATIVE AND QUANTITATIVE COMPOSITION

Zokinvy 50 mg hard capsules

Each capsule contains 50 mg lonafarnib.

Zokinvy 75 mg hard capsules

Each capsule contains 75 mg lonafarnib.

For the full list of excipients, see section 6.1.

3. PHARMACEUTICAL FORM

Hard capsule (capsule)

Zokinvy 50 mg hard capsules

Size 4 hard capsule (5 mm x 14 mm), opaque yellow with "LNF" and "50" printed in black.

Zokinvy 75 mg hard capsules

Size 3 hard capsule (6 mm x 16 mm), opaque light orange with "LNF and "75" printed in black.

4. CLINICAL PARTICULARS

4.1 Therapeutic indications

Zokinvy is indicated for the treatment of patients 12 months of age and older with a genetically confirmed diagnosis of Hutchinson-Gilford progeria syndrome or a processing-deficient progeroid laminopathy associated with either a heterozygous *LMNA* mutation with progerin-like protein accumulation or a homozygous or compound heterozygous *ZMPSTE24* mutation.

4.2 Posology and method of administration

Treatment should be initiated by a physician experienced in the treatment of patients with progeroid syndromes or patients with rare genetic metabolic syndromes.

Posology

Starting dose

For all indications, the recommended starting dose is 115 mg/m² twice daily. The Du Bois formula was used in clinical trials and should be used to calculate body surface area for dosing. All total daily

doses should be rounded to the nearest 25 mg increment and divided into two equal, or near equal, doses (see Table 1). Doses should be taken approximately 12 hours apart from one another (morning and evening).

Table 1: Recommended starting dose and administration schedule for 115 mg/m² body surface

area-based dosing

Body surface	Total daily	Morning dose		Evening dose	
area (m²)	dose rounded	number of capsule(s)		number of capsule(s)	
	to nearest	lonafarnib	lonafarnib lonafarnib		lonafarnib
	25 mg	50 mg	75 mg	50 mg	75 mg
0.30 - 0.38	75		1*		1*
0.39 - 0.48	100	1		1	
0.49 - 0.59	125		1	1	
0.6 - 0.7	150		1		1
0.71 - 0.81	175	2			1
0.82 - 0.92	200	2		2	
0.93 - 1	225	1	1	2	

^{*} For patients with a body surface area of 0.30 m² to 0.38 m², the contents of a 75 mg capsule must be mixed with 10 mL of orange juice. Half of the mixture (5 mL) equates to a 37.5 mg dose of lonafarnib. This dose will be prepared and consumed twice daily (see section 6.6).

Maintenance dose

After 4 months of treatment using the starting dose of 115 mg/m² twice daily, the dose should be increased to the maintenance dose of 150 mg/m² twice daily (morning and evening). All total daily doses should be rounded to the nearest 25 mg increment and divided into two equal, or near equal, doses (see Table 2).

Table 2: Recommended maintenance dose and administration schedule for 150 mg/m² body

surface area-based dosing

Body surface area (m ²)	Total daily dose rounded	Morning dose number of capsule(s)		Evening dose number of capsule(s)	
, ,	to nearest 25 mg	lonafarnib 50 mg	lonafarnib 75 mg	lonafarnib 50 mg	lonafarnib 75 mg
0.30 - 0.37	100	1		1	
0.38 - 0.45	125		1	1	
0.46 - 0.54	150		1		1
0.55 - 0.62	175	2			1
0.63 - 0.7	200	2		2	
0.71 - 0.79	225	1	1	2	
0.8 - 0.87	250	1	1	1	1
0.88 - 0.95	275		2	1	1
0.96 - 1	300		2		2

Missed dose

If a dose is missed, the dose should be taken as soon as possible, up to 8 hours prior to the next scheduled dose with food. If less than 8 hours remain before the next scheduled dose, the missed dose should be skipped and the dose regimen should be resumed at the next scheduled dose.

Patients taking the starting dose of 115 mg/m² with a body surface area of 0.30 m² to 0.38 m² Patients will need to receive a daily dose of 75 mg (37.5 mg twice daily). The contents of a lonafarnib 75 mg capsule should be mixed with 10 mL of orange juice. Only half of the 10-mL mixture will be consumed (see section 6.6).

Dose adjustment for patients with persistent vomiting and/or diarrhoea leading to dehydration or weight loss

For patients who have increased their dose to 150 mg/m² twice daily and are experiencing repeated episodes of vomiting and/or diarrhoea resulting in dehydration or weight loss (see section 4.4), the dose can be reduced to the starting dose of 115 mg/m² twice daily. All daily doses should be rounded to the nearest 25 mg increment and divided into two equal, or near equal, doses (see Table 1).

Prevention or treatment of vomiting and/or diarrhoea leading to dehydration or weight loss Prevention or treatment of vomiting and/or diarrhoea with an anti-emetic and/or anti-diarrhoeal medicinal product can be considered (see section 4.4).

Dose adjustment for patients who cannot avoid taking a concomitant moderate CYP3A inhibitor (see section 4.5)

The patient's daily dose of lonafarnib should be reduced by 50% and the reduced daily dose should be divided into two equal doses. Each dose should be rounded to the nearest 25 mg increment. The dosing regimen will be either 25 mg twice daily, 50 mg twice daily or 75 mg twice daily. Patients who have a reduced daily dose of 50 mg (25 mg twice daily) should mix the contents of a lonafarnib 50 mg capsule with 10 mL of orange juice to achieve the correct dose. Only half (5 mL) of the 10-mL mixture will be consumed (see section 6.6). QTc monitoring is recommended while the patient is taking a concomitant moderate CYP3A inhibitor and being treated with 50% of the indicated dose of lonafarnib. The patient should resume the body surface area indicated dose of lonafarnib 14 days after discontinuation of the moderate CYP3A inhibitor.

Dose adjustment for patients taking a concomitant weak CYP3A inhibitor and experiencing a persistent toxicity (see section 4.5)

The patient's daily dose of lonafarnib should be reduced by 50% and the reduced daily dose should be divided into two equal doses. Each dose should be rounded to the nearest 25 mg increment. The dosing regimen will be either 25 mg twice daily, 50 mg twice daily or 75 mg twice daily. Patients who have a reduced daily dose of 50 mg (25 mg twice daily) should mix the contents of a lonafarnib 50 mg capsule with 10 mL of orange juice to achieve the correct dose. Only half (5 mL) of the 10-mL mixture will be consumed (see section 6.6). QTc monitoring is recommended while the patient is taking a concomitant weak CYP3A inhibitor and being treated with 50% of the indicated dose of lonafarnib due to the presence of a persistent toxicity. The patient should resume the body surface area indicated dose of lonafarnib 14 days after the toxicity has fully resolved or discontinuation of the weak CYP3A inhibitor.

Dose adjustment for patients with known dysfunctional polymorphisms in CYP3A4 The patient's daily dose of lonafarnib should be reduced by 50% and the reduced daily dose should be divided into two equal doses. Each dose should be rounded to the nearest 25 mg increment. The dosing regimen will be either 25 mg twice daily, 50 mg twice daily or 75 mg twice daily. Patients who have a reduced daily dose of 50 mg (25 mg twice daily) should mix the contents of a lonafarnib 50 mg capsule with 10 mL of orange juice to achieve the correct dose. Only half (5 mL) of the 10-mL mixture will be consumed (see section 6.6). QTc monitoring is recommended.

Dose adjustment for patients requiring parenteral midazolam for a surgical procedure Concomitant use of midazolam is contraindicated (see sections 4.3 and 4.5). Patients requiring parenteral midazolam for a surgical procedure should discontinue lonafarnib for 14 days before and 2 days after administration of midazolam.

Specific interactions with foods and drinks

Lonafarnib should not be taken with foods or juices that contain grapefruit, cranberries, pomegranates or Seville oranges (*e.g.*, orange marmalade), otherwise known as sour or bitter oranges (see section 4.5). Taking lonafarnib with food or drinks containing these fruits or fruit juices may increase adverse reactions associated with lonafarnib.

Special populations

Patients with hepatic impairment

No dose adjustments are required in patients with mild or moderate hepatic impairment (Child-Pugh Class A or B, respectively). Lonafarnib is contraindicated in patients with severe hepatic impairment (Child-Pugh Class C) (see sections 4.3, 4.4 and 5.2).

Patients with renal impairment

Lonafarnib has not been studied in patients with renal impairment. Because lonafarnib and metabolite HM21 are only excreted to a limited extent via urine, no dose adjustments are required in patients with renal impairment (see sections 4.4 and 5.2).

Paediatric population

The posology is the same in adults and children 12 months of age and older.

The safety and efficacy of lonafarnib in children less than 12 months of age have not been established. No data are available (see section 5.1).

Method of administration

Lonafarnib is intended for oral use. The capsule should be swallowed whole. The capsule should not be chewed. Each dose is to be taken with food.

For patients unable to swallow the capsule whole, instructions on mixing the capsule contents with orange juice are provided in section 6.6.

4.3 Contraindications

Hypersensitivity to the active substance or any other member of the farnesyltransferase class, or to any of the excipients listed in section 6.1.

Concomitant use with strong CYP3A inhibitors (see section 4.5).

Concomitant use of medicinal products that are predominantly metabolised by CYP3A4, such as midazolam, atorvastatin, lovastatin and simvastatin (see sections 4.2, 4.4 and 4.5).

Patients with severe hepatic impairment (Child-Pugh Class C) (see section 5.2).

4.4 Special warnings and precautions for use

Age at start of treatment

Treatment with lonafarnib should be initiated as soon as a diagnosis has been made. The clinical data indicate that the expected survival benefit of lonafarnib treatment in Hutchinson-Gilford progeria syndrome (HGPS) patients who started treatment at 10 years of age or above is less compared to those who started at a younger age (see section 5.1).

Treatment initiation with lonafarnib in older patients should be balanced against the side effects (*i.e.*, vomiting, nausea and diarrhoea) in the first few months of treatment.

Gastrointestinal adverse reactions and dehydration

Electrolyte abnormalities (hypermagnesaemia, hypokalaemia, hyponatraemia) have been reported (see section 4.8). The severity of gastrointestinal adverse reactions, especially during the first 4 months of treatment, should be closely monitored. When gastrointestinal adverse reactions occur, monitoring the patient's weight, caloric consumption and fluid volume intake should be done on a regular basis. In some cases, persistent diarrhoea can result in hypovolaemia, which should be treated by infusion or orally.

Patients experiencing diarrhoea and treated with the anti-diarrhoeal loperamide should be monitored for adverse reactions associated with increased exposure to loperamide (see section 4.5).

Patients requiring parenteral midazolam for a surgical procedure

Concomitant administration of lonafarnib and midazolam is contraindicated (see sections 4.3 and 4.5) due to an increased risk of extreme sedation and respiratory depression. For patients requiring midazolam as a component of anaesthesia for a surgical procedure, lonafarnib treatment should be discontinued for 14 days before and 2 days after parenteral midazolam is administered.

Abnormal liver function

Increased liver enzymes, such as aspartate aminotransferase or alanine aminotransferase, have been reported (see section 4.8). Signs and symptoms of reduced liver function should be assessed on a consistent basis. Liver function should be measured annually or at the onset of any new or worsening signs or symptoms of liver dysfunction.

Nephrotoxicity

Lonafarnib caused nephrotoxicity in rats with clinical chemistry and urinalysis changes, at plasma exposures approximately equal to the human dose (see section 5.3). Signs and symptoms of reduced renal function should be assessed on a consistent basis. Renal function should be measured annually or at the onset of any new or worsening signs or symptoms associated with renal dysfunction.

Retinal toxicity

Lonafarnib caused rod-dependent, low-light vision decline in monkeys at plasma exposures similar to the human dose (see section 5.3). An ophthalmological evaluation should be performed annually and at the onset of any new visual disturbances during therapy.

Concomitant use of moderate and strong CYP3A inducers

Concomitant use of moderate and strong CYP3A inducers may reduce the efficacy of lonafarnib and they should be avoided (see section 4.5).

Concomitant use of moderate CYP3A inhibitors

Concomitant use of lonafarnib and moderate CYP3A inhibitors should be avoided. If concomitant use is unavoidable, the dose of lonafarnib should be reduced by 50% and QTc monitoring is recommended (see sections 4.2 and 4.5).

Concomitant use of weak CYP3A inducers

Concomitant use of weak CYP3A inducers may reduce the efficacy of lonafarnib and should be avoided. If their use is unavoidable, no dose adjustment of lonafarnib is needed (see section 4.5).

Subjects with known dysfunctional polymorphisms in CYP3A4

Subjects with a known dysfunctional polymorphism in CYP3A4 should start therapy at 50% of the indicated dose. QTc monitoring is necessary (see section 4.2 and 4.5).

Other progeroid syndromes

Lonafarnib is not expected to be effective for the treatment of progeroid syndromes caused by mutations in genes other than *LMNA* or *ZMPSTE24* and laminopathies not associated with the accumulation of progerin-like proteins. Lonafarnib is not expected to be effective in the treatment of the following progeroid syndromes: Werner syndrome, Bloom syndrome, Rothmund–Thomson

syndrome, Cockayne syndrome, xeroderma pigmentosum, trichothiodystrophy and ataxiatelangiectasia.

Excipients with known effect

Zokinvy contains less than 1 mmol sodium (23 mg) per dose, that is to say essentially 'sodium-free'.

4.5 Interaction with other medicinal products and other forms of interaction

Strong CYP3A inhibitors

When lonafarnib was co-administered with ketoconazole, a strong CYP3A inhibitor, in healthy adult subjects, ketoconazole (200 mg for 5 doses) increased lonafarnib (single dose of 50 mg) C_{max} by 270% and AUC by 425%. This may lead to an increased risk of adverse reactions. Therefore, concomitant use of lonafarnib and strong CYP3A inhibitors is contraindicated (see section 4.3).

Moderate CYP3A inhibitors

No interaction studies have been conducted with a moderate CYP3A inhibitor. Concomitant use of lonafarnib and a moderate CYP3A inhibitor should be avoided (see sections 4.2 and 4.4).

Select HMG-CoA reductase inhibitors

No interaction studies have been conducted. The HMG-CoA reductase inhibitors atorvastatin, lovastatin and simvastatin are all dependent on CYP3A for metabolism. Lonafarnib is a potent *in vivo* CYP3A mechanism-based inhibitor and, when given concomitantly with either atorvastatin, lovastatin or simvastatin, is expected to increase the plasma concentrations of these statins. This results in an increased risk of myopathy including rhabdomyolysis. Therefore, concomitant use of lonafarnib and atorvastatin, lovastatin and simvastatin are contraindicated (see section 4.3).

Midazolam

When lonafarnib was co-administered with midazolam in healthy adult subjects, multiple dose lonafarnib (100 mg twice daily for 5 consecutive days) increased midazolam (single 3 mg oral dose) C_{max} by 180% and AUC by 639%. This interaction thereby increases the risk of extreme sedation and respiratory depression. Therefore, concomitant use of lonafarnib and midazolam is contraindicated (see sections 4.2, 4.3 and 4.4).

Strong CYP3A inducers

Co-administration of a single oral dose of 50 mg lonafarnib (combined with a single oral dose of 100 mg ritonavir) following 600 mg rifampin once daily for 8 days resulted in the C_{max} of lonafarnib being reduced by 92% and the AUC being reduced by 98%, when compared to rifampin alone in healthy adult subjects. There is no efficacy data available that demonstrates lonafarnib remains effective when administered concomitantly with a strong CYP3A inducer. Therefore, the concomitant use of lonafarnib and a strong CYP3A inducer should be avoided, and therapeutic alternatives sought (see section 4.4).

Moderate CYP3A inducers

No interaction studies have been conducted with a moderate CYP3A inducer. There is no efficacy data available demonstrating that lonafarnib remains effective when given concomitantly with a moderate CYP3A inducer. Therefore, the concomitant use of lonafarnib and a moderate CYP3A inducer should be avoided, and therapeutic alternatives sought (see section 4.4).

Weak CYP3A inducers

No interaction studies have been conducted with a weak CYP3A inducer. There is no efficacy data available demonstrating that lonafarnib remains effective when given concomitantly with a weak CYP3A inducer. Therefore, the concomitant use of lonafarnib and a weak CYP3A inducer should be avoided, and therapeutic alternatives sought (see sections 4.2 and 4.4). If co-administration with a weak CYP3A inducer is unavoidable, maintain the current dose of lonafarnib. If the patient has not already been escalated to the maintenance dose of 150 mg/m² twice daily, the timing of their scheduled dose increase should be maintained.

Foods and select juices that affect the metabolism of lonafarnib

Grapefruit, cranberries, pomegranate and Seville oranges (*e.g.*, orange marmalade), otherwise known as sour or bitter oranges, inhibit the CYP3A system. Ingestion of food or juices containing these fruits should be avoided while taking lonafarnib (see section 4.2).

Weak CYP3A inhibitors

No interaction studies have been conducted with a weak CYP3A inhibitor. No dose adjustment is considered necessary; however, if the concomitant use of a weak CYP3A inhibitor induces a persistent toxicity, the dose of lonafarnib should be reduced by 50% and QTc monitoring is recommended (see sections 4.2 and 6.6).

Loperamide

When lonafarnib was co-administered with loperamide in healthy adult subjects, multiple dose lonafarnib (100 mg twice daily for 5 consecutive days) increased loperamide (single 2 mg oral dose) C_{max} by 214% and AUC by 299%. The dose of loperamide should not exceed 1 mg daily (see section 4.4). In the event more than 1 mg of loperamide daily is to be administered, the dose should be slowly increased with caution as needed to treat diarrhoea.

CYP2C19 substrates

When lonafarnib was co-administered with the CYP2C19 substrate omeprazole in healthy adult subjects, multiple dose lonafarnib (75 mg twice daily for 5 consecutive days) increased omeprazole (single 40 mg oral dose) C_{max} by 28% and AUC by 60%. Patients taking medicinal products that are CYP2C19 substrates should be monitored during this period for potential adverse reactions, with dose adjustments made as necessary.

MATE1 and MATE2-K

Based on *in vitro* data, lonafarnib is a MATE1/MATE2-K inhibitor at clinically relevant maximal systemic concentrations and could potentially precipitate a clinically relevant interaction. Currently, the only identified clinically relevant substrate of MATE1/MATE2-K is metformin. Concomitant use of metformin and lonafarnib should be avoided. If metformin is required, clinicians should carefully monitor the patient for interactions with lonafarnib.

P-glycoprotein substrates

When lonafarnib was co-administered with the P-glycoprotein substrate fexofenadine in healthy adult subjects, multiple dose lonafarnib (100 mg twice daily for 5 consecutive days) increased fexofenadine (single 180 mg oral dose) C_{max} by 21% and AUC by 24%. When lonafarnib is co-administered with P-glycoprotein substrates (*e.g.*, digoxin, dabigatran) where minimal concentration changes may lead to serious or life-threatening toxicities, monitor for adverse reactions and reduce the dose of the P-glycoprotein substrate in accordance with its approved product labelling.

OCT1 substrates

In vitro studies indicate that lonafarnib is an OCT1 inhibitor at clinically relevant systemic concentrations. However, the clinical relevance is currently unknown.

Oral contraceptives

There have been no studies assessing the interaction of concomitant lonafarnib and an oral contraceptive. Females of childbearing potential must use effective contraception during treatment with Zokinvy and for at least 1 week after the final dose (see section 4.6).

Paediatric population

Interaction studies have only been performed in adults.

4.6 Fertility, pregnancy and lactation

Women of childbearing potential/Contraception in males and females

Females of childbearing potential must use effective contraception during treatment with Zokinvy and for at least 1 week after the final dose. Males with female partners of reproductive potential must use effective contraception during treatment with Zokinvy and for at least 3 months after the final dose.

Effects of Zokinvy on contraceptive steroids have not been studied. A barrier method must be added if systemic steroids are used for contraception.

Pregnancy

There are no or limited data from the use of lonafarnib in pregnant women. Studies in animals have shown reproductive toxicity (see section 5.3). Lonafarnib is not recommended during pregnancy and in women of childbearing potential not using contraception.

Breast-feeding

It is unknown whether lonafarnib is excreted in human milk. Animal studies have shown excretion of lonafarnib in milk (for details see section 5.3). A risk to the newborns/infants cannot be excluded.

A decision must be made whether to discontinue breast-feeding or to discontinue therapy with lonafarnib taking into account the benefit of breast-feeding to the child and the benefit of therapy to the woman

Fertility

There are no data on the effects of lonafarnib on fertility in humans. In animal studies, lonafarnib resulted in changes in the male and female reproductive tracts and resorptions (see section 5.3). The potential effect of lonafarnib on fertility in humans is currently unknown.

4.7 Effects on ability to drive and use machines

Lonafarnib has a minor influence on the ability to drive and use machines. Fatigue may occur following the administration of lonafarnib (see section 4.8).

4.8 Undesirable effects

Summary of the safety profile

The most frequently occurring adverse reactions are: vomiting (86%), diarrhoea (78%), increased aspartate aminotransferase (64%), increased alanine aminotransferase (50%), decreased appetite (41%), nausea (38%), abdominal pain (35%), fatigue (29%), decreased weight (27%), constipation

(18%) and upper respiratory tract infection (11%). Most adverse reactions occurred within the first 4 weeks following initiation of treatment and in general steadily decreased with increasing duration of treatment.

The most serious adverse reactions are increased alanine aminotransferase (3.6%), increased aspartate aminotransferase (3.6%), cerebral ischaemia (3.2%), pyrexia (1.6%) and dehydration (1.6%).

Tabulated list of adverse reactions

Adverse reactions occurring in the clinical trials are presented in Table 3 by System Organ Class and Preferred Term. Frequencies are defined as: very common ($\geq 1/10$), common ($\geq 1/100$ to <1/10), uncommon ($\geq 1/1,000$ to <1/100), rare ($\geq 1/10,000$ to <1/1,000), very rare (<1/10,000) or not known (cannot be estimated from the available data). Within each frequency grouping, adverse reactions are presented in order of decreasing frequency within each System Organ Class.

Table 3: Adverse reactions

System organ class	Very common	Common
Infections and	Upper respiratory tract infection	Infection
infestations		Rhinitis
		Gastroenteritis
		Influenza
		Oral pustule
		Perirectal abscess
		Pneumonia
		Sinusitis
Blood and lymphatic	Haemoglobin decreased	White blood cell count decreased
system disorders		
Metabolism and	Decreased appetite	Dehydration
nutrition disorders	Weight decreased	Hypermagnesaemia
		Hypokalaemia
		Hypoalbuminaemia
		Hyponatraemia
Psychiatric disorders		Depressed mood
Nervous system		Cerebral ischaemia
disorders		Headache
disorders		Dizziness
		Paraesthesia
Descriptory theresis		
Respiratory, thoracic		Cough
and mediastinal		Epistaxis
disorders		Laryngeal/oropharyngeal pain
	77	Nasal congestion
Gastrointestinal	Vomiting	Flatulence
disorders	Diarrhoea	Colitis
	Nausea	Dyspepsia
	Abdominal pain ^a	Gastritis
	Constipation	Lower gastrointestinal haemorrhage
Hepatobiliary disorders	Aspartate aminotransferase	Blood creatinine decreased
	increased	
	Alanine aminotransferase	
	increased	
	Blood bicarbonate decreased	
Skin and subcutaneous		Rash
tissue disorders		Pruritus
		Dry skin
		Skin hyperpigmentation
Musculoskeletal and		Musculoskeletal pain
connective tissue		Back pain
disorders		Pain in extremity
General disorders and	Fatigue	Fever
administration site		Chest pain
conditions		Chills
Injury, poisoning and		Tooth fracture
procedural		
complications		
Complications	I	<u>I</u>

^a Abdominal pain includes abdominal pain and abdominal pain upper

Description of selected adverse reactions

Gastrointestinal adverse reactions

Gastrointestinal adverse reactions (vomiting [85.7%], diarrhoea [77.8%], nausea [38.1%]) were the most frequently reported adverse reactions. Of the patients with treatment related vomiting, 29 (53.7%) patients had Grade 1 vomiting (defined as no intervention required) and 25 (46.3%) had

Grade 2 vomiting (defined as outpatient intravenous hydration; medical intervention required). Of these patients with treatment related nausea, 23 (95.8%) had Grade 1 nausea (defined as loss of appetite without alteration in eating habits) and 1 (4.2%) patient had Grade 2 nausea (defined as oral intake decreased without significant weight loss, dehydration or malnutrition). During the first 4 months of treatment in ProLon1, 19 (67.9%) patients had vomiting and 10 (35.7%) patients had nausea. By the end of therapy, 4 (14.3%) patients required anti-emetics or anti-nauseants (see section 4.4). A total of 4 patients discontinued treatment, mostly due to nausea or vomiting.

Most patients with treatment related diarrhoea (approximately 94%) experienced mild or moderate diarrhoea; 38 (77.6%) patients reported Grade 1 (defined as an increase of less than 4 stools per day over baseline) and 8 (16.3%) patients reported Grade 2 treatment related diarrhoea (defined as an increase of 4 to 6 stools per day over baseline; limiting instrumental activities of daily living). Three (6.1%) patients reported Grade 3 diarrhoea (defined as an increase of 7 or more stools per day over baseline; hospitalisation indicated; severe increase in ostomy output compared to baseline; limiting self-care activities of daily living). During the first 4 months of treatment in ProLon1, 23 (82.1%) patients had diarrhoea; by the end of therapy, 3 (10.7%) patients had diarrhoea. Twelve (42.9%) patients were treated with loperamide.

Electrolyte abnormalities

Electrolyte abnormalities (hypermagnesaemia, hypokalaemia, hyponatraemia) were experienced by 4 (6.3%) patients. Of the 2 patients who experienced hypermagnesaemia, 2 (100%) patients had Grade 1 hypermagnesaemia (defined as > upper limit of normal [ULN] to 3.0 mg/dL; >ULN to 1.23 mmol/L). Of the 2 patients who experienced hypokalaemia, 1 (50%) patient had Grade 1 hypokalaemia (defined as < lower limit of normal [LLN] to 3.0 mmol/L) and 1 (50%) patient had Grade 3 hypokalaemia (defined as <3.0 to 2.5 mmol/L; hospitalisation indicated). Of the 1 patient that experienced hyponatraemia, 1 (100%) patient had Grade 1 hyponatraemia (defined as <LLN to 130 mmol/L). Dehydration was experienced by 3 (4.8%) patients. Of the 3 patients who experienced dehydration, 1 (33.3%) patient had Grade 1 dehydration (defined as increased oral fluids indicated; dry mucous membranes; diminished skin turgor) and 2 (66.7%) patients had Grade 2 dehydration (defined as intravenous fluids indicated).

Aminotransferase increases

Increased alanine aminotransferase was recorded for 14 (50.0% of patients) ProLon1 patients. Of the patients with increased alanine aminotransferase, 11 (78.6%) patients experienced a Grade 1 increase (defined as greater than ULN to 3.0 times ULN if baseline was normal; 1.5 to 3.0 times baseline if baseline was abnormal), 1 (7.1%) patient experienced a Grade 2 increase (defined as >3.0 to 5.0 times ULN if baseline was normal; >3.0 to 5.0 x baseline if baseline was abnormal), and 2 (14.3%) patients experienced a Grade 3 increase (defined as >5.0 to 20.0 x ULN if baseline was normal; >5.0 to 20.0 x baseline if baseline was abnormal).

Increased aspartate aminotransferase was recorded for 18 (64.3%) ProLon1 patients. Of these patients, 17 (94.4%) patients experienced a Grade 1 increase (defined as greater than ULN to 3.0 times ULN if baseline was normal; 1.5 to 3.0 times baseline if baseline was abnormal) and 1 (5.6%) patient experienced a Grade 3 increase (defined as >5.0 to 20.0 x ULN if baseline was normal; >5.0 to 20.0 x baseline if baseline was abnormal).

Reporting of suspected adverse reactions

Reporting suspected adverse reactions after authorisation of the medicinal product is important. It allows continued monitoring of the benefit/risk balance of the medicinal product. Healthcare professionals are asked to report any suspected adverse reactions via the national reporting system listed in Appendix V.

4.9 Overdose

In the event of acute overdose, supportive medical care should be given as clinically indicated, including fluid replacement to avoid electrolyte imbalance and close monitoring of vital signs. There is no antidote to lonafarnib to reverse overdose.

5. PHARMACOLOGICAL PROPERTIES

5.1 Pharmacodynamic properties

Pharmacotherapeutic group: Other alimentary tract and metabolism products, Various alimentary tract and metabolism products, ATC code: A16AX20

Mechanism of action

Lonafarnib is a disease modifying agent that prevents farnesylation, thereby reducing the accumulation of aberrant progerin and progerin-like proteins in the cell's inner nuclear membrane. This results in maintaining cell integrity and normal function. The accumulation of progerin and progerin-like proteins in the cells within the walls of large blood vessels causes inflammation and fibrosis.

Clinical efficacy and safety

The clinical efficacy and safety of lonafarnib have been evaluated in two Phase 2 studies (ProLon1 and ProLon2). Both studies were single-centre, open-label, single-arm trials that evaluated the efficacy and safety of lonafarnib in patients with genetically confirmed HGPS or a processing-deficient progeroid laminopathy. Analysis was done by combining the studies into a pooled analysis to evaluate differences in survival between those HGPS patients treated with lonafarnib and those that were lonafarnib-naïve. Survival analyses were conducted at 1, 2 and 3 years based upon the period of lonafarnib monotherapy in either ProLon1 or ProLon2 and using vital status as of August 1, 2021, otherwise called last follow-up.

There were 28 patients in ProLon1 (26 patients with classic HGPS, 1 patient with non-classic HGPS and 1 patient with a progeroid laminopathy with a *LMNA* heterozygous mutation with progerin-like protein accumulation). Patients received lonafarnib over 24 to 30 months. Patients initiated treatment with lonafarnib 115 mg/m² twice daily. After 4 months of treatment, patients who tolerated treatment had an increase in dose to 150 mg/m² twice daily. Among the 28 patients treated, 27 patients with HGPS (16 females, 11 males) were included in the survival assessment. The median age at treatment initiation for the 27 patients was 7.5 years (range: 3 to 16 years). At the start of the study all patients were less than 18 years of age.

There were 35 patients in ProLon2 (34 patients with classic HGPS and 1 patient with non-classic HGPS). Patients received lonafarnib over 12 to 36 months. Patients were treated with lonafarnib 150 mg/m² twice daily. Among the 35 patients treated, all were included in the survival assessment. The median age at treatment initiation was 6.0 years (range: 2 to 17 years). At the start of the study all patients were less than 18 years of age.

Of the 63 patients in ProLon1 and ProLon2, 15 (24%) required some form of dosing adjustment. One (2%) patient discontinued, 11 (17%) patients had their dose interrupted, and 3 (5%) patients reduced dose. For 10 patients (10/63, 16%), the action taken was associated with a gastrointestinal disturbance, a known and common side-effect of lonafarnib.

The retrospective 3-year survival analysis was based on the mortality data from 62 HGPS patients (27 treatment-naïve patients in ProLon1 and 35 treatment-naïve patients in ProLon2) treated with lonafarnib monotherapy and data from matched, untreated patients in a separate natural history cohort.

The mean lifespan of HGPS patients treated with lonafarnib increased by an average of 0.44 to 0.47 years (without and with adjustment for age at start of treatment, respectively) through the first

3 years of follow-up. However due to the uncertainties of the available data this might be as low as 2.4 months.

At last follow-up time (*i.e.*, August 1, 2021) the mean lifespan of HGPS patients treated with lonafarnib increased by an average of 4.3 years. Given the limited information in the datasets this can be as low as 2.6 years. The results for the last follow-up time should be interpreted with some caution as patients underwent additional (potentially beneficial) treatments.

The survival analysis summary is provided in Table 4.

Table 4: Survival analysis summary for patients with Hutchinson-Gilford progeria syndrome (lonafarnib treated versus external natural history cohort)

(Ionarai nib treated versus externar natural nistory conort)			
	Difference in RMST* in years	Hazard ratio* (95%-CI)	
	(95%-CI)		
3-year follow-up	0.466 (0.204, 0.728) P1+P2	0.28 (0.107, 0.756) P1+P2	
	0.414 (0.042, 0.785) P1	0.15 (0.017, 1.263) P1	
	0.172 (-0.101, 0.445) P2	0.71 (0.199, 2.556) P2	
last follow-up (August 1, 2021)	4.338 (2.551, 6.126) P1+P2	0.28 (0.154, 0.521) P1+P2	
2-year follow-up	0.237 (0.074, 0.401) P1+P2	0.29 (0.097, 0.838) P1+P2	
1-year follow-up	0.094 (0.034, 0.154) P1+P2	0.20 (0.054, 0.732) P1+P2	

CI = confidence interval; P1 = ProLon1; P2 = ProLon2; RMST = restricted mean survival time There were 27 patients in ProLon1 and 35 patients in ProLon2.

This medicinal product has been authorised under 'exceptional circumstances'. This means that due to the rarity of the disease it has not been possible to obtain complete information on this medicinal product. The European Medicines Agency will review any new information which may become available every year and this SmPC will be updated as necessary.

5.2 Pharmacokinetic properties

Absorption

Absolute bioavailability has not been assessed. Lonafarnib is absorbed via the oral route. The median time to maximum peak concentration (t_{max}) was 2 to 4 hours. Following multiple dose administration of lonafarnib (100 mg twice daily for 5 days) in healthy volunteers, the mean maximum peak concentration was 964 ng/mL observed at a median time of 4 hours (2 to 5 hours range).

In healthy volunteers, the exposure following a single oral dose of 75 mg lonafarnib taken as an intact capsule was compared to the exposure following a single oral dose of 75 mg lonafarnib capsule contents mixed with orange juice (for instructions on mixing the capsule contents with orange juice see section 6.6). When the capsule contents were mixed with orange juice the C_{max} of lonafarnib was reduced by 9% and the AUC was reduced by 8% as compared to when administered as an intact capsule.

In healthy volunteers, following a single oral dose of 100 mg lonafarnib, food decreased the absorption of lonafarnib and the relative oral bioavailability under fed conditions as compared to fasted conditions was 48% and 77% based on C_{max} and AUC, respectively. Multiple-dose administration of lonafarnib with food in healthy adult subjects did not have a significant effect on bioavailability and resulted in lower inter-subject variability (\sim 16%).

^{*} Estimates are based on matching as follows: for each lonafarnib patient a random match untreated patient was selected with the same sex and same continent. Lonafarnib patients were matched sequentially from the lonafarnib patient with oldest age at start to the youngest. The age at start of treatment of the untreated patient within a matched pair was set to that of the lonafarnib patient. If an untreated patient had a longer follow-up than the lonafarnib treated patient in a matched pair, then this follow-up was censored at the length of the follow-up of the lonafarnib treated patient. Regression analysis for the RMST and Cox proportional hazard regression for the hazard ratio had sex and continent as stratification factors and age at start of treatment as covariate.

In healthy volunteers, the accumulation ratio is estimated to be 4.46 for AUC_{TAU}/AUC_{0-12} and 3.36 for C_{max} .

The intra-individual variability is 20.79% for C_{max} and 21.13% for AUC_{TAU} and the inter-individual variability is 36.92% for C_{max} and 50.75% for AUC_{TAU} .

Distribution

In vitro plasma protein binding of lonafarnib was ≥99% over the concentration range between 0.5 to 40.0 micrograms/mL. The blood-to-plasma ratio was 0.992 to 1.56.

Lonafarnib exhibits time-dependent pharmacokinetics. Comparing studies in healthy adult volunteers of single-dose 75 mg lonafarnib to 75 mg lonafarnib twice daily for 5 days shows the lonafarnib apparent volume of distribution is reduced by 60% (242 L and 97.4 L, respectively) following multiple dose lonafarnib for 5 days.

Biotransformation

Lonafarnib is extensively metabolised via hepatic means. Lonafarnib accounted for 50% to 57% of the profiled plasma radioactivity. Total plasma recovery for the two metabolites of interest: HM17 (15.1%) and HM21 (13.9%); therefore, a total of 79% to 86% of the plasma radioactivity was recovered. The common metabolic pathways included oxidation, dehydrogenation and combinations of these two processes. Most of the metabolites resulted from structural changes in the pendant piperidine ring region of lonafarnib.

HM21 is a pharmacologically active metabolite. Following oral administration of 100 mg lonafarnib twice daily for 5 days, HM21 has a peak plasma concentration of 94.8 ng/mL occurring after approximately 4 hours (range: 3 to 6), with an AUC_{TAU} of 864 ng·h/mL. Following oral administration of 75 mg lonafarnib twice daily for 5 days, HM21 has a peak plasma concentration of 82.1 ng/mL after approximately 3 hours (range: 3 to 5), with an AUC_{TAU} of 767 ng·h/mL.

In vitro metabolism studies indicate that CYP3A4 and CYP3A5 are mainly responsible for the oxidative metabolism of lonafarnib and that lonafarnib is an *in vivo*-sensitive CYP3A4 substrate.

Twenty-one metabolites were characterised/identified in urine and faeces. No single uncharacterised metabolite represented greater than 5% of the dose.

Transporters

Based on the *in vitro* data, lonafarnib is most likely a substrate of P-glycoprotein and not a substrate of BCRP, OCT1, OATP1B1 and OATP1B3.

Elimination

A 14 C- absorption, metabolism and excretion trial conducted in healthy volunteers following single-dose administration of lonafarnib revealed that drug-derived radioactivity was primarily excreted via the faeces. Mean cumulative excretion of radioactivity was 61% in faeces and less than 1% in urine up to 24 hours post-dose (total recovery was \sim 62% in the mass balance study).

Lonafarnib exhibits time-dependent pharmacokinetics. Comparing studies in healthy adult volunteers of single-dose 75 mg lonafarnib to 75 mg lonafarnib twice daily for 5 days shows lonafarnib clearance was reduced by 75% (48.2 L/h and 12.1 L/h, respectively) and the t_{1/2} increased by 60% (3.5 h versus 5.6 h, respectively) following multiple dose lonafarnib for 5 days.

Special populations

Hepatic impairment

Lonafarnib has not been studied in patients with hepatic impairment. Co-administration of a single oral dose of 50 mg lonafarnib (combined with a single oral dose of 100 mg ritonavir) in mild and moderate hepatically impaired subjects showed similar lonafarnib exposures relative to the matched normal control group (normal hepatic function). These results indicate no dose adjustments are warranted in patients with mild or moderate hepatic impairment (see section 4.2). Lonafarnib is contraindicated in patients with severe hepatic impairment (see section 4.3) due to the predicted safety issue of decompensation due to the risk of diarrhoea (see sections 4.4 and 4.8). Lonafarnib (and most likely HM21) is extensively metabolised in the liver. Therefore, decreased hepatic function will most likely lead to an increase in exposure to lonafarnib (effect on HM21 is unknown) (see section 4.4).

Renal impairment

Lonafarnib has not been studied in patients with renal impairment (see section 4.4). Lonafarnib and HM21 are only excreted to a limited extent via urine. Therefore, it is not expected that renal impairment will affect the exposure to lonafarnib and HM21.

Gender

In healthy volunteers, following a single oral dose of 100 mg lonafarnib, the pharmacokinetic data suggest lonafarnib exposures (AUC_{0-inf}) are higher in female subjects (44% higher) as compared to male subjects. Gender had less of an effect (26%) on the C_{max} as compared with AUC_{0-inf} .

Age

In healthy volunteers, following a single oral dose of 100 mg lonafarnib, the pharmacokinetic data show lonafarnib exposures (AUC_{0-inf}) are higher in elderly subjects (59% higher in those aged 65 years or older) as compared to younger subjects aged 18 to 45 years. Age had less of an effect (27%) on the C_{max} as compared with AUC_{0-inf} .

5.3 Preclinical safety data

Lonafarnib had no effects on QT or QTc interval in guinea pigs and no electrocardiogram (ECG) changes were observed in monkeys. Lonafarnib produced modest and isolated effects on the QT interval of ECG in rats at estimated exposures similar to that seen in humans.

A no-observed-adverse-effect level (NOAEL) could not be established in studies of up to 1-year duration in monkeys. Systemic toxicity was observed in 3-month and 1-year toxicity studies in rats and monkeys following repeated oral administration of lonafarnib at doses ≥30 and ≥10 mg/kg/day, respectively, corresponding to exposures lower than what is seen in patients. Toxicity findings included bone marrow suppression, testicular toxicity and lymphoid toxicity in rats and monkeys; kidney changes in rats (vacuolisation, mineralisation and necrosis of the inner renal medulla); and diarrhoea and electroretinographic changes in monkeys. In a 3-month toxicity study in monkeys, acute morbidity due to haemorrhage in multiple organs was observed in a small number of monkeys administered 60 mg/kg/day, corresponding to exposures similar to that seen in humans (at 150 mg/m² twice daily). In toxicity studies in monkeys, ocular findings of single cell necrosis of retinal photoreceptors were observed at ≥40 mg/kg/day. In a 3-month follow up study, changes in electroretinography were noted at ≥15 mg/kg/day, including substantial changes in scotopic amplitudes at 60 mg/kg/day indicating perturbation of rod cells and impairment of night vision. The NOAEL for ocular toxicity for lonafarnib was considered to be 20 mg/kg/day, corresponding to exposures similar to those seen in humans (at 150 mg/m² twice daily).

Lonafarnib increased pre- and post-implantation loss and decreased the number of live foetuses in female rats at doses ≥ 30 mg/kg/day. Decreased maternal body weight and lower foetal body weights were also observed at this dose level. The NOAEL for maternal toxicity and F1 litters was considered 10 mg/kg/day, with an estimated exposure level lower than what is seen in humans at 150 mg/m² twice daily.

Reproductive organ toxicity was observed in male rats and monkeys, including lower testicular and epididymal weight, aspermia, altered spermatogenesis and spermatogonial debris in male rats at

≥90 mg/kg/day, and lower testes weights in male monkeys at the lowest tested dose 10 mg/kg/day. The NOAEL or the lowest tested dose regarding these effects corresponds to exposure levels lower than what is seen in humans at 150 mg/m² twice daily.

Lonafarnib demonstrated teratogenic potential at clinically relevant exposures in rabbits in the absence of maternal toxicity, with increased incidence of malformations and variations in foetal skeletal development observed at the lowest tested dose 10 mg/kg/day, corresponding to an exposure level lower than what is seen in humans at 150 mg/m² twice daily. Maternal toxicity was observed at ≥40 mg/kg/day and both maternal and embryofoetal toxicity, including abortion, discoloured urine, body weight loss, increased post-implantation loss and decreased foetal body weight, were observed at 120 mg/kg/day, corresponding to exposures greater than those seen in humans (~2- and 25-times the human exposure at 150 mg/m² twice daily, respectively). In rats, lonafamib had no adverse effects on F1 and F2 generations in a pre- and post-natal development study. Lonafamib is excreted in milk following oral administration in lactating rats, with a mean milk to plasma concentration ratio of 1.5 at 12 hours.

Overall, lonafarnib does not represent a genotoxic concern based on results of *in vitro* tests, including bacterial reverse mutation assays and a chromosome aberration assay using human peripheral blood lymphocytes. In the *in vivo* mouse bone micronucleus assay, lonafarnib was not genotoxic at doses up to 50 and 60 mg/kg/day (intraperitoneal injection) in male and female mice, respectively. However, these dose levels are lower than the clinical relevant dose.

The carcinogenic potential of lonafarnib has not been studied.

6. PHARMACEUTICAL PARTICULARS

6.1 List of excipients

Capsule contents

Povidone Poloxamer Croscarmellose sodium Silica, colloidal anhydrous Magnesium stearate

Capsule shell

Gelatin (E 171)
Titanium dioxide
Yellow iron oxide (E 172)
Red iron oxide (E 172) (75 mg capsules only)
Sunflower lecithin (E 322)

Black ink

Shellac Iron oxide black (E 172) Propylene glycol Ammonia solution Potassium hydroxide

6.2 Incompatibilities

Not applicable.

6.3 Shelf life

3 years

6.4 Special precautions for storage

Store in the original package. Keep the bottle tightly closed in order to protect from moisture. This medicinal product does not require any special temperature storage conditions.

6.5 Nature and contents of container

HDPE bottle, containing desiccant in a cannister and capsules, with induction seal and polypropylene cap. Pack size of 30 hard capsules.

6.6 Special precautions for disposal and other handling

Any unused medicinal product or waste material should be disposed of in accordance with local requirements.

Patients unable to swallow capsules whole

If capsules cannot be swallowed whole, capsules can be opened and the contents of the capsule can be mixed with orange juice.

- Step 1: Using a clean medicine cup, measure either 5 mL or 10 mL of orange juice. You can choose to use 5 mL or 10 mL of orange juice.
- Step 2: Pour the orange juice measured in **Step 1** into a clean cup.
- Step 3: Hold a capsule above the cup containing the orange juice. Hold the capsule between your thumb and forefinger on both sides. Gently twist and pull apart the capsule.
- Step 4: Empty the contents of the capsule directly into the cup containing the orange juice.
- Step 5: Using a clean spoon, mix the capsule contents and orange juice well. If only 1 capsule is to be taken, skip to **Step 7**. If 2 capsules are to be taken proceed to **Step 6**.
- Step 6: If 2 capsules will be taken, repeat Steps 1 through 5 for the second capsule. After finishing, go to **Steps 7, 8 and 9**.
- Step 7: Take all of the mixture with food within about 10 minutes of preparing. Each dose must be mixed and consumed within 10 minutes. The mixture should only be prepared at the time it is to be consumed.
- Step 8: Rinse the medicine cup used to measure the orange juice and fill it with 5 mL of water for each capsule mixed with orange juice.
- Step 9: Pour the water measured in **Step 8** into the cup used to mix the Zokinvy and orange juice. Gently swirl the water around the cup. Consume the water.

Patients with a body surface area between 0.30 m² and 0.38 m², patients taking Zokinvy and a weak CYP3A inhibitor and experiencing persistent toxicities or patients with a dysfunctional CYP3A4 polymorphism requiring a reduced daily dose of less than or equal to 50 mg

Patients with a body surface area between 0.30 m² and 0.38 m² require a daily dose of 75 mg (37.5 mg twice daily). Some patients taking Zokinvy concomitantly with a weak CYP3A inhibitor and experiencing persistent side-effects or patients with a dysfunctional CYP3A4 polymorphism may

require a daily dose of 50 mg (25 mg twice daily). In these instances, a Zokinvy 75 mg or 50 mg capsule will need to be mixed with 10 mL of orange juice to reach the appropriate dose. Only half of the 10-mL mixture will be consumed yielding a dose of either 25 mg or 37.5 mg.

- Step 1: Use a clean medicine cup and fill it with 10 mL of orange juice.
- Step 2: Pour the orange juice measured in **Step 1** into a clean cup for mixing.
- Step 3: Depending on your doctor's direction, hold either a Zokinvy 75 mg or 50 mg capsule above the cup containing the orange juice. Hold the capsule between your thumb and forefinger on both sides. Gently twist and pull apart the capsule.
- Step 4: Empty the contents of the capsule directly into the cup containing the orange juice.
- Step 5: Using a clean spoon, mix the capsule contents and orange juice well.
- Step 6: Pour 5 mL of the orange juice and mixture from the mixing cup into a clean medicine cup.
- Step 7: Take the 5-mLmixture with food and within about 10 minutes of preparing. Each dose must be mixed and consumed within 10 minutes. The mixture should only be prepared at the time it is to be consumed.
- Step 8: Fill the medicine cup used to consume the mixture with 5 mL of water.
- Step 9: Gently swirl the water around the medicine cup. Consume the water.

7. MARKETING AUTHORISATION HOLDER

EigerBio Europe Ltd. 1 Castlewood Avenue Rathmines, D06 H685, Ireland

8. MARKETING AUTHORISATION NUMBER(S)

Zokinvy 50 mg hard capsules

EU/1/22/1660/001

Zokinvy 75 mg hard capsules

EU/1/22/1660/002

9. DATE OF FIRST AUTHORISATION/RENEWAL OF THE AUTHORISATION

Date of first authorisation:

10. DATE OF REVISION OF THE TEXT

Detailed information on this medicinal product is available on the website of the European Medicines Agency http://www.ema.europa.eu.

ANNEX II

- A. MANUFACTURER RESPONSIBLE FOR BATCH RELEASE
- B. CONDITIONS OR RESTRICTIONS REGARDING SUPPLY AND USE
- C. OTHER CONDITIONS AND REQUIREMENTS OF THE MARKETING AUTHORISATION
- D. CONDITIONS OR RESTRICTIONS WITH REGARD TO THE SAFE AND EFFECTIVE USE OF THE MEDICINAL PRODUCT
- E. SPECIFIC OBLIGATION TO COMPLETE POST-AUTHORISATION MEASURES FOR THE MARKETING AUTHORISATION UNDER EXCEPTIONAL CIRCUMSTANCES

A. MANUFACTURER RESPONSIBLE FOR BATCH RELEASE

Name and address of the company responsible for product release

ABF Pharmaceutical Services GmbH Brunner Straße 63/18-19 A-1230 Vienna Austria

B. CONDITIONS OR RESTRICTIONS REGARDING SUPPLY AND USE

Medicinal product subject to restricted medical prescription (see Annex I: Summary of Product Characteristics, section 4.2).

C. OTHER CONDITIONS AND REQUIREMENTS OF THE MARKETING AUTHORISATION

• Periodic safety update reports (PSURs)

The requirements for submission of PSURs for this medicinal product are set out in the list of Union reference dates (EURD list) provided for under Article 107c(7) of Directive 2001/83/EC and any subsequent updates published on the European medicines web-portal.

The marketing authorisation holder (MAH) shall submit the first PSUR for this product within 6 months following authorisation.

D. CONDITIONS OR RESTRICTIONS WITH REGARD TO THE SAFE AND EFFECTIVE USE OF THE MEDICINAL PRODUCT

• Risk management plan (RMP)

The MAH shall perform the required pharmacovigilance activities and interventions detailed in the agreed RMP presented in Module 1.8.2 of the marketing authorisation and any agreed subsequent updates of the RMP.

An updated RMP should be submitted:

- At the request of the European Medicines Agency;
- Whenever the risk management system is modified, especially as the result of new
 information being received that may lead to a significant change to the benefit/risk profile
 or as the result of an important (pharmacovigilance or risk minimisation) milestone being
 reached.

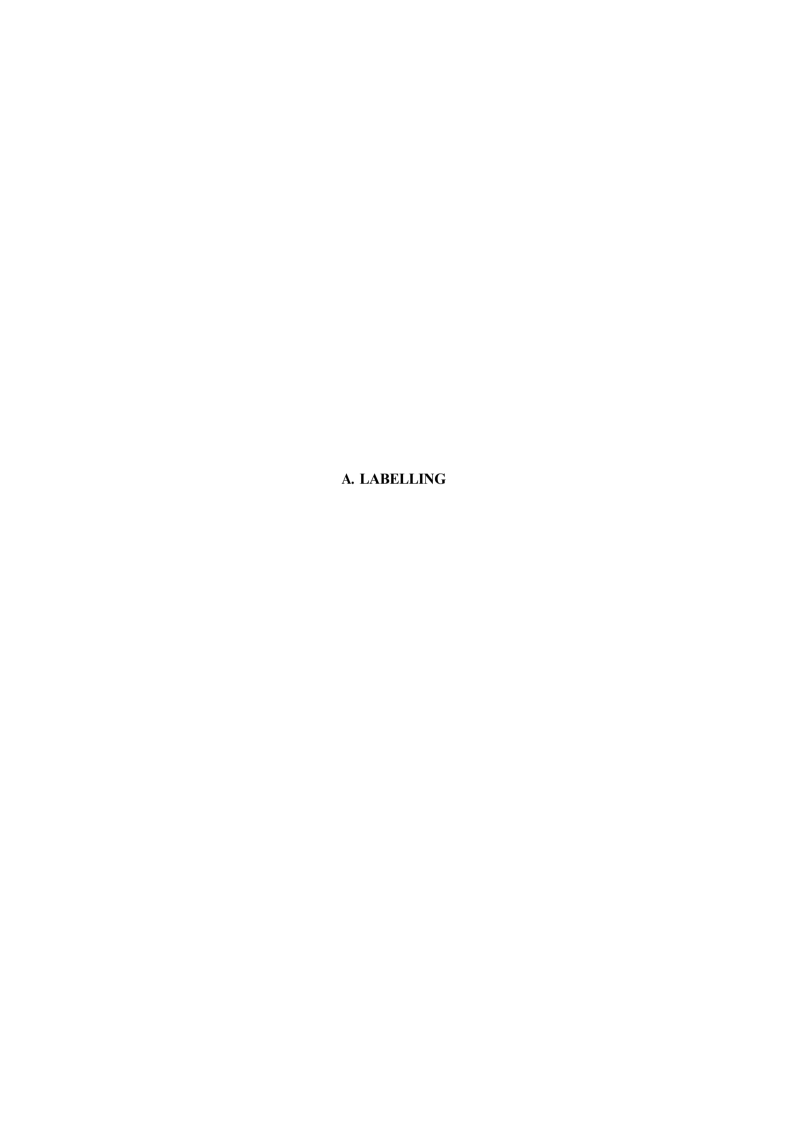
E. SPECIFIC OBLIGATION TO COMPLETE POST-AUTHORISATION MEASURES FOR THE MARKETING AUTHORISATION UNDER EXCEPTIONAL CIRCUMSTANCES

This being an approval under exceptional circumstances and pursuant to Article 14(8) of Regulation (EC) No 726/2004, the MAH shall conduct, within the stated timeframe, the following measures:

Description	Due date
Non-interventional Post authorisation safety	Annual study reports will be submitted with the
study (PASS): in order to further characterise the	annual re-assessment

I	safety, effectiveness and health-related quality of
	life of Zokinvy in patients with Hutchinson-
	Gilford Progeria Syndrome and Processing
	Deficient Progeroid Laminopathies, the MAH
	shall submit the results of a prospective
	observational cohort study based on a registry.

ANNEX III LABELLING AND PACKAGE LEAFLET



CARTON (ZOKINVY 50 mg)
1. NAME OF THE MEDICINAL PRODUCT
Zokinvy 50 mg hard capsules lonafarnib
2. STATEMENT OF ACTIVE SUBSTANCE(S)
Each capsule contains 50 mg lonafarnib.
3. LIST OF EXCIPIENTS
4 DHADMACEUTICAL EODM AND CONTENTS
4. PHARMACEUTICAL FORM AND CONTENTS
30 hard capsules
5. METHOD AND ROUTE(S) OF ADMINISTRATION
Read the package leaflet before use. Oral use
6. SPECIAL WARNINGS THAT THE MEDICINAL PRODUCT MUST BE STORED OUT OF THE SITE AND REACH OF CHILDREN
Keep out of the sight and reach of children.
7. OTHER SPECIAL WARNING(S), IF NECESSARY
Do not swallow the desiccant.
8. EXPIRY DATE
EXP
9. SPECIAL STORAGE CONDITIONS
Store in the original package. Keep the bottle tightly closed in order to protect from moisture. This medicinal product does not require any special temperature storage conditions.

PARTICULARS TO APPEAR ON THE OUTER PACKAGING

10. SPECIAL PRECAUTIONS FOR DISPOSAL OF UNUSED MEDICINAL PRODUCTS OR WASTE MATERIALS DERIVED FROM SUCH MEDICINAL PRODUCTS, IF APPROPRIATE
11. NAME AND ADDRESS OF THE MARKETING AUTHORISATION HOLDER
EigerBio Europe Ltd. I Castlewood Avenue Rathmines, D06 H685, Ireland
12. MARKETING AUTHORISATION NUMBER(S)
EU/1/22/1660/001
13. BATCH NUMBER
Lot
14. GENERAL CLASSIFICATION FOR SUPPLY
15. INSTRUCTIONS ON USE
16. INFORMATION IN BRAILLE
Zokinvy 50 mg
17. UNIQUE IDENTIFIER – 2D BARCODE
2D barcode carrying the unique identifier included

18. UNIQUE IDENTIFIER – HUMAN READABLE DATA

PC SN NN

PARTICULARS TO APPEAR ON THE IMMEDIATE PACKAGING
BOTTLE LABEL (ZOKINVY 50 mg)
1. NAME OF THE MEDICINAL PRODUCT
Zokinvy 50 mg hard capsules lonafarnib
2. STATEMENT OF ACTIVE SUBSTANCE(S)
Each capsule contains 50 mg lonafarnib.
3. LIST OF EXCIPIENTS
4. PHARMACEUTICAL FORM AND CONTENTS
30 hard capsules
5. METHOD AND ROUTE(S) OF ADMINISTRATION
Read the package leaflet before use. Oral use
6. SPECIAL WARNINGS THAT THE MEDICINAL PRODUCT MUST BE STORED OUT OF THE SITE AND REACH OF CHILDREN
Keep out of the sight and reach of children.
7. OTHER SPECIAL WARNING(S), IF NECESSARY
Do not swallow the desiccant.
8. EXPIRY DATE
EXP
9. SPECIAL STORAGE CONDITIONS
Store in the original package. Keep the bottle tightly closed in order to protect from moisture. This medicinal product does not require any special temperature storage conditions.

PRODU	CTS, IF APPROPRIATE
11. NAME A	ND ADDRESS OF THE MARKETING AUTHORISATION HOLDER
12. MARKET	TING AUTHORISATION NUMBER(S)
EU/1/22/1660/00)1
13. BATCH N	NUMBER
Lot	
14. GENERA	L CLASSIFICATION FOR SUPPLY
15. INSTRUC	CTIONS ON USE
16. INFORM	ATION IN BRAILLE
17. UNIQUE	IDENTIFIER – 2D BARCODE
18. UNIQUE	IDENTIFIER – HUMAN READABLE DATA

10. SPECIAL PRECAUTIONS FOR DISPOSAL OF UNUSED MEDICINAL

PRODUCTS OR WASTE MATERIALS DERIVED FROM SUCH MEDICINAL

PARTICULARS TO APPEAR ON THE OUTER PACKAGING
CARTON (ZOKINVY 75 mg)
1. NAME OF THE MEDICINAL PRODUCT
Zokinvy 75 mg hard capsules lonafarnib
2. STATEMENT OF ACTIVE SUBSTANCE(S)
Each capsule contains 75 mg lonafarnib.
3. LIST OF EXCIPIENTS
4. PHARMACEUTICAL FORM AND CONTENTS
30 hard capsules
5. METHOD AND ROUTE(S) OF ADMINISTRATION
Read the package leaflet before use. Oral use
6. SPECIAL WARNINGS THAT THE MEDICINAL PRODUCT MUST BE STORED OUT OF THE SITE AND REACH OF CHILDREN
Keep out of the sight and reach of children.
7. OTHER SPECIAL WARNING(S), IF NECESSARY
Do not swallow the desiccant.
8. EXPIRY DATE
EXP
9. SPECIAL STORAGE CONDITIONS
Store in the original package. Keep the bottle tightly closed in order to protect from moisture. This medicinal product does not require any special temperature storage conditions.

	PRODUCTS, IF APPROPRIATE
11.	NAME AND ADDRESS OF THE MARKETING AUTHORISATION HOLDER
1 Cas	rBio Europe Ltd. stlewood Avenue mines, D06 H685, Ireland
12.	MARKETING AUTHORISATION NUMBER(S)
EU/1	/22/1660/002
13.	BATCH NUMBER
Lot	
14.	GENERAL CLASSIFICATION FOR SUPPLY
15.	INSTRUCTIONS ON USE
16.	INFORMATION IN BRAILLE
Zokii	nvy 75 mg
17.	UNIQUE IDENTIFIER – 2D BARCODE
2D ba	arcode carrying the unique identifier included
18.	UNIQUE IDENTIFIER – HUMAN READABLE DATA
PC SN NN	

10. SPECIAL PRECAUTIONS FOR DISPOSAL OF UNUSED MEDICINAL

PRODUCTS OR WASTE MATERIALS DERIVED FROM SUCH MEDICINAL

PARTICULARS TO APPEAR ON THE IMMEDIATE PACKAGING			
BOTTLE LABEL (75 mg)			
1.	NAME OF THE MEDICINAL PRODUCT		
	anvy 75 mg hard capsules farnib		
2.	STATEMENT OF ACTIVE SUBSTANCE(S)		
Each	n capsule contains 75 mg lonafarnib.		
3.	LIST OF EXCIPIENTS		
4.	PHARMACEUTICAL FORM AND CONTENTS		
30 h	ard capsules		
5.	METHOD AND ROUTE(S) OF ADMINISTRATION		
Read Oral	I the package leaflet before use. use		
6.	SPECIAL WARNINGS THAT THE MEDICINAL PRODUCT MUST BE STORED OUT OF THE SITE AND REACH OF CHILDREN		
Keep	o out of the sight and reach of children.		
7.	OTHER SPECIAL WARNING(S), IF NECESSARY		
Do n	not swallow the desiccant.		
8.	EXPIRY DATE		
EXP			
9.	SPECIAL STORAGE CONDITIONS		
	e in the original package. Keep the bottle tightly closed in order to protect from moisture. This icinal product does not require any special temperature storage conditions.		

PRODUCTS OR WASTE MATERIALS DERIVED FROM SUCH MEDICINAL PRODUCTS, IF APPROPRIATE		
11.	NAME AND ADDRESS OF THE MARKETING AUTHORISATION HOLDER	
12.	MARKETING AUTHORISATION NUMBER(S)	
EU/1	/22/1660/002	
13.	BATCH NUMBER	
Lot		
14.	GENERAL CLASSIFICATION FOR SUPPLY	
15.	INSTRUCTIONS ON USE	
16.	INFORMATION IN BRAILLE	
17.	UNIQUE IDENTIFIER – 2D BARCODE	
10		
18.	UNIQUE IDENTIFIER – HUMAN READABLE DATA	

10. SPECIAL PRECAUTIONS FOR DISPOSAL OF UNUSED MEDICINAL

B. PACKAGE LEAFLET

Package leaflet: Information for the patient

Zokinvy 50 mg hard capsules Zokinvy 75 mg hard capsules lonafarnib

This medicine is subject to additional monitoring. This will allow quick identification of new safety information. You can help by reporting any side-effects you may get. See the end of section 4 for how to report side-effects.

Read this entire 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, pharmacist or nurse.
- 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, pharmacist or nurse. This includes any possible side effects not listed in this leaflet. See section 4.

What is in this leaflet

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

1. What Zokinvy is and what it is used for

What Zokinvy is

Zokinvy contains the active substance lonafarnib.

What Zokinvy is used for

This medicine is used to treat patients aged 12 months and older with the following rare disorders:

- Hutchinson-Gilford progeria syndrome
- processing-deficient progeroid laminopathies

These illnesses are caused by changes in genes needed to make certain proteins. Normal versions of these proteins help to keep cells strong and stable. However, the altered genes cause a build-up of harmful forms of the proteins called progerin or progerin-like proteins. These harmful proteins lead to cell damage that resembles the effects of aging.

How Zokinvy works

Zokinvy works by helping to reduce the build-up of the harmful progerin or progerin-like proteins.

2. What you need to know before you take Zokinvy

Do not take Zokinvy

• if you are allergic to lonafarnib or any of the other ingredients in this medicine (listed in section 6).

- with medicines known as strong CYP3A inhibitors (these can reduce the breakdown of Zokinvy in the body, leading to more side effects, see Other medicines and Zokinvy, below).
- with the medicine midazolam
- with the medicines atorvastatin, lovastatin, simvastatin
- if you have severe hepatic (liver) impairment

Warnings and precautions

Talk to your doctor, pharmacist or nurse before taking Zokinvy.

Please tell your doctor immediately if you:

- are over 10 years of age. Treatment results can vary depending on the age when you start taking Zokinvy.
- have persistent vomiting or diarrhoea and prolonged loss of appetite or weight (see section 4).
- start to take the anti-diarrhoeal medicine loperamide. Because of the interaction between Zokinvy and loperamide, it will be important for your doctor to provide dosing guidance and monitor your use of this medicine.
- are having surgery. You must not use midazolam, a medicine commonly used during surgery, with Zokinvy. Your doctor can provide instructions for this situation.
- experience increased levels of liver enzymes shown by blood tests. Your doctor should monitor your liver function while taking this medicine.
- develop any symptoms of kidney problems. Your doctor should monitor your kidney function while taking this medicine.
- experience any new visual changes. Your doctor should monitor your vision and eye health while taking this medicine.
- are taking a medicine that is a moderate or strong CYP3A inducer. These types of medicines should be avoided (see Other medicines and Zokinvy, below).
- are taking a medicine that is a moderate CYP3A inhibitor. These types of medicines should be avoided (see Other medicines and Zokinvy, below).
- have a known dysfunctional polymorphism in CYP3A4.
- have a progeroid syndrome caused by a mutation in a gene other than *LMNA* or *ZMPSTE24* and do not cause a build-up of the harmful proteins called progerin or progerin-like proteins. Zokinvy is not expected to be effective for these types of progeroid syndromes. Examples of progeroid syndromes that Zokinvy is not expected to provide benefit include, Werner syndrome, Bloom syndrome, Rothmund–Thomson syndrome, Cockayne syndrome, xeroderma pigmentosum, trichothiodystrophy and ataxia-telangiectasia.

Children

Do not give this medicine to children under 12 months of age because it has not been studied in this age group.

Other medicines and Zokinvv

Tell your doctor or pharmacist if you are taking or might take any other medicines.

Some medicines may interact with Zokinvy when taken together. The following medicines must not be combined with Zokinvy:

- medicines that are strong CYP3A inhibitors (they can reduce the breakdown of Zokinvy in the body, leading to more side effects; ask your pharmacist or doctor if any of your other medicines are of this kind)
- medicines that are weak or moderate CYP3A inhibitors (they can reduce the breakdown of Zokinvy in the body, leading to more side effects; ask your pharmacist or doctor if any of your other medicines are of this kind). Your doctor may temporarily reduce your dose of Zokinvy while taking a weak or moderate CYP3A inhibitor

- midazolam (used to treat seizures and for surgical procedures tell your doctor you are taking Zokinvy if you are planning to have surgery)
- atorvastatin, lovastatin or simvastatin (used to lower blood cholesterol)
- medicines that are strong, moderate or weak CYP3A inducers (these can increase the breakdown of Zokinvy in the body, making the medicine less effective; ask your pharmacist or doctor if any of your other medicines are of this kind)
- loperamide (used to treat diarrhoea). The dose of loperamide should not exceed 1 mg daily. Children less than 2 years of age should not take loperamide.
- metformin (used to treat type 2 diabetes)
- medicines that are CYP2C19 substrates (ask your pharmacist or doctor if any of your other medicines are of this kind). If you must take a CYP2C19 substrate, your doctor may need to adjust your dose of the CYP2C19 substrate and monitor your side effects more closely.
- St. John's wort or St. John's wort-containing products (a herbal medicine used to treat mild depression)
- medicines that are P-glycoprotein substrates (ask your pharmacist or doctor if any of your other medicines are this kind). If you must take a P-glycoprotein substrate, your doctor may need to adjust your dose of the P-glycoprotein substrate and monitor your side effects more closely.
- medicines that are OCT1 substrates (ask your pharmacist or doctor if any of your other medicines are this kind)
- oral contraceptives

Zokinvy with food and drink

Do not take Zokinvy with foods or drinks that contain grapefruit, cranberries, pomegranates or Seville (bitter) orange (such as orange marmalade). Foods or drinks containing these fruits may increase the side effects of Zokinvy.

Pregnancy, breast-feeding and fertility

If you are pregnant or breast-feeding, think you may be pregnant or are planning to have a baby, ask your doctor or pharmacist for advice before taking this medicine.

Zokinvy has not been tested in pregnant women.

Zokinvy is not recommended during pregnancy.

Females of childbearing potential must use effective contraception while taking Zokinvy and for at least 1 week after the final dose. Males with female partners of childbearing potential must use effective contraception while taking Zokinvy and for at least 3 months after the final dose. Add a barrier method of contraception if systemic steroids are used for contraception.

It is not known if Zokinvy passes into breast milk and could affect a breast-fed baby. If you wish to breast feed, discuss the benefits and possible risks of doing so or of stopping Zokinvy with your doctor first.

It is not yet known if this medicine affects fertility in men or women.

Driving and using tools or machines

Zokinvy has a minor influence on the ability to drive and use machines. Fatigue may occur following administration of Zokinvy.

Zokinvy contains sodium

This medicine contains less than 1 mmol sodium (23 mg) per dose, that is to say essentially 'sodium-free'.

3. How to take Zokinvy

Always take this medicine exactly as your doctor or pharmacist has told you. Check with your doctor or pharmacist if you are not sure.

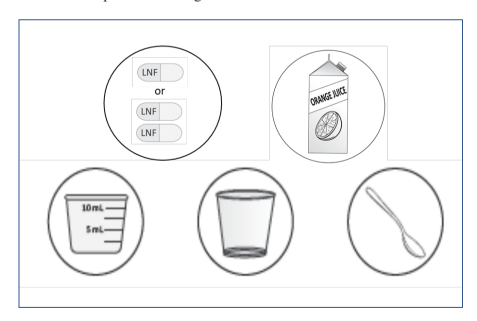
- Zokinvy is taken as 1 or 2 capsules twice a day, about 12 hours apart (morning and evening) with food. The dose of Zokinvy is based on your height and weight.
- Your doctor will work out the right starting dose of Zokinvy for you. This may mean taking capsules
 of different strengths to make up the right amount. After 4 months of treatment with Zokinvy, your
 doctor may increase your dose.
- Make sure you know how many capsules you need to take at every dose and the strength of each capsule you need. Ask your doctor, pharmacist or nurse to write it down (including the colour of the capsule(s) to be taken for each dose).
- Take the capsules with food, washing them down with enough water to help you swallow them. Taking Zokinvy with food may help to reduce side effects.

If you cannot swallow a Zokinvy capsule whole

• If you cannot swallow a Zokinvy capsule whole, use the following directions for mixing the capsule content with orange juice.

What you need to mix Zokinvy with orange juice

- Mix a fresh dose of Zokinvy for each use.
- Set aside the right number of Zokinvy capsules for your dose. Place the capsule or capsules on a clean flat surface.
- Use orange juice only. Do not use other drinks to mix Zokinvy.
- A clean medicine cup with 5 mL and 10 mL measurement levels.
- A clean cup for each Zokinvy capsule to be mixed.
- A clean spoon for stirring the mixture.



How to mix Zokinvy with orange juice

Using a clean medicine cup, measure either 5 mL or 10 mL of orange juice. You can choose to use 5 mL or 10 mL of orange juice.

Step 2: Pour the orange juice measured in Step 1 into a clean cup.	\$ m		
 Hold a Zokinvy capsule above the cup containing the orange juice. Hold the capsule between your thumb and forefinger on both sides. Gently twist and pull apart the capsule. 			
Step 4: Empty all the contents of the capsule into the cup containing the orange juice.			
 Using a clean spoon, mix the capsule contents and orange juice well. If only 1 capsule is to be taken, skip to Step 7. If 2 capsules are to be taken, proceed to Step 6. 			
 Step 6: If 2 capsules will be taken, repeat Steps 1 through 5 for the second capsule. Once the second capsule is mixed - the 2 servings can either be combined in a single cup or remain in 2 serving cups. After you finish, go to Steps 7, 8 and 9. 			
 Step 7: Take all of the Zokinvy mixture: with food. within about 10 minutes of preparing. Each dose must be mixed and consumed within 10 minutes. The mixture should only be prepared at the time it is to be consumed.	TAKE WITHIN 10 Minutes		

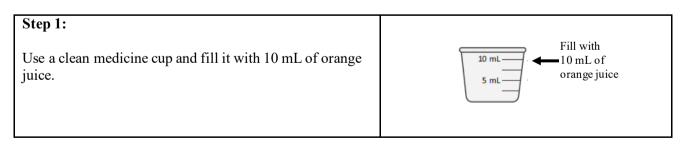
Step 9: Pour the water measured in Step 8 into the cup used to mix the Zokinvy and orange juice (a). Gently swirl the water around the cup (b). Consume the water.

Patients starting Zokinvy with a body surface area between 0.30 m² and 0.38 m², patients taking a weak CYP3A inhibitor concomitantly with Zokinvy and experiencing persistent side-effects and patients with a dysfunctional CYP3A4 polymorphism requiring a reduced daily dose less than or equal to 50 mg

Before beginning to mix Zokinvy with orange juice, read What you need to mix Zokinvy with orange juice.

For patients with a body surface area between 0.30 m² and 0.38 m², the daily starting dose of Zokinvy is 75 mg (37.5 mg twice daily with food). Each dose is prepared fresh using a 75 mg capsule and mixed with 10 mL of orange juice. Half of the 10-mL mixture will be consumed.

For patients taking Zokinvy and a weak CYP3A inhibitor and experiencing persistent side-effects or with a dysfunctional CYP3A4 polymorphism, your doctor may reduce your daily dose of Zokinvy by 50%. Prepare each dose fresh. If your reduced daily dose is 50 mg (25 mg twice daily), mix the contents of a 50 mg capsule in 10 mL of orange juice. Half of the 10-mL mixture will be consumed. Generally, 14 days after ending treatment with the weak CYP3A inhibitor or upon your side-effects ending, your doctor will increase your dose to its previous level. Your doctor will provide you with specific guidance.



30 mi
V
Fill to 5 mL

Step 7: Take the 5-mL Zokinvy and orange juice mixture from the TAKE WITHIN medicine cup: with food within about 10 minutes of preparing Each dose must be mixed and consumed within 10 minutes. The mixture should only be prepared at the time it is to be consumed. Step 8: Fill the medicine cup used to consume the Zokinvy and orange juice mixture with 5 mL of water. Fill to 5 mL Step 9: Gently swirl the water around the medicine cup. Consume the water.

Drink lots of water while taking Zokinvy

It is important to drink lots of water and other liquids while taking Zokinvy. This may help to reduce problems linked to diarrhoea or vomiting.

Ask your doctor about the amount of water or other liquids that you should drink each day.

Your doctor will discuss with you which liquids you can drink to make sure that you are getting the correct amount of fluids each day.

Do not eat foods or drink juices that contain grapefruit, cranberries, pomegranates or Seville oranges (known as sour or bitter oranges).

If you take more Zokinvy than you should

If you take more capsules than you should, stop taking the medicine and contact your doctor.

If you forget to take Zokinvy

If you forget to take a dose, and 8 hours or more remain until your next planned dose, take the missed dose as soon as possible with some food. If less than 8 hours remains before the next scheduled dose, skip the missed dose and resume taking Zokinvy at the next scheduled dose.

If you stop taking Zokinvy

Do not stop taking Zokinvy without talking to your doctor.

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

4. Possible side effects

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

Serious side effects. Tell your doctor immediately if you:

• have persistent nausea, vomiting or diarrhoea that leads to loss of appetite, weight loss or dehydration. Vomiting or diarrhoea are very common (may affect more than 1 in 10 people) and may lead to electrolyte deficiencies requiring supportive care. Your doctor may monitor your weight, appetite and how much you eat and drink to help detect any of these possible electrolyte conditions.

Tell your doctor immediately if you notice any of the serious side effects above.

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

- increased liver enzymes shown by blood tests, which indicate stress on the liver
- stomach pain
- fatigue (tiredness)
- constipation
- sinus infections or other upper respiratory tract infections
- decreased haemoglobin shown by blood tests
- decreased bicarbonate shown by blood tests

Common (may affect up to 1 in 10 people)

- body aches and body pain, including back pain and pain in extremity
- fever
- decease in levels of sodium, potassium, albumin, creatinine shown by blood tests
- increase in levels of magnesium shown by blood tests
- cough
- flatulence
- rash
- pruritus (itchy skin)
- cerebral ischaemia (stroke)
- headache
- runny nose
- nasal congestion (blocked nose)
- nosebleed
- sore throat
- depression
- oral pustule (mouth sores)
- painful boil-like swelling near the anus (perirectal abscess)
- pneumonia
- influenza
- decrease in blood cell counts (such as white blood cell count) shown by blood tests
- tingling in hands and feet
- dizziness
- irritation, inflammation or ulcers of the large intestine (colitis)
- indigestion (may include feelings of bloating, discomfort, feeling too full or gas)
- inflammation of the stomach lining (gastritis)
- bleeding of the large intestine, rectum or anus

- dry skin
- skin darkening (hyperpigmentation)
- chest pain
- chills
- tooth fracture

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 via the national reporting system listed in Appendix V. By reporting side effects you help provide more information on the safety of this medicine.

5. How to store Zokinvy

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 carton and on the bottle after "EXP". The expiry date refers to the last day of that month.

Store in the original package. Keep the bottle tightly closed in order to protect from moisture. This medicinal product does not require any special temperature storage conditions.

Do not throw away any medicines via wastewater or household waste. Ask your pharmacist how to throw away medicines you no longer use. These measures will help protect the environment.

6. Contents of the pack and other information

What Zokinvy contains

- The active substance is lonafarnib

Zokinvy 50 mg hard capsules: each capsule contains 50 mg lonafarnib. Zokinvy 75 mg hard capsules: each capsule contains 75 mg lonafarnib.

- The other ingredients are:

<u>Capsule contents</u>: croscarmellose sodium (see section 2 "Zokinvy contains sodium"), magnesium stearate, poloxamer, povidone and silica, colloidal anhydrous

Capsule shell:

Zokinvy 50 mg hard capsules: gelatin, titanium dioxide, yellow iron oxide and sunflower lecithin Zokinvy 75 mg hard capsules: gelatin, titanium dioxide, yellow iron oxide, red iron oxide and sunflower lecithin

Printing ink: shellac, iron oxide black

What Zokinvy looks like and contents of the pack

Zokinvy 50 mg hard capsules are opaque yellow hard capsules, marked with "LNF" and "50" in black ink.

Zokinvy 75 mg hard capsules are opaque light orange hard capsules, marked with "LNF" and "75" in black ink.

The bottle pack contains 30 hard capsules and a desiccant. The desiccant is in a cannister and that cannister is included in the bottle, containing the capsules.

Marketing Authorisation Holder

EigerBio Europe Ltd. 1 Castlewood Avenue Rathmines, D06 H685, Ireland

Manufacturer

Patheon, part of Thermo Fisher Scientific Inc. 2100 Syntex Court Mississauga, Ontario L5N 7K9 Canada

This leaflet was last revised in MONTH YEAR

This medicine has been authorised under 'exceptional circumstances'.

This means that because of the rarity of this disease it has been impossible to get complete information on this medicine.

The European Medicines Agency will review any new information on this medicine every year and this leaflet will be updated as necessary.

Other sources of information

Detailed information on this medicine is available on the European Medicines Agency web site: http://www.ema.europa.eu. There are also links to other websites about rare diseases and treatments. This leaflet is available in all EU/EEA languages on the European Medicines Agency website.

ANNEX IV

CONCLUSIONS ON THE GRANTING OF THE MARKETING AUTHORISATION UNDER EXCEPTIONAL CIRCUMSTANCES PRESENTED BY THE EUROPEAN MEDICINES AGENCY

Conclusions presented by the European Medicines Agency on:

• Marketing authorisation under exceptional circumstances

The CHMP having considered the application is of the opinion that the risk-benefit balance is favourable to recommend the granting of the marketing authorisation under exceptional circumstances as further explained in the European Public Assessment Report.