

LenvimaTM 4,10mg

Lenvatinib

Hard capsules

1. NAME OF THE MEDICINAL PRODUCT

LENVIMA 4 mg hard capsules
LENVIMA 10 mg hard capsules

2. QUALITATIVE AND QUANTITATIVE COMPOSITION

LENVIMA 4 mg hard capsules

Each hard capsule contains 4 mg of lenvatinib (as mesilate).

LENVIMA 10 mg hard capsules

Each hard capsule contains 10 mg of lenvatinib (as mesilate).

For the full list of excipients, see section 6.1.

3. PHARMACEUTICAL FORM

Hard capsule.

LENVIMA 4 mg hard capsules

A yellowish-red body and yellowish-red cap, approximately 14.3 mm in length, marked in black ink with “C” on the cap, and “LENV 4 mg” on the body.

LENVIMA 10 mg hard capsules

A yellow body and yellowish-red cap, approximately 14.3 mm in length, marked in black ink with “C” on the cap, and “LENV 10 mg” on the body.

4. CLINICAL PARTICULARS

4.1 Therapeutic indications

LENVIMA is indicated for the treatment of adult patients with progressive, locally advanced or metastatic, differentiated (papillary/follicular/Hürthle cell) thyroid carcinoma (DTC), refractory to radioactive iodine (RAI).

LENVIMA is indicated as monotherapy for the treatment of adult patients with advanced or unresectable hepatocellular carcinoma (HCC) who have received no prior systemic therapy (see section 5.1).

4.2 Posology and method of administration

LENVIMA treatment should be initiated and supervised by a health care professional experienced in the use of anticancer therapies.

If a patient misses a dose, and it cannot be taken within 12 hours, then that dose should be skipped and the next dose should be taken at the usual time of administration.

Treatment should continue as long as clinical benefit is observed or until unacceptable toxicity occurs.

Optimal medical management (i.e. treatment or therapy) for nausea, vomiting, and diarrhoea should be initiated prior to any lenvatinib therapy interruption or dose reduction; gastrointestinal toxicity should be actively treated in order to reduce the risk of development of renal impairment or failure (see section 4.4, Renal failure and impairment).

Posology

Differentiated Thyroid Cancer (DTC)

The recommended daily dose of lenvatinib is 24 mg taken once daily. The daily dose is to be modified as needed according to the dose/toxicity management plan (see dose adjustment section below).

If a patient misses a dose, and it cannot be taken within 12 hours, then that dose should be skipped and the next dose should be taken at the usual time of administration.

Treatment should continue as long as clinical benefit is observed or until unacceptable toxicity occurs.

Dose adjustment

Management of adverse reactions may require dose interruption, adjustment, or discontinuation of lenvatinib (see section 4.4). Mild to moderate adverse reactions (e.g., Grade 1 or 2) generally do not warrant interruption of lenvatinib, unless intolerable to the patient despite optimal management. Severe (e.g., Grade 3) or intolerable adverse reactions require interruption of lenvatinib until resolution or improvement of the reaction, after which treatment should be resumed at a reduced dose as suggested in Table 1. LENVIMA should be discontinued in case of life-threatening reactions (e.g., Grade 4) with the exception of laboratory abnormality judged to be non-life-threatening, in which case they should be managed as severe reaction (e.g., Grade 3).

Grades are based on the National Cancer Institute (NCI) Common Terminology Criteria for Adverse Events (CTCAE).

Optimal medical management for nausea, vomiting, and diarrhoea should be initiated prior to any interruption or dose reduction of lenvatinib. Gastrointestinal toxicity should be actively managed in order to reduce the risk of development of renal impairment or failure (see section 4.4, Renal failure and impairment).

Table 1 Dose modifications from recommended daily dose

Dose level	Daily dose	Number of capsules
Recommended daily dose	24 mg orally once daily	Two 10 mg capsules plus one 4 mg capsule
First dose reduction	20 mg orally once daily	Two 10 mg capsules
Second dose reduction	14 mg orally once daily	One 10 mg capsule plus one 4 mg capsule
Third dose reduction	10 mg orally once daily ^a	One 10 mg capsule

a: Further dose reductions should be considered on an individual patient basis as limited data are available for doses below 10 mg.

Hepatocellular Carcinoma

The recommended daily dose of lenvatinib is 8 mg (two 4 mg capsules) once daily for patients with a body weight of < 60 kg and 12 mg (three 4 mg capsules) once daily for patients with a body weight of ≥ 60 kg. Dose adjustments are based only on toxicities observed and not on body weight changes during treatment. The daily dose is to be modified, as needed, according to the dose/toxicity management plan.

Dose adjustments and Discontinuation for HCC

Management of some adverse reactions may require dose interruption, adjustment, or discontinuation of lenvatinib therapy. Mild to moderate adverse reactions (e.g., Grade 1 or 2) generally do not warrant interruption of lenvatinib, unless intolerable to the patient despite optimal management. Details for monitoring, dose adjustment and discontinuation are provided in Table 2.

Table 2 Dose modifications from recommended lenvatinib daily dose in HCC patients

Starting Dose		≥60 kg BW 12 mg (three 4 mg capsules orally once daily)	<60 kg BW 8 mg (two 4 mg capsules orally once daily)
Persistent and Intolerable Grade 2 or Grade 3 Toxicities^a			
Adverse Reaction	Modification	Adjusted Dose ^b (≥60 kg BW)	Adjusted Dose ^b (<60 kg BW)
First occurrence ^c	Interrupt until resolved to Grade 0-1 or baseline ^d	8 mg (two 4 mg capsules) orally once daily	4 mg (one 4 mg capsule) orally once daily
Second occurrence (same reaction or new reaction)	Interrupt until resolved to Grade 0-1 or baseline ^d	4 mg (one 4 mg capsule) orally once daily	4 mg (one 4 mg capsule) orally every other day
Third occurrence (same reaction or new reaction)	Interrupt until resolved to Grade 0-1 or baseline ^d	4 mg (one 4 mg capsule) orally every other day	Discontinue
Life-threatening toxicities (Grade 4): Discontinue^e			
a. Initiate medical management for nausea, vomiting, or diarrhoea prior to interruption or dose reduction. b. Reduce dose in succession based on the previous dose level (12 mg, 8 mg, 4 mg or 4 mg every other day). c. Haematologic toxicity or proteinuria-no dose adjustment required for first occurrence. d. For haematologic toxicity, dosing can restart when resolved to Grade 2; proteinuria, resume when resolves to less than 2g/24 hours e. Excluding laboratory abnormalities judged to be nonlife-threatening, which should be managed as Grade 3.			

Grades are based on the National Cancer Institute (NCI) Common Terminology Criteria for Adverse Events (CTCAE).

Table 3 Adverse reactions requiring dose modification of lenvatinib in DTC and HCC

Adverse reaction	Severity	Action	Dose reduce and resume lenvatinib
Hypertension	Grade 3 (despite optimal antihypertensive therapy)	Interrupt	Resolves to Grade 0, 1 or 2. See detailed guidance in Table 3 in section 4.4.
	Grade 4	Discontinue	Do not resume
Proteinuria	≥ 2 g / 24 hours	Interrupt	Resolves to less than 2 g / 24 hours.
Nephrotic syndrome	-----	Discontinue	Do not resume

Table 3 Adverse reactions requiring dose modification of lenvatinib in DTC and HCC

Adverse reaction	Severity	Action	Dose reduce and resume lenvatinib
Renal impairment or failure	Grade 3	Interrupt	Resolves to Grade 0-1 or baseline.
	Grade 4*	Discontinue	Do not resume
Cardiac dysfunction	Grade 3	Interrupt	Resolves to Grade 0-1 or baseline.
	Grade 4	Discontinue	Do not resume
PRES/RPLS	Any grade	Interrupt	Consider resuming at reduced dose if resolves to Grade 0-1.
Hepatotoxicity	Grade 3	Interrupt	Resolves to Grade 0-1 or baseline.
	Grade 4*	Discontinue	Do not resume
Arterial thromboembolisms	Any grade	Discontinue	Do not resume
Haemorrhage	Grade 3	Interrupt	Resolves to Grade 0-1.
	Grade 4	Discontinue	Do not resume
GI perforation or fistula	Grade 3	Interrupt	Resolves to Grade 0-1 or baseline.
	Grade 4	Discontinue	Do not resume
Non-GI fistula	Grade 4	Discontinue	Do not resume
QT interval prolongation	>500 ms	Interrupt	Resolves to <480 ms or baseline
Diarrhoea	Grade 3	Interrupt	Resolves to Grade 0-1 or baseline.
	Grade 4 (despite medical management)	Discontinue	Do not resume

*Grade 4 laboratory abnormalities judged to be non-life-threatening, may be managed as severe reactions (e.g., Grade 3)

Special populations

Elderly population

DTC

Patients of age ≥ 75 years, of Asian race, with comorbidities (such as hypertension, and hepatic or renal impairment), or body weight below 60 kg appear to have reduced tolerability to lenvatinib (see section 4.8, Other special populations). All patients other than those with severe hepatic or renal impairment (see below) should initiate treatment at the recommended 24 mg dose, following which the dose should be further adjusted on the basis of individual tolerability.

HCC

Patients ≥ 75 years, of white race or female sex or those with worse baseline hepatic impairment (Child-Pugh A score of 6 compared to score of 5) appear to have reduced tolerability to lenvatinib.

HCC patients other than those with moderate and severe hepatic impairment or severe renal impairment should initiate treatment at the recommended starting dose of 8 mg (two 4 mg capsules) for body weight < 60 kg and 12 mg (three 4 mg capsules) for body weight ≥ 60 kg, following which the dose should be further adjusted on the basis of individual tolerability.

Patients with hypertension

Blood pressure should be well controlled prior to treatment with lenvatinib, and should be regularly monitored during treatment (see Table 4).

Table 4 Recommended management of hypertension

Blood Pressure (BP) level	Recommended action
Systolic BP ≥ 140 mmHg up to <160 mmHg or diastolic BP ≥ 90 mmHg up to <100 mmHg	Continue lenvatinib and initiate antihypertensive therapy, if not already receiving OR Continue lenvatinib and increase the dose of the current antihypertensive therapy or initiate additional antihypertensive therapy
Systolic BP ≥ 160 mmHg or diastolic BP ≥ 100 mmHg despite optimal antihypertensive therapy	1. Withhold lenvatinib 2. When systolic BP ≤ 150 mmHg, diastolic BP ≤ 95 mmHg, and patient has been on a stable dose of antihypertensive therapy for at least 48 hours, resume lenvatinib at a reduced dose
Life-threatening consequences (malignant hypertension, neurological deficit, or hypertensive crisis)	Urgent intervention is indicated. Discontinue lenvatinib and institute appropriate medical management.

Patient with Proteinuria

If urine dipstick proteinuria $\geq 2+$ is detected, dose interruptions, adjustments, or discontinuation may be necessary (see Table 1). LENVIMA should be discontinued in the event of nephrotic syndrome.

Patients with hepatic impairment

DTC

No adjustment of starting dose is required on the basis of hepatic function in patients with mild (Child-Pugh A) or moderate (Child-Pugh B) hepatic impairment. In patients with severe (Child-Pugh C) hepatic impairment, the recommended starting dose is 14 mg taken once daily. Further dose adjustments may be necessary on the basis of individual tolerability. Refer also to section 4.8, Other special populations.

HCC

In the patient populations enrolled in the HCC study no dose adjustments were required on the basis of hepatic function in those patients who had mild hepatic impairment (Child-Pugh A). The available very limited data are not sufficient to allow for a dosing recommendation for HCC patients with moderate hepatic impairment (Child-Pugh B). Close monitoring of overall safety is recommended in these patients (see sections 4.4 and 5.2). Lenvatinib has not been studied in patients with severe hepatic impairment (Child-Pugh C) and is not recommended for use in these patients.

Patients with renal impairment

DTC

No adjustment of starting dose is required on the basis of renal function in patients with mild or moderate renal impairment. In patients with severe renal impairment, the recommended starting dose is 14 mg taken once daily. Further dose adjustments may be necessary based on individual tolerability. Patients with end-stage renal disease were not studied, therefore the use of lenvatinib in these patients is not recommended. Refer also to section 4.8, Other special populations.

HCC

No dose adjustments are required on the basis of renal function in patients with mild or moderate renal impairment. The available data do not allow for a dosing recommendation for patients with HCC and severe renal impairment.

Elderly population

No adjustment of starting dose is required on the basis of age. Limited data are available on use in patients aged ≥ 75 years (see also section 4.8, Other special populations).

Paediatric population

Lenvatinib should not be used in children younger than 2 years of age because of safety concerns identified in animal studies (see section 5.3). The safety and efficacy of lenvatinib in children aged 2 to <18 years have not yet been established (see section 5.1). No data are available.

Race

No adjustment of starting dose is required on the basis of race (see section 5.2). Limited data are available on use in patients from ethnic origins other than Caucasian or Asian (see also section 4.8, Other special populations).

Method of administration

Lenvatinib is for oral use. The capsules should be taken at about the same time each day, with or without food (see section 5.2). The capsules should be swallowed whole with water. Caregivers should not open the capsule, in order to avoid repeated exposure to the contents of the capsule.

Alternatively, the lenvatinib capsules may be added without breaking or crushing them to a tablespoon of water or apple juice in a small glass to produce a suspension. The capsules must be left in the liquid for at least 10 minutes and stirred for at least 3 minutes to dissolve the capsule shells. The suspension is to be swallowed. After drinking, the same amount of water or apple juice (one tablespoon) must be added to the glass and swirled a few times. The additional liquid must be swallowed.

4.3 Contraindications

Hypersensitivity to the active substance or to any of the excipients listed in section 6.1.
Breast-feeding (see section 4.6).

4.4 Special warnings and precautions for use

Hypertension

Hypertension has been reported in patients treated with lenvatinib, usually occurring early in the course of treatment (see section 4.8, Description of selected adverse reactions). Blood pressure (BP) should be well controlled prior to treatment with lenvatinib and, if patients are known to be hypertensive, they should be on a stable dose of antihypertensive therapy for at least 1 week prior to treatment with lenvatinib. Serious complications of poorly controlled hypertension, including aortic dissection, have been reported. The early detection and effective management of hypertension are important to minimise the need for lenvatinib dose interruptions and reductions. Antihypertensive agents should be started as soon as elevated BP is confirmed. BP should be monitored after 1 week of treatment with lenvatinib, then every 2 weeks for the first 2 months, and monthly thereafter. The choice of antihypertensive treatment should be individualised to the

patient's clinical circumstances and follow standard medical practice. For previously normotensive subjects, monotherapy with one of the classes of antihypertensives should be started when elevated BP is observed. For those patients already on antihypertensive medication, the dose of the current agent may be increased, if appropriate, or one or more agents of a different class of antihypertensive should be added. When necessary, manage hypertension as recommended in Table 4.

Table 4 Recommended management of hypertension

Blood Pressure (BP) level	Recommended action
Systolic BP ≥ 140 mmHg up to <160 mmHg or diastolic BP ≥ 90 mmHg up to <100 mmHg	Continue lenvatinib and initiate antihypertensive therapy, if not already receiving OR Continue lenvatinib and increase the dose of the current antihypertensive therapy or initiate additional antihypertensive therapy
Systolic BP ≥ 160 mmHg or diastolic BP ≥ 100 mmHg despite optimal antihypertensive therapy	1. Withhold lenvatinib 2. When systolic BP ≤ 150 mmHg, diastolic BP ≤ 95 mmHg, and patient has been on a stable dose of antihypertensive therapy for at least 48 hours, resume lenvatinib at a reduced dose (see section 4.2)
Life-threatening consequences (malignant hypertension, neurological deficit, or hypertensive crisis)	Urgent intervention is indicated. Discontinue lenvatinib and institute appropriate medical management.

Proteinuria

Proteinuria has been reported in patients treated with lenvatinib, usually occurring early in the course of treatment (see section 4.8, Description of selected adverse reactions). Urine protein should be monitored regularly. If urine dipstick proteinuria $\geq 2+$ is detected, dose interruptions, adjustments, or discontinuation may be necessary (see section 4.2). Lenvatinib should be discontinued in the event of nephrotic syndrome.

Hepatotoxicity

In DTC, liver-related adverse reactions most commonly reported in patients treated with lenvatinib included increases in alanine aminotransferase (ALT), increases in aspartate - aminotransferase (AST), and increases in blood bilirubin. Hepatic failure and acute hepatitis ($<1\%$) have been reported in patients treated with lenvatinib. The hepatic failure cases were generally reported in patients with progressive liver metastases. Liver function tests should be monitored before initiation of treatment, then every 2 weeks for the first 2 months and monthly thereafter during treatment. In the case of hepatotoxicity, dose interruptions, adjustments, or discontinuation may be necessary.

If patients have severe hepatic impairment, the initial dose of lenvatinib should be adjusted

In HCC patients treated with lenvatinib in the REFLECT trial, liver-related adverse reactions including hepatic encephalopathy and hepatic failure (including fatal reactions) were reported at a higher frequency (see Section 4.8) compared to patients treated with sorafenib . Patients with

worse hepatic impairment and/or greater liver tumour burden at baseline had a higher risk of developing hepatic encephalopathy and hepatic failure. Hepatic encephalopathy also occurred more frequently in patients aged 75 years and older. Approximately half of the events of hepatic failure and one third of the events of the hepatic encephalopathy were reported in patients with disease progression.

Data in HCC patients with moderate hepatic impairment (Child-Pugh B) are very limited and there are currently no data available in HCC patients with severe hepatic impairment (Child-Pugh C). Since lenvatinib is mainly eliminated by hepatic metabolism, an increase in exposure in patients with moderate to severe hepatic impairment is expected.

Close monitoring of the overall safety is recommended in patients with mild or moderate hepatic impairment (see also sections 4.2 and 5.2). Liver function tests should be monitored before initiation of treatment, then every 2 weeks for the first 2 months and monthly thereafter during treatment. Patients with HCC should be monitored for worsening liver function including hepatic encephalopathy. In the case of hepatotoxicity, dose interruptions, adjustments, or discontinuation may be necessary (see section 4.2).

Renal failure and impairment

Renal impairment and renal failure have been reported in patients treated with lenvatinib (see section 4.8, Description of selected adverse reactions). The primary risk factor identified was dehydration and/or hypovolemia due to gastrointestinal toxicity. Gastrointestinal toxicity should be actively managed in order to reduce the risk of development of renal impairment or renal failure. Dose interruptions, adjustments, or discontinuation may be necessary (see section 4.2).

If patients have severe renal impairment, the initial dose of lenvatinib should be adjusted (see sections 4.2 and 5.2).

Diarrhoea

Diarrhoea has been reported frequently in patients treated with lenvatinib, usually occurring early in the course of treatment (see section 4.8, Description of selected adverse reactions). Prompt medical management of diarrhoea should be instituted in order to prevent dehydration. Lenvatinib should be discontinued in the event of persistence of Grade 4 diarrhoea despite medical management.

Cardiac dysfunction

Cardiac failure (<1%) and decreased left ventricular ejection fraction have been reported in patients treated with lenvatinib (see section 4.8, Description of selected adverse reactions). Patients should be monitored for clinical symptoms or signs of cardiac decompensation, as dose interruptions, adjustments, or discontinuation may be necessary (see section 4.2).

Posterior reversible encephalopathy syndrome (PRES) / Reversible posterior leucoencephalopathy syndrome (RPLS)

PRES, also known as RPLS, has been reported in patients treated with lenvatinib (<1%; see section 4.8, Description of selected adverse reactions). PRES is a neurological disorder which can present with headache, seizure, lethargy, confusion, altered mental function, blindness, and other visual or neurological disturbances. Mild to severe hypertension may be present. Magnetic resonance imaging is necessary to confirm the diagnosis of PRES. Appropriate measures should be taken to control blood pressure (see section 4.4, Hypertension). In patients with signs or symptoms of PRES, dose interruptions, adjustments, or discontinuation may be necessary (see section 4.2).

Arterial thromboembolisms

Arterial thromboembolisms (cerebrovascular accident, transient ischaemic attack, and myocardial infarction) have been reported in patients treated with lenvatinib (see section 4.8, Description of selected adverse reactions). Lenvatinib has not been studied in patients who have had an arterial thromboembolism within the previous 6 months, and therefore should be used with caution in such patients. A treatment decision should be made based upon an assessment of the individual patient's benefit/risk. Lenvatinib should be discontinued following an arterial thrombotic event.

Women of childbearing potential

Women of childbearing potential must use highly effective contraception while taking lenvatinib and for one month after stopping treatment (see section 4.6). It is currently unknown if lenvatinib increases the risk of thromboembolic events when combined with oral contraceptives.

Haemorrhage

Serious tumour related bleeds, including fatal haemorrhagic events have occurred in clinical trials and have been reported in post-marketing experience (see section 4.8, Description of selected adverse reactions). In post-marketing surveillance, serious and fatal carotid artery haemorrhages were seen more frequently in patients with anaplastic thyroid carcinoma (ATC) than in DTC or other tumour types. The degree of tumour invasion/infiltration of major blood vessels (e.g. carotid artery) should be considered because of the potential risk of severe haemorrhage associated with tumour shrinkage/necrosis following lenvatinib therapy. Some cases of bleeding have occurred secondarily to tumour shrinkage and fistula formation, e.g. tracheo-oesophageal fistulae. Cases of fatal intracranial haemorrhage have been reported in some patients with or without brain metastases. Bleeding in sites other than the brain (e.g. trachea, intra-abdominal, lung) has also been reported. One fatal case of hepatic tumour haemorrhage in a patient with HCC has been reported.

Screening for and subsequent treatment of oesophageal varices in patients with liver cirrhosis should be performed as per standard of care before starting treatment with lenvatinib

In the case of bleeding, dose interruptions, adjustments, or discontinuation may be required (see Section 4.2, Table 3).

Gastrointestinal perforation and fistula formation

Gastrointestinal perforation or fistulae have been reported in patients treated with lenvatinib (see section 4.8). In most cases, gastrointestinal perforation and fistulae occurred in patients with risk factors such as prior surgery or radiotherapy. In the case of a gastrointestinal perforation or fistula, dose interruptions, adjustments, or discontinuation may be necessary (see section 4.2).

Non-Gastrointestinal fistula

Patients may be at increased risk for the development of fistulae when treated with lenvatinib. Cases of fistula formation or enlargement that involve areas of the body other than stomach or intestines were observed in clinical trials and in post-marketing experience (e.g. tracheal, tracheo-oesophageal, oesophageal, cutaneous, female genital tract fistulae). Prior surgery and radiotherapy may be contributing risk factors. Lenvatinib should not be started in patients with fistula to avoid worsening and lenvatinib should be permanently discontinued in patients with oesophageal or tracheobronchial tract involvement and any Grade 4 fistula (see section 4.2); limited information is available on the use of dose interruption or reduction in management of other events, but worsening was observed in some cases and caution should be taken. Lenvatinib may adversely affect the wound healing process as for other agents of the same class.

QT interval prolongation

QT/QTc interval prolongation has been reported at a higher incidence in patients treated with lenvatinib than in patients treated with placebo (see section 4.8, Description of selected adverse reactions). Electrocardiograms should be monitored at baseline and periodically during treatment in all patients with particular attention to those with congenital long QT syndrome, congestive heart failure, bradyarrhythmias, and those taking medicinal products known to prolong the QT interval, including Class Ia and III antiarrhythmics. Lenvatinib should be withheld in the event of development of QT interval prolongation greater than 500 ms. Lenvatinib should be resumed at a reduced dose when QTc prolongation is resolved to < 480 ms or baseline.

Electrolyte disturbances such as hypokalaemia, hypocalcaemia, or hypomagnesaemia increase the risk of QT prolongation; therefore electrolyte abnormalities should be monitored and corrected in all patients before starting treatment. Electrolytes (magnesium, potassium and calcium) should be monitored periodically during treatment. Blood calcium levels should be monitored at least monthly and calcium should be replaced as necessary during lenvatinib treatment. Lenvatinib dose should be interrupted or dose adjusted as necessary depending on severity, presence of ECG changes, and persistence of hypocalcaemia.

Impairment of thyroid stimulating hormone suppression/ Thyroid dysfunction

Hypothyroidism has been reported in patients treated with lenvatinib (see section 4.8, Description of selected adverse reactions). Thyroid function should be monitored before initiation of, and periodically throughout, treatment with lenvatinib. Hypothyroidism should be treated according to standard medical practice to maintain euthyroid state.

Lenvatinib impairs exogenous thyroid suppression (see section 4.8, Description of selected adverse reactions). Thyroid stimulating hormone (TSH) levels should be monitored on a regular basis and thyroid hormone administration should be adjusted to reach appropriate TSH levels, according to the patient's therapeutic target.

Wound Healing Complications

No formal studies of the effect of lenvatinib on wound healing have been conducted. Impaired wound healing has been reported in patients receiving lenvatinib. Temporary interruption of lenvatinib should be considered in patients undergoing major surgical procedures. There is limited clinical experience regarding the timing of reinitiation of lenvatinib following a major surgical procedure. Therefore, the decision to resume lenvatinib following a major surgical procedure should be based on clinical judgment of adequate wound healing.

Special populations

Limited data are available for patients of ethnic origin other than Caucasian or Asian, and in patients aged ≥ 75 years. Lenvatinib should be used with caution in such patients, given the reduced tolerability of lenvatinib in Asian and elderly patients (see section 4.8, Other special populations).

There are no data on the use of lenvatinib immediately following sorafenib or other anticancer treatments and there may be a potential risk for additive toxicities unless there is an adequate washout period between treatments. The minimal washout period in clinical trials was 4 weeks.

4.5 Interaction with other medicinal products and other forms of interaction

Effect of other medicinal products on lenvatinib

Chemotherapeutic agents

Concomitant administration of lenvatinib, carboplatin, and paclitaxel has no significant impact on the pharmacokinetics of any of these 3 substances.

Effect of lenvatinib on other medicinal products

No data is available that can be used to exclude the risk that lenvatinib could be an inducer of CYP3A4 or Pgp in the gastrointestinal tract. This could potentially lead to decreased exposure to oral CYP3A4/Pgp substrates. This should be considered if co-administering oral CYP3A4/Pgp substrates for which retained efficacy is very important. CYP3A4 substrates known to have a narrow therapeutic index (e.g. astemizole, terfenadine, cisapride, pimozide, quinidine, bepridil or ergot alkaloids (ergotamine, dihydroergotamine)) should therefore be administered with caution in patients receiving lenvatinib.

Oral contraceptives

It is currently unknown whether lenvatinib may reduce the effectiveness of hormonal contraceptives, and therefore women using oral hormonal contraceptives should add a barrier method (see section 4.6).

4.6 Fertility, pregnancy and lactation

Women of childbearing potential

Women of childbearing potential should avoid becoming pregnant and use highly effective contraception while on treatment with lenvatinib and for at least one month after finishing treatment. It is currently unknown whether lenvatinib may reduce the effectiveness of hormonal contraceptives, and therefore women using oral hormonal contraceptives should add a barrier method.

Pregnancy

There are no data on the use of lenvatinib in pregnant women. Lenvatinib was embryotoxic and teratogenic when administered to rats and rabbits (see section 5.3).

Lenvatinib should not be used during pregnancy unless clearly necessary and after a careful consideration of the needs of the mother and the risk to the foetus.

Breast-feeding

It is not known whether lenvatinib is excreted in human milk. Lenvatinib and its metabolites are excreted in rat milk (see section 5.3). A risk to newborns or infants cannot be excluded and, therefore, lenvatinib is contraindicated during breast-feeding (see section 4.3).

Fertility

Effects in humans are unknown. However, testicular and ovarian toxicity has been observed in rats, dogs, and monkeys (see section 5.3).

4.7 Effects on ability to drive and use machines

Lenvatinib has a minor influence on the ability to drive and use machines, due to undesirable effects such as fatigue and dizziness. Patients who experience these symptoms should use caution when driving or operating machines.

4.8 Undesirable effects

Summary of the safety profile

The safety profile of lenvatinib is based on data from 452 DTC patients and 496 HCC patients; allowing characterisation only of common adverse drug reactions in DTC and HCC patients. The adverse reactions presented in this section are based on safety data of both DTC and HCC patients (see section 5.1).

DTC

The most frequently reported adverse reactions (occurring in $\geq 30\%$ of patients) are hypertension (68.6%), diarrhoea (62.8%), decreased appetite (51.5%), decreased weight (49.1%), fatigue (45.8%), nausea (44.5%), proteinuria 36.9%), stomatitis (35.8%), vomiting (34.5%), dysphonia (34.1%), headache (34.1%), and palmar-plantar erythrodysesthesia syndrome (PPE) (32.7%). Hypertension and proteinuria tend to occur early during lenvatinib treatment (see sections 4.4 and 4.8, Description of selected adverse reactions). The majority of Grade 3 to 4 adverse reactions occurred during the first 6 months of treatment except for diarrhoea, which occurred throughout treatment, and weight loss, which tended to be cumulative over time.

The most important serious adverse reactions were renal failure and impairment (2.4%), arterial thromboembolisms (3.9%), cardiac failure (0.7%), intracranial tumour haemorrhage (0.7%), PRES / RPLS (0.2%), hepatic failure (0.2%), and arterial thromboembolisms (cerebrovascular accident (1.1%), transient ischaemic attack (0.7%), and myocardial infarction (0.9%).

In 452 patients with RAI-refractory DTC, dose reduction and discontinuation were the actions taken for an adverse reaction in 63.1% and 19.5% of patients, respectively. Adverse reactions that most commonly led to dose reductions (in $\geq 5\%$ of patients) were hypertension, proteinuria, diarrhoea, fatigue, PPE, decreased weight, and decreased appetite. Adverse reactions that most commonly led to discontinuation of lenvatinib were proteinuria, asthenia, hypertension, cerebrovascular accident, diarrhoea, and pulmonary embolism.

HCC

The most frequently reported adverse reactions (occurring in $\geq 30\%$ of patients) are hypertension (44.0%), diarrhoea (38.1%), decreased appetite (34.9%), fatigue (30.6%), and decreased weight (30.4%).

The most important serious adverse reactions were hepatic failure (2.8%), hepatic encephalopathy (4.6%), oesophageal varices haemorrhage (1.4%), cerebral haemorrhage (0.6%), arterial thromboembolic events (2.0%) including myocardial infarction (0.8%), cerebral infarction (0.4%) and cerebrovascular accident (0.4%) and renal failure/impairment events (1.4%). There was a higher incidence of decreased neutrophil count in patients with HCC (8.7% on lenvatinib than in other non- HCC tumour types (1.4%)), which was not associated with infection, sepsis or bacterial peritonitis.

In 496 patients with HCC, dose modification (interruption or reduction) and discontinuation were the actions taken for an adverse reaction in 62.3% and 20.2% of patients, respectively. Adverse reactions that most commonly led to dose modifications (in $\geq 5\%$ of patients) were decreased appetite, diarrhoea, proteinuria, hypertension, fatigue, PPE and decreased platelet count. Adverse reactions that most commonly led to discontinuation of lenvatinib were hepatic encephalopathy, fatigue, increased blood bilirubin, proteinuria and hepatic failure.

Tabulated list of adverse reactions

Similar adverse reactions were observed in clinical trials in DTC and HCC.

Table 5 shows the frequency categories of adverse reactions observed in clinical trials in DTC and HCC. The adverse reaction frequency category represents the most conservative estimate of frequency from the two individual populations.

Frequencies are defined as:

- Very common ($\geq 1/10$)
- Common ($\geq 1/100$ to $< 1/10$)
- Uncommon ($\geq 1/1,000$ to $< 1/100$)
- Not known (cannot be estimated from the available data)

Within each frequency category, undesirable effects are presented in order of decreasing seriousness.

Table 5 Adverse reactions reported in patients in clinical trials

System Organ Class (MedDRA terminology*)	Very Common	Common	Uncommon	Not known
Infections and infestation	Urinary tract infection		Perineal abscess	
Blood and lymphatic disorders	Thrombocytopenia ^a Leukopenia ^a Neutropenia ^a	Lymphopenia ^a	Splenic infarction	
Endocrine disorders	Hypothyroidism	Increased blood thyroid stimulating hormone ‡		
Metabolism and nutrition disorders	Hypocalcaemia‡ Hypokalaemia Decreased weight Decreased appetite	Dehydration Hypomagnesaemia ^b Hypercholesterolaemia ^{a,b}		
Psychiatric disorders	Insomnia			
Nervous system disorders	Dizziness Headache Dysgeusia	Cerebrovascular accident †	Posterior reversible encephalopathy syndrome Monoparesis Transient ischaemic attack	
Cardiac disorders		Myocardial infarction ^{c,†} Cardiac failure Prolonged electrocardiogram QT Decreased ejection fraction		

System Organ Class (MedDRA terminology [*])	Very Common	Common	Uncommon	Not known
Vascular disorders	Haemorrhage ^{d, †, ‡} Hypertension ^{e, ‡} Hypotension		Aortic Dissection [*]	
Respiratory, thoracic and mediastinal disorders	Dysphonia	Pulmonary embolism ^{†, ‡}	Pneumothorax	
Gastrointestinal disorders	Diarrhoea Gastrointestinal and abdominal pains ^f Vomiting Nausea Oral inflammation ^g Oral pain ^h Constipation Dyspepsia Dry mouth	Anal fistula Flatulence Increased lipase Increased amylase	Pancreatitis ^{i, †}	
Hepatobiliary disorders	Increased blood bilirubin ^{j, ‡} Hypoalbuminaemia ^{j, ‡} Increased alanine aminotransferase [‡] Increased aspartate aminotransferase [‡]	Hepatic failure ^{k, ‡, †} Hepatic encephalopathy ^{l, ‡, †} Increased blood alkaline phosphatase Hepatic function abnormal Increased gamma-glutamyltransferase Cholecystitis	Hepatocellular damage/hepatitis	
Skin and subcutaneous tissue disorders	Palmar-plantar erythrodysesthesia syndrome Rash Alopecia	Hyperkeratosis		
Musculoskeletal and connective tissue disorders	Back pain Arthralgia Myalgia Pain in extremity Musculoskeletal pain			
Renal and urinary disorders	Proteinuria [‡]	Renal failure cases ^{n, †} Renal impairment Increased blood creatinine Increased blood urea	Nephrotic syndrome	
General disorders and administration site conditions	Fatigue Asthenia Peripheral oedema	Malaise	Impaired healing [*]	Non-gastrointestinal fistula ^o

^{*} Identified from post-marketing use of lenvatinib

[†]: Includes cases with a fatal outcome.

[‡]: See section 4.8 Description of selected adverse reactions for further characterisation.

The following terms have been combined:

- a: Thrombocytopenia includes thrombocytopenia and decreased platelet count. Neutropenia includes neutropenia and decreased neutrophil count decreased. Leukopenia includes leukopenia and decreased white blood cell count Lymphopenia includes lymphopenia and lymphocyte count decreased.
- b: Hypomagnesaemia includes hypomagnesaemia and decreased blood magnesium. Hypercholesterolaemia includes hypercholesterolaemia and increased blood cholesterol.
- c: Myocardial infarction includes myocardial infarction and acute myocardial infarction.
- d: Includes all haemorrhage terms.
Haemorrhage terms that occurred in 5 or more subjects with DTC were: epistaxis, haemoptysis, haematuria, contusion, haematochezia, gingival bleeding, petechial, pulmonary haemorrhage, rectal haemorrhage, blood urine present, haematoma and vaginal haemorrhage.
Haemorrhage terms that occurred in 5 or more subjects with HCC were: epistaxis, haematuria, gingival bleeding, haemoptysis, oesophageal varices haemorrhage, haemorrhoidal haemorrhage, mouth haemorrhage, rectal haemorrhage and upper gastrointestinal haemorrhage.
- e: Hypertension includes: hypertension, hypertensive crisis, increased diastolic blood pressure , , orthostatic hypertension, and increased blood pressure.
- f: Gastrointestinal and abdominal pain includes: abdominal discomfort, abdominal pain, abdominal pain lower, abdominal pain upper, abdominal tenderness, epigastric discomfort, and gastrointestinal pain.
- g: Oral inflammation includes: aphthous stomatitis, aphthous ulcer, gingival erosion, gingival ulceration, oral mucosal blistering, stomatitis, glossitis, mouth ulceration, and mucosal inflammation.
- h: Oral pain includes: oral pain, glossodynia, gingival pain, oropharyngeal discomfort, oropharyngeal pain and tongue discomfort.
- i: Pancreatitis includes: pancreatitis and acute pancreatitis.
- j: Hyperbilirubinaemia includes: hyperbilirubinaemia, increased blood bilirubin, jaundice and increased bilirubin conjugated. Hypoalbuminaemia includes hypoalbuminaemia and decreased blood albumin.
- k: Hepatic failure includes: hepatic failure, acute hepatic failure and chronic hepatic failure.
- l: Hepatic encephalopathy includes: hepatic encephalopathy, coma hepatic, metabolic encephalopathy and encephalopathy.
- m: Hepatocellular damage and hepatitis includes: drug-induced liver injury, hepatic steatosis, and cholestatic liver injury.
- m: Renal failure cases includes: acute prerenal failure, renal failure, renal failure acute, acute kidney injury and renal tubular necrosis.
- o: Non-gastrointestinal fistula includes cases of fistula occurring outside of the stomach and intestines such as tracheal, tracheo-oesophageal, oesophageal, female genital tract fistula, and cutaneous fistula.

Description of selected adverse reactions

DTC

Hypertension (see section 4.4)

In the pivotal Phase 3 SELECT trial (see section 5.1), hypertension (including hypertension, hypertensive crisis, increased diastolic blood pressure, and increased blood pressure) was reported in 72.8% of lenvatinib-treated patients and 16.0% of patients in the placebo-treated group. The median time to onset in lenvatinib-treated patients was 16 days. Reactions of Grade 3 or higher (including 1 reaction of Grade 4) occurred in 44.4% of lenvatinib-treated patients compared with 3.8% of placebo-treated patients. The majority of cases recovered or resolved following dose interruption or reduction, which occurred in 13.0% and 13.4% of patients, respectively. In 1.1% of patients, hypertension led to permanent treatment discontinuation.

HCC

In the Phase 3 -REFLECT trial (see section 5.1), hypertension (including hypertension, increased blood pressure, increased diastolic blood pressure and orthostatic hypertension) was reported in 44.5% of lenvatinib-treated patients and Grade 3 hypertension occurred in 23.5%. The median time to onset was 26 days. The majority of cases recovered following dose interruption or reduction, which occurred in 3.6% and 3.4% of patients respectively. One subject (0.2%) discontinued lenvatinib due to hypertension.

Proteinuria (see section 4.4)

DTC

In the pivotal Phase 3 SELECT trial (see section 5.1), proteinuria was reported in 33.7% of lenvatinib-treated patients and 3.1% of patients in the placebo-treated group. The median time to onset was 6.7 weeks. Grade 3 reactions occurred in 10.7% of lenvatinib-treated patients and none in placebo-treated patients. The majority of cases had an outcome of recovered or resolved following dose interruption or reduction, which occurred in 16.9% and 10.7% of patients, respectively. Proteinuria led to permanent treatment discontinuation in 0.8% of patients.

HCC

In the Phase 3 REFLECT trial (see section 5.1), proteinuria was reported in 26.3% of lenvatinib-treated patients and Grade 3 reactions occurred in 5.9%. The median time to onset was 6.1 weeks. The majority of cases recovered following dose interruption or reduction, which occurred in 6.9% and 2.5% of patients respectively. Proteinuria led to permanent treatment discontinuation in 0.6% of patients.

Renal failure and impairment (see section 4.4)

DTC

In the pivotal Phase 3 SELECT trial (see section 5.1), 5.0% of patients developed renal failure and 1.9% developed renal impairment (3.1% of patients had a Grade ≥ 3 event of renal failure or impairment). In the placebo group 0.8% of patients developed renal failure or impairment (0.8% were Grade ≥ 3).

HCC

In the Phase 3 REFLECT trial (see section 5.1), 7.1% of lenvatinib-treated patients developed a renal failure/impairment event. Grade 3 or greater reactions occurred in 1.9% of lenvatinib-treated patients.

Cardiac dysfunction (see section 4.4)

DTC

In the pivotal Phase 3 SELECT trial (see section 5.1), decreased ejection fraction/cardiac failure was reported in 6.5% of patients (1.5% were Grade ≥ 3) in the lenvatinib treated group, and 2.3% in the placebo group (none were Grade ≥ 3).

HCC

In the Phase 3 REFLECT trial (see section 5.1), cardiac dysfunction (including congestive cardiac failure, cardiogenic shock, and cardiopulmonary failure) was reported in 0.6% of patients (0.4% were Grade ≥ 3) in the lenvatinib-treated group.

Posterior reversible encephalopathy syndrome (PRES) / Reversible posterior leucoencephalopathy syndrome (RPLS) (see section 4.4)

DTC

In the pivotal Phase 3 SELECT trial (see section 5.1), there was 1 event of PRES (Grade 2) in the lenvatinib-treated group and no reports in the placebo group.

Amongst 1,166 patients treated with lenvatinib, there were 4 cases (0.3%) of PRES (0.3% were Grade 3 or 4), all of which resolved after treatment and/or dose interruption, or permanent discontinuation.

HCC

In the Phase 3 REFLECT trial (see section 5.1), there was 1 event of PRES (Grade 2) in the lenvatinib-treated group.

Amongst 1,823 patients treated with lenvatinib monotherapy in clinical trials, there were 5 cases (0.3%) of PRES (0.2% were Grade 3 or 4), all of which resolved after treatment and/or dose interruption, or permanent discontinuation.

Hepatotoxicity (see section 4.4)

DTC

In the pivotal Phase 3 SELECT trial (see section 5.1), the most commonly reported liver-related adverse reactions were hypoalbuminaemia (9.6% lenvatinib vs. 1.5% placebo) and elevations of liver enzyme levels, including increases in alanine aminotransferase (7.7% lenvatinib vs. 0 placebo), aspartate aminotransferase (6.9% lenvatinib vs. 1.5% placebo), and blood bilirubin (1.9% lenvatinib vs. 0 placebo). The median time to onset of liver reactions in lenvatinib-treated patients was 12.1 weeks. Liver-related reactions of Grade 3 or higher (including 1 Grade 5 case of hepatic failure) occurred in 5.4% of lenvatinib-treated patients compared with 0.8% in placebo-treated patients. Liver-related reactions led to dose interruptions and reductions in 4.6% and 2.7% of patients, respectively, and to permanent discontinuation in 0.4%.

Amongst 1,166 patients treated with lenvatinib, there were 3 cases (0.3%) of hepatic failure, all with a fatal outcome. One occurred in a patient with no liver metastases. There was also a case of acute hepatitis in a patient without liver metastases.

HCC

In the Phase 3 REFLECT trial (see section 5.1), the most commonly reported hepatotoxicity adverse reactions were increased blood bilirubin (14.9%), increased aspartate aminotransferase (13.7%), increased alanine aminotransferase (11.1%), hypoalbuminaemia (9.2%), hepatic

encephalopathy (8.0%), increased gamma-glutamyltransferase (7.8%) and increased blood alkaline phosphatase (6.7%). The median time to onset of hepatotoxicity adverse reactions was 6.4 weeks. Hepatotoxicity reactions of $\geq$ Grade 3 occurred in 26.1% of lenvatinib-treated patients. Hepatic failure (including fatal events in 12 patients) occurred in 3.6% of patients (all were $\geq$ Grade 3). Hepatic encephalopathy (including fatal events in 4 patients) occurred in 8.4% of patients (5.5% were $\geq$ Grade 3). There were 17 (3.6%) deaths due to hepatotoxicity events in the lenvatinib arm and 4 (0.8%) deaths in the sorafenib arm. Hepatotoxicity adverse reactions led to dose interruptions and reductions in 12.2% and 7.4% of lenvatinib-treated patients respectively, and to permanent discontinuation in 5.5%.

Across clinical studies in which 1327 patients received lenvatinib monotherapy in indications other than HCC, hepatic failure (including fatal events) was reported in 4 patients (0.3%), liver injury in 2 patients (0.2%), acute hepatitis in 2 patients (0.2%), and hepatocellular injury in 1 patient (0.1%).

Arterial thromboembolisms (see section 4.4)

DTC

In the pivotal Phase 3 SELECT trial (see section 5.1), arterial thromboembolic events were reported in 5.4% of lenvatinib-treated patients and 2.3% of patients in the placebo group.

Amongst 1,166 patients treated with lenvatinib, there were 5 cases (0.4%) of arterial thromboembolisms (3 cases of myocardial infarction and 2 cases of cerebrovascular accident) with a fatal outcome.

HCC

In the Phase 3 REFLECT trial (see section 5.1), arterial thromboembolic events were reported in 2.3% of patients treated with lenvatinib.

Amongst 1,823 patients treated with lenvatinib monotherapy in clinical studies, there were 10 cases (0.5%) of arterial thromboembolisms (5 cases of myocardial infarction and 5 cases of cerebrovascular accident) with a fatal outcome.

Haemorrhage (see section 4.4)

DTC

In the pivotal Phase 3 SELECT trial (see section 5.1), haemorrhage was reported in 34.9% (1.9% were Grade $\geq$ 3) of lenvatinib-treated patients versus 18.3% (3.1% were Grade $\geq$ 3) of placebo-treated patients. Reactions that occurred at an incidence of $\geq$ 0.75% above placebo were: epistaxis (11.9%), haematuria (6.5%), contusion (4.6%), gingival bleeding (2.3%), haematochezia (2.3%), rectal haemorrhage (1.5%), haematoma (1.1%), haemorrhoidal haemorrhage (1.1%), laryngeal haemorrhage (1.1%), petechiae (1.1%), and intracranial tumour haemorrhage (0.8%). In this trial, there was 1 case of fatal intracranial haemorrhage among 16 patients who received lenvatinib and had CNS metastases at baseline.

The median time to first onset in lenvatinib-treated patients was 10.1 weeks. No differences between lenvatinib- and placebo-treated patients were observed in the incidences of serious reactions (3.4% vs. 3.8%), reactions leading to premature discontinuation (1.1% vs. 1.5%), or reactions leading to dose interruption (3.4% vs. 3.8%) or reduction (0.4% vs. 0).

Amongst 1,166 patients treated with lenvatinib, Grade 3 or greater haemorrhage was reported in 2% of patients, 3 patients (0.3%) had a Grade 4 haemorrhage and 5 patients (0.4%) had a Grade 5 reaction including arterial haemorrhage, haemorrhagic stroke, intracranial tumour haemorrhage, haemoptysis and tumour haemorrhage.

HCC

In the Phase 3 REFLECT trial (see section 5.1), haemorrhage was reported in 24.6% of patients and 5.0% were Grade ≥ 3 . Grade 3 reactions occurred in 3.4%, Grade 4 reactions in 0.2% and 7 patients (1.5%) had a grade 5 reaction including cerebral haemorrhage, upper gastrointestinal haemorrhage, intestinal haemorrhage and tumour haemorrhage. The median time to first onset was 11.9 weeks. A haemorrhage event led to dose interruption or reduction in 3.2% and 0.8% patients respectively and to treatment discontinuation in 1.7% of patients.

Across clinical studies in which 1,327 patients received lenvatinib monotherapy in indications other than HCC, Grade ≥ 3 or greater haemorrhage was reported in 2% of patients, 3 patients (0.2%) had a Grade 4 haemorrhage and 8 patients (0.6%) had a Grade 5 reaction including arterial haemorrhage, haemorrhagic stroke, intracranial haemorrhage, intracranial tumour haemorrhage, haematemesis, melaena, haemoptysis and tumour haemorrhage.

Hypocalcaemia (see section 4.4, QT interval prolongation)

DTC

In the pivotal Phase 3 SELECT trial (see section 5.1), hypocalcaemia was reported in 12.6% of lenvatinib-treated patients vs. no cases in the placebo arm. The median time to first onset in lenvatinib-treated patients was 11.1 weeks. Reactions of Grade 3 or 4 severity occurred in 5.0% of lenvatinib-treated vs 0 placebo-treated patients. Most reactions resolved following supportive treatment, without dose interruption or reduction, which occurred in 1.5% and 1.1% of patients, respectively; 1 patient with Grade 4 hypocalcaemia discontinued treatment permanently.

HCC

In the Phase 3 REFLECT trial (see section 5.1), hypocalcaemia was reported in 1.1% of patients, with grade 3 reactions occurring in 0.4%. Lenvatinib dose interruption due to hypocalcaemia occurred in one subject (0.2%) and there were no dose reductions or discontinuations.

Gastrointestinal perforation and fistula formation (see section 4.4)

DTC

In the pivotal Phase 3 SELECT trial (see section 5.1), events of gastrointestinal perforation or fistula were reported in 1.9% of lenvatinib-treated patients and 0.8% of patients in the placebo group.

HCC

In the Phase 3 REFLECT trial (see section 5.1), events of gastrointestinal perforation or fistula were reported in 1.9% of lenvatinib-treated patients.

Non-Gastrointestinal fistulae (see section 4.4)

Lenvatinib use has been associated with cases of fistulae including reactions resulting in death. Reports of fistulae that involve areas of the body other than stomach or intestines were observed across various indications. Reactions were reported at various time points during treatment ranging from two weeks to greater than 1 year from initiation of lenvatinib, with median latency of about 3 months.

QT interval prolongation (see section 4.4)

DTC

In the pivotal Phase 3 SELECT trial (see section 5.1), QT/QTc interval prolongation was reported in 8.8% of lenvatinib-treated patients and 1.5% of patients in the placebo group. The incidence of QT interval prolongation of greater than 500 ms was 2% in the lenvatinib-treated patients compared to no reports in the placebo group.

HCC

In the Phase 3 REFLECT trial (see section 5.1), QT/QTc interval prolongation was reported in 6.9% of lenvatinib-treated patients. The incidence of QTcF interval prolongation of greater than 500ms was 2.4%.

Increased blood thyroid stimulating hormone (see section 4.4 Impairment of thyroid stimulating hormone suppression / Thyroid dysfunction)

DTC

In the pivotal Phase 3 SELECT trial (see section 5.1), 88% of all patients had a baseline TSH level less than or equal to 0.5 mU/L. In those patients with a normal TSH at baseline, elevation of TSH level above 0.5 mU/L was observed post baseline in 57% of lenvatinib-treated patients as compared with 14% of placebo-treated patients.

HCC

In the Phase 3 REFLECT trial (see section 5.1), 89.6% of patients had a baseline TSH level of less than the upper limit of normal. Elevation of TSH above the upper limit of normal was observed post baseline in 69.6% of lenvatinib-treated patients.

Diarrhoea (see section 4.4)

DTC

In the pivotal Phase 3 SELECT trial (see section 5.1), diarrhoea was reported in 67.4% of patients in the lenvatinib-treated group (9.2% were Grade $\geq$ 3) and in 16.8% of patients in the placebo group (none were Grade $\geq$ 3).

HCC

In the Phase 3 REFLECT trial (see section 5.1), diarrhoea was reported in 38.7% of patients treated with lenvatinib (4.2% were Grade $\geq$ 3).

Paediatric population

Clinical data are not available in this population (see section 4.2).

Other special populations

Elderly

DTC

Patients of age ≥ 75 years were more likely to experience Grade 3 or 4 hypertension, proteinuria, decreased appetite, and dehydration.

HCC

Patients of age ≥ 75 years were more likely to experience hypertension, proteinuria, decreased appetite, asthenia, dehydration, dizziness, malaise, peripheral oedema, pruritus and hepatic encephalopathy. Hepatic encephalopathy occurred at more than twice the incidence in patients aged ≥ 75 years (17.2%) than in those < 75 years (7.1%). Hepatic encephalopathy tended to be associated with adverse disease characteristics at baseline or with the use of concomitant medications. Arterial thromboembolic events also occurred at an increased incidence in this age group.

Gender

DTC

Females had a higher incidence of hypertension (including Grade 3 or 4 hypertension), proteinuria, and PPE, while males had a higher incidence of decreased ejection fraction and gastrointestinal perforation and fistula formation.

HCC

Females had a higher incidence of hypertension, fatigue, ECG QT prolongation and alopecia. Men had a higher incidence (26.5%) of dysphonia than women (12.3%), decreased weight and decreased platelet count. Hepatic failure events were observed in male patients only.

Ethnic origin

DTC

Asian patients had a higher incidence than Caucasian patients of peripheral oedema, hypertension, fatigue, PPE, proteinuria, thrombocytopenia, and increased blood thyroid stimulating hormone.

HCC

Asian patients had a higher incidence than Caucasian patients of proteinuria, decreased neutrophil count, decreased platelet count, decreased white blood count and PPE syndrome, while Caucasian patients had a higher incidence of fatigue, hepatic encephalopathy, acute kidney injury, anxiety, asthenia, nausea, thrombocytopenia and vomiting.

Baseline hypertension

DTC

Patients with baseline hypertension had a higher incidence of Grade 3 or 4 hypertension, proteinuria, diarrhoea, and dehydration, and experienced more serious cases of dehydration, hypotension, pulmonary embolism, malignant pleural effusion, atrial fibrillation, and GI symptoms (abdominal pain, diarrhoea, vomiting).

Hepatic impairment

DTC

Patients with baseline hepatic impairment had a higher incidence of hypertension and PPE, and a higher incidence of Grade 3 or 4 hypertension, asthenia, fatigue, and hypocalcaemia compared with patients with normal hepatic function.

HCC

Patients with a baseline Child Pugh (CP) score of 6 (about 20% patients in the REFLECT study) had a higher incidence of decreased appetite, fatigue, proteinuria, hepatic encephalopathy and hepatic failure compared to patients with a baseline CP score of 5. Hepatotoxicity events and haemorrhage events also occurred at a higher incidence in CP score 6 patients compared to CP score 5 patients.

Renal impairment

DTC

Patients with baseline renal impairment had a higher incidence of Grade 3 or 4 hypertension, proteinuria, fatigue, stomatitis, oedema peripheral, thrombocytopenia, dehydration, prolonged QT, hypothyroidism, hyponatraemia, increased blood thyroid stimulating hormone, pneumonia compared with subjects with normal renal function. These patients also had a higher incidence of renal reactions and a trend towards a higher incidence of liver reactions.

HCC

Patients with baseline renal impairment had a higher incidence of fatigue, hypothyroidism, dehydration, diarrhoea, decreased appetite, proteinuria and hepatic encephalopathy. These patients also had a higher incidence of renal reactions and arterial thromboembolic events.

Patients with body weight <60 kg

DTC

Patients with low body weight (<60 kg) had a higher incidence of PPE, proteinuria, of Grade 3 or 4 hypocalcaemia and hyponatraemia, and a trend towards a higher incidence of Grade 3 or 4 decreased appetite.

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

The highest doses of lenvatinib studied clinically were 32 mg and 40 mg per day. Accidental medication errors resulting in single doses of 40 to 48 mg have occurred in clinical trials. The most frequently observed adverse drug reactions at these doses were hypertension, nausea, diarrhoea, fatigue, stomatitis, proteinuria, headache, and aggravation of PPE. There have also been reports of overdose with lenvatinib involving single administrations of 6 to 10 times the recommended daily dose. These cases were associated with adverse reactions consistent with the known safety profile of lenvatinib (i.e., renal and cardiac failure), or were without adverse reactions.

Symptoms and Management

There is no specific antidote for overdose with lenvatinib. In case of suspected overdose, lenvatinib should be withheld and appropriate supportive care given as required.

5. PHARMACOLOGICAL PROPERTIES

5.1 Pharmacodynamic properties

Pharmacotherapeutic group: antineoplastic agents, protein kinase inhibitors, ATC code: L01XE29

Lenvatinib is a multikinase inhibitor which has shown mainly antiangiogenic properties in vitro and in vivo, and direct inhibition of tumour growth was also observed in in vitro models.

Mechanism of action

Lenvatinib is a receptor tyrosine kinase (RTK) inhibitor that selectively inhibits the kinase activities of vascular endothelial growth factor (VEGF) receptors VEGFR1 (FLT1), VEGFR2 (KDR), and VEGFR3 (FLT4), in addition to other proangiogenic and oncogenic pathway-related RTKs including fibroblast growth factor (FGF) receptors FGFR1, 2, 3, and 4, the platelet derived growth factor (PDGF) receptor PDGFR α , KIT, and RET.

In addition, lenvatinib had selective, direct antiproliferative activity in hepatocellular cell lines dependent on activated FGFR signalling, which is attributed to the inhibition of FGFR signalling by lenvatinib.

Although not studied directly with lenvatinib, the mechanism of action (MOA) for hypertension is postulated to be mediated by the inhibition of VEGFR2 in vascular endothelial cells. Similarly, although not studied directly, the MOA for proteinuria is postulated to be mediated by downregulation of VEGFR1 and VEGFR2 in the podocytes of the glomerulus.

The mechanism of action for hypothyroidism is not fully elucidated.

Clinical efficacy

Radioiodine-refractory differentiated thyroid cancer

The SELECT study was a multicentre, randomised, double-blind, placebo-controlled trial that was conducted in 392 patients with radioiodine-refractory differentiated thyroid cancer with independent, centrally reviewed, radiographic evidence of disease progression within 12 months (+1 month window) prior to enrollment. Radioiodine-refractory was defined as one or more measurable lesions either with a lack of iodine uptake or with progression in spite of radioactive-iodine (RAI) therapy, or having a cumulative activity of RAI of >600 mCi or 22 GBq with the last dose at least 6 months prior to study entry. Randomisation was stratified by geographic region (Europe, North America, and Other), prior VEGF/VEGFR-targeted therapy (patients may have received 0 or 1 prior VEGF/VEGFR-targeted therapy), and age (≤ 65 years or > 65 years). The main efficacy outcome measure was progression-free survival (PFS) as determined by blinded independent radiologic review using Response Evaluation Criteria in Solid Tumours (RECIST) 1.1. Secondary efficacy outcome measures included overall response rate and overall survival. Patients in the placebo arm could opt to receive lenvatinib treatment at the time of confirmed disease progression.

Eligible patients with measurable disease according to RECIST 1.1 were randomised 2:1 to receive lenvatinib 24 mg once daily (n=261) or placebo (n=131). Baseline demographics and disease characteristics were well balanced for both treatment groups. Of the 392 patients randomised, 76.3% were naïve to prior VEGF/VEGFR-targeted therapies, 49.0% were female, 49.7% were European, and the median age was 63 years. Histologically, 66.1% had a confirmed diagnosis of papillary thyroid cancer and 33.9% had follicular thyroid cancer which included Hürthle cell 14.8% and clear cell 3.8%. Metastases were present in 99% of the patients: lungs in

89.3%, lymph nodes in 51.5%, bone in 38.8%, liver in 18.1%, pleura in 16.3%, and brain in 4.1%. The majority of patients had an ECOG performance status of 0; 42.1% had a status of 1; 3.9% had a status above 1. The median cumulative RAI activity administered prior to study entry was 350 mCi (12.95 GBq).

A statistically significant prolongation in PFS was demonstrated in lenvatinib-treated patients compared with those receiving placebo ($p < 0.0001$) (see figure 1). The positive effect on PFS was seen across the subgroups of age (above or below 65 years), sex, race, histological subtype, geographic region, and those who received 0 or 1 prior VEGF/VEGFR-targeted therapies. Following independent review confirmation of disease progression, 109 (83.2%) patients randomised to placebo had crossed over to open-label lenvatinib at the time of the primary efficacy analysis.

The objective response rate (complete response [CR] plus partial response [PR]) per independent radiological review was significantly ($p < 0.0001$) higher in the lenvatinib-treated group (64.8%) than in the placebo-treated group (1.5%). Four (1.5%) subjects treated with lenvatinib attained a CR and 165 subjects (63.2%) had a PR, while no subjects treated with placebo had a CR and 2 (1.5%) subjects had a PR.

The median time to first dose reduction was 2.8 months. The median time to objective response was 2.0 (95% CI: 1.9, 3.5) months; however, of the patients who experienced a complete or partial response to lenvatinib, 70.4% were observed to develop the response on or within 30 days of being on the 24-mg dose.

The overall survival analysis was confounded by the fact that placebo-treated subjects with confirmed disease progression had the option to cross over to open-label lenvatinib. There was no statistically significant difference in overall survival between the treatment groups at the time of the primary efficacy analysis (HR=0.73; 95%CI: 0.50, 1.07, $p = 0.1032$). The median OS had not been reached for either the lenvatinib group or the placebo crossover group.

Table 6 Efficacy results in DTC patients

	Lenvatinib N=261	Placebo N=131
Progression-Free Survival (PFS)^a		
Number of progressions or deaths (%)	107 (41.0)	113 (86.3)
Median PFS in months (95% CI)	18.3 (15.1, NE)	3.6 (2.2, 3.7)
Hazard ratio (99% CI) ^{b,c}	0.21 (0.14, 0.31)	
P-value ^b	<0.0001	
Patients who had received 0 prior VEGF/VEGFR-targeted therapy (%)	195 (74.7)	104 (79.4)
Number of progressions or deaths	76	88
Median PFS in months (95% CI)	18.7 (16.4, NE)	3.6 (2.1, 5.3)
Hazard ratio (95% CI) ^{b,c}	0.20 (0.14, 0.27)	
Patients who had received 1 prior VEGF/VEGFR-targeted therapy (%)	66 (25.3)	27 (20.6)
Number of progressions or deaths	31	25
Median PFS in months (95% CI)	15.1 (8.8, NE)	3.6 (1.9, 3.7)
Hazard ratio (95% CI) ^{b,c}	0.22 (0.12, 0.41)	
Objective Response Rate^a		
Number of objective responders (%)	169 (64.8)	2 (1.5)
(95% CI)	(59.0, 70.5)	(0.0, 3.6)
P-value ^b	<0.0001	
Number of complete responses	4	0
Number of partial responses	165	2
Median time to objective response, ^d months (95% CI)	2.0 (1.9, 3.5)	5.6 (1.8, 9.4)
Duration of response, ^d months, median (95% CI)	NE (16.8, NE)	NE (NE, NE)

Overall Survival		
Number of deaths (%)	71 (27.2)	47 (35.9)
Median OS in months (95% CI)	NE (22.0, NE)	NE (20.3, NE)
Hazard ratio (95% CI) ^{b, e}	0.73 (0.50, 1.07)	
P-value ^{b, e}	0.1032	

CI, confidence interval; NE, not estimable; OS, overall survival; PFS, progression-free survival; RPSFT, rank preserving structural failure time model; VEGF/VEGFR, vascular endothelial growth factor / vascular endothelial growth factor receptor.

a: Independent radiologic review.

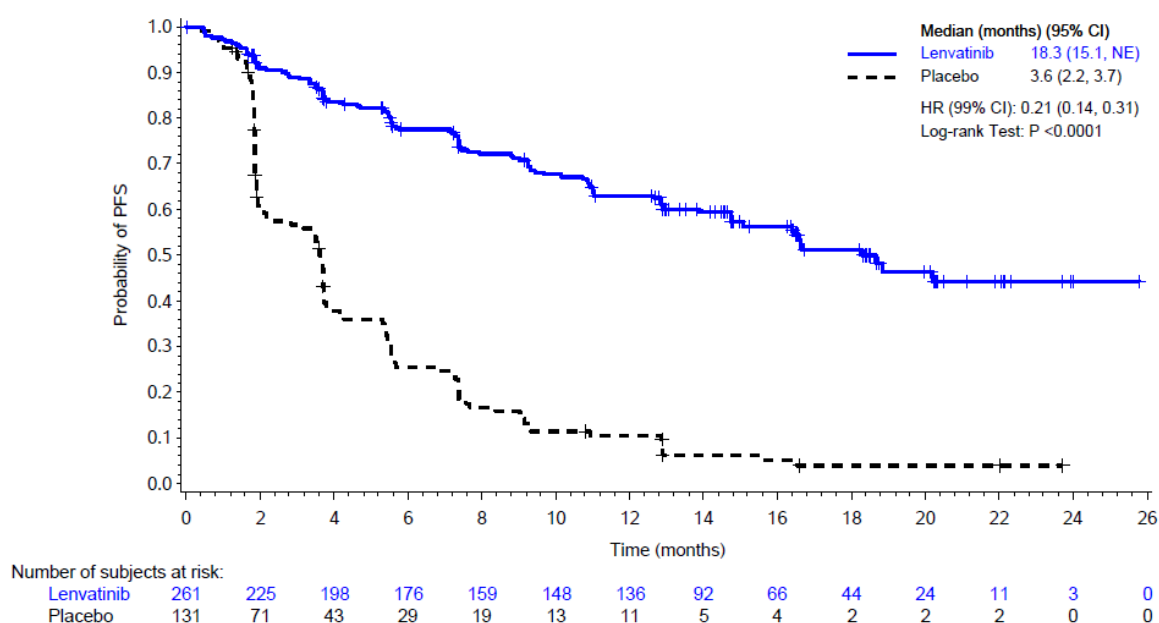
b: Stratified by region (Europe vs. North America vs. Other), age group (≤ 65 years vs > 65 years), and previous VEGF/VEGFR-targeted therapy (0 vs. 1).

c: Estimated with Cox proportional hazard model.

d: Estimated using the Kaplan-Meier method; the 95% CI was constructed with a generalised Brookmeyer and Crowley method in patients with a best overall response of complete response or partial response.

e: Not adjusted for crossover effect.

Figure 1 Kaplan-Meier Curve of Progression-Free Survival - DTC



CI, confidence interval; NE, not estimable.

Hepatocellular Carcinoma

The clinical efficacy and safety of lenvatinib have been evaluated in an international, multicenter, open-label, randomised phase 3 study (REFLECT) in patients with unresectable hepatocellular carcinoma (HCC).

In total, 954 patients were randomised 1:1 to receive either lenvatinib (12 mg [baseline body weight ≥ 60 kg] or 8 mg [baseline body weight < 60 kg]) given orally once daily or sorafenib 400 mg given orally twice daily.

Patients were eligible to participate if they had a liver function status of Child-Pugh class A and Eastern Cooperative Oncology Group Performance Status (ECOG PS) 0 or 1. Patients were excluded who had prior systemic anticancer therapy for advanced/unresectable HCC or any prior anti-VEGF therapy. Target lesions previously treated with radiotherapy or locoregional therapy had to show radiographic evidence of disease progression. Patients with $\geq 50\%$ liver occupation, clear invasion into the bile duct or a main branch of the portal vein (Vp4) on imaging were also excluded.

- Demographic and baseline disease characteristics were similar between the lenvatinib and the sorafenib groups and are shown below for all 954 randomised patients:
- Median age: 62 years
- Male: 84%
- White: 29%, Asian: 69%, Black or African American: 1.4%
- Body weight: <60 kg -31%, 60-80 kg – 50%, >80 kg - 19%
- Eastern Cooperative Oncology Group Performance Status (ECOG PS) of 0: 63%, ECOG PS of 1: 37%
- Child-Pugh A: 99%, Child-Pugh B: 1%
- Aetiology: Hepatitis B (50%), Hepatitis C (23%), alcohol (6%)
- Absence of macroscopic portal vein invasion (MPVI): 79%
- Absence of MPVI, extra-hepatic tumour spread (EHS) or both: 30%
- Underlying cirrhosis (by independent imaging review): 75%
- Barcelona Clinic Liver Cancer (BCLC) stage B: 20%; BCLC stage C: 80%
- Prior treatments: hepatectomy (28%), radiotherapy (11%), loco-regional therapies including transarterial (chemo)embolisation (52%), radiofrequency ablation (21%) and percutaneous ethanol injection (4%)

The primary efficacy endpoint was Overall Survival (OS). Lenvatinib was non-inferior for OS to sorafenib with HR = 0.92 [95% CI of (0.79, 1.06)] and a median OS of 13.6 months vs 12.3 months (see Table 7 and Figure 2). The results for surrogate endpoints (PFS and ORR) are presented in Table 7 below.

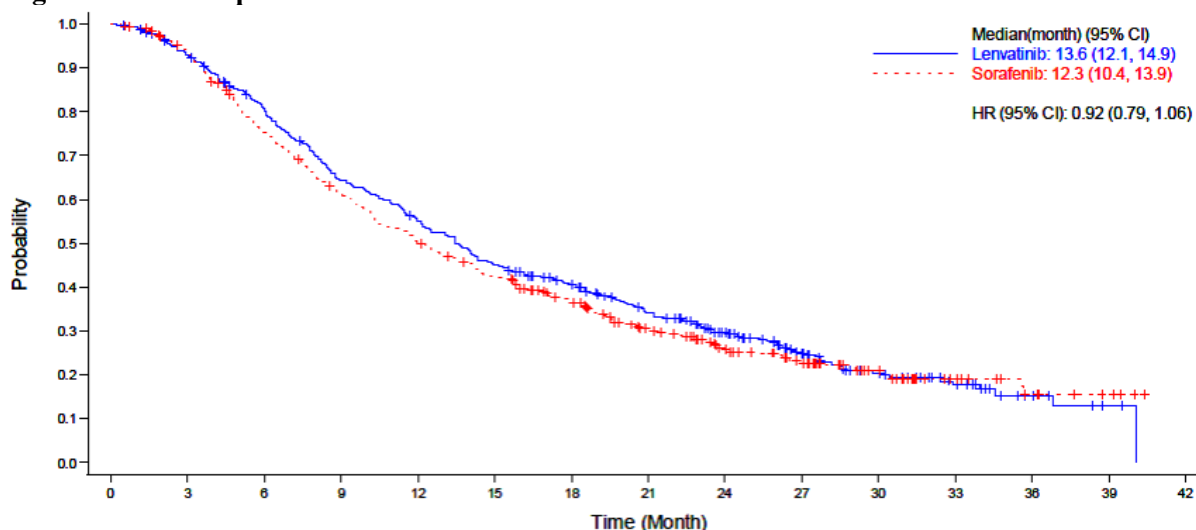
Table 7: Efficacy Results from the REFLECT study in HCC

Efficacy parameter	Hazard ratio ^{a, b} (95% CI)	P-value ^d	Median (95% CI) ^e	
			Lenvatinib (N= 478)	Sorafenib (N=476)
OS	0.92 (0.79,1.06)	NA	13.6 (12.1, 14.9)	12.3 (10.4, 13.9)
PFS ^g (mRECIST)	0.64 (0.55, 0.75)	<0.00001	7.3 (5.6, 7.5)	3.6 (3.6, 3.7)
			Percentages (95% CI)	
ORR ^{c, f, g} (mRECIST)	NA	<0.00001	41% (36%, 45%)	12% (9%, 15%)

Data cut-off date: 13 Nov 2016.

- Hazard ratio is for lenvatinib vs. sorafenib, based on a Cox model including treatment group as a factor.
- Stratified by region (Region 1: Asia-Pacific; Region 2: Western), macroscopic portal vein invasion or extrahepatic spread or both (yes, no), ECOG PS (0, 1) and body weight (<60 kg, ≥60 kg).
- Results are based on confirmed and unconfirmed responses.
- P-value is for the superiority test of lenvatinib versus sorafenib.
- Quartiles are estimated by the Kaplan-Meier method, and the 95% CIs are estimated with a generalized Brookmeyer and Crowley method
- Response rate (complete or partial response)
- Per independent radiology review retrospective analysis. The median duration of objective response was 7.3 (95% CI 5.6, 7.4) months in the lenvatinib arm and 6.2 (95% CI 3.7, 11.2) months in the sorafenib arm.

Figure 2 Kaplan-Meier Curve of Overall Survival - HCC



Number of subjects at risk:

Lenvatinib	478	436	374	297	253	207	178	140	102	67	40	21	8	2	0
Sorafenib	476	440	348	282	230	192	156	116	83	57	33	16	8	4	0

1. Data cut-off date = 13 Nov 2016.
2. Noninferiority margin for hazard ratio (HR: lenvatinib vs sorafenib = 1.08).
3. Median was estimated with the Kaplan-Meier method and the 95% confidence interval was constructed with a generalised Brookmeyer and Crowley method.
4. HR was estimated from the Cox proportional hazard model with treatment as independent variable and stratified by IxRS stratification factors. The Efron method was used for ties.
5. += censored observations.

In subgroup analyses by stratification factors (presence or absence of MPVI or EHS or both, ECOG PS 0 or 1, BW <60 kg or ≥60 kg and region) the HR consistently favoured lenvatinib over sorafenib, with the exception of Western region [HR of 1.08 (95% CI 0.82, 1.42)], patients without EHS [HR of 1.01 (95% CI 0.78, 1.30)] and patients without MPVI, EHS or both [HR of 1.05 (0.79, 1.40)]. The results of subgroup analyses should be interpreted with caution.

The median duration of treatment was 5.7 months (Q1: 2.9, Q3: 11.1) in the lenvatinib arm and 3.7 months (Q1: 1.8, Q3: 7.4) in the sorafenib arm.

In both treatment arms in the REFLECT study, median OS was approximately 9 months longer in subjects who received post-treatment anticancer therapy than in those who did not. In the lenvatinib arm, median OS was 19.5 months (95% CI: 15.7, 23.0) for subjects who received post-treatment anticancer therapy (43%) and 10.5 months (95% CI: 8.6, 12.2) for those who did not. In the sorafenib arm, median OS was 17.0 months (95% CI: 14.2, 18.8) for subjects who received posttreatment anticancer therapy (51%) and 7.9 months (95% CI: 6.6, 9.7) for those who did not. Median OS was longer by approximately 2.5 months in the lenvatinib compared with the sorafenib arm in both subsets of subjects (with or without post-treatment anticancer therapy).

QT interval prolongation

A single 32-mg dose of lenvatinib did not prolong the QT/QTc interval based on results from a thorough QT study in healthy volunteers; however, QT/QTc interval prolongation has been reported at a higher incidence in patients treated with lenvatinib than in patients treated with placebo (see sections 4.4 and 4.8).

Paediatric population

The European Medicines Agency (EMA) has deferred the obligation to submit the results of a study with lenvatinib in one or more subsets of the paediatric population in the treatment of radioiodine-refractory differentiated thyroid cancer and has waived the obligation to submit the results of studies with lenvatinib in one or more subsets of the paediatric population in the treatment of hepatocellular carcinoma (HCC).

5.2 Pharmacokinetic properties

Pharmacokinetic parameters of lenvatinib have been studied in healthy adult subjects, adult subjects with hepatic impairment, renal impairment, and solid tumours.

Absorption

Lenvatinib is rapidly absorbed after oral administration with $t_{\max}$ typically observed from 1 to 4 hours postdose. Food does not affect the extent of absorption, but slows the rate of absorption. When administered with food to healthy subjects, peak plasma concentrations are delayed by 2 hours. Absolute bioavailability has not been determined in humans; however, data from a mass-balance study suggest that it is in the order of 85%. Lenvatinib exhibited good oral bioavailability in dogs (70.4%) and monkeys (78.4%).

Distribution

In vitro binding of lenvatinib to human plasma proteins is high and ranged from 98% to 99% (0.3 - 30 µg/mL, mesilate). This binding was mainly to albumin with minor binding to α1-acid glycoprotein and γ-globulin.

In vitro, the lenvatinib blood-to-plasma concentration ratio ranged from 0.589 to 0.608 (0.1 – 10 µg/mL, mesilate).

Lenvatinib is a substrate for P-gp and BCRP. Lenvatinib is not a substrate for OAT1, OAT3, OATP1B1, OATP1B3, OCT1, OCT2, MATE1, MATE2-K or the bile salt export pump BSEP.

In patients, the median apparent volume of distribution (V_z/F) of the first dose ranged from 50.5 L to 92 L and was generally consistent across the dose groups from 3.2 mg to 32 mg. The analogous median apparent volume of distribution at steady-state (V_z/F_{ss}) was also generally consistent and ranged from 43.2 L to 121 L.

Biotransformation

In vitro, cytochrome P450 3A4 was demonstrated as the predominant (>80%) isoform involved in the P450-mediated metabolism of lenvatinib. However, *in vivo* data indicated that non-P450-mediated pathways contributed to a significant portion of the overall metabolism of lenvatinib. Consequently, *in vivo*, inducers and inhibitors of CYP 3A4 had a minimal effect on lenvatinib exposure (see section 4.5).

In human liver microsomes, the demethylated form of lenvatinib (M2) was identified as the main metabolite. M2' and M3', the major metabolites in human faeces, were formed from M2 and lenvatinib, respectively, by aldehyde oxidase.

In plasma samples collected up to 24 hours after administration, lenvatinib constituted 97% of the radioactivity in plasma radiochromatograms while the M2 metabolite accounted for an additional 2.5%. Based on $AUC_{(0 - \infty)}$, lenvatinib accounted for 60% and 64% of the total radioactivity in plasma and blood, respectively.

Data from a human mass balance/excretion study indicate lenvatinib is extensively metabolised in humans. The main metabolic pathways in humans were identified as oxidation by aldehyde oxidase, demethylation via CYP3A4, glutathione conjugation with elimination of the O-aryl group (chlorophenyl moiety), and combinations of these pathways followed by further biotransformations (e.g., glucuronidation, hydrolysis of the glutathione moiety, degradation of the cysteine moiety, and intramolecular rearrangement of the cysteinylglycine and cysteine conjugates with subsequent dimerisation). These *in vivo* metabolic routes align with the data provided in the *in vitro* studies using human biomaterials.

In vitro transporter studies

For the following transporters, OAT1, OAT3, OATP1B1, OCT1, OCT2, and BSEP, clinically relevant inhibition was excluded based on a cutoff of $IC_{50} > 50 \times C_{\max, \text{unbound}}$.

Lenvatinib showed minimal or no inhibitory activities toward P-gp-mediated and breast cancer resistance protein (BCRP)-mediated transport activities. Similarly, no induction of P-gp mRNA expression was observed.

Lenvatinib showed minimal or no inhibitory effect on OATP1B3 and MATE2-K. Lenvatinib weakly inhibits MATE1. In human liver cytosol, lenvatinib did not inhibit aldehyde oxidase activity.

Elimination

Plasma concentrations decline bi-exponentially following $C_{\max}$. The mean terminal exponential half-life of lenvatinib is approximately 28 hours.

Following administration of radiolabelled lenvatinib to 6 patients with solid tumours, approximately two-thirds and one-quarter of the radiolabel were eliminated in the faeces and urine, respectively. The M3 metabolite was the predominant analyte in excreta (~17% of the dose), followed by M2' (~11% of the dose) and M2 (~4.4 of the dose).

Linearity/non-linearity

Dose proportionality and accumulation

In patients with solid tumours administered single and multiple doses of lenvatinib once daily, exposure to lenvatinib ($C_{\max}$ and AUC) increased in direct proportion to the administered dose over the range of 3.2 to 32 mg once-daily.

Lenvatinib displays minimal accumulation at steady state. Over this range, the median accumulation index (Rac) ranged from 0.96 (20 mg) to 1.54 (6.4 mg). The Rac in HCC subjects with mild and moderate liver impairment was similar to that reported for other solid tumours.

Special populations

Hepatic impairment

The pharmacokinetics of lenvatinib following a single 10-mg dose were evaluated in 6 subjects each with mild and moderate hepatic impairment (Child-Pugh A and Child-Pugh B, respectively). A 5-mg dose was evaluated in 6 subjects with severe hepatic impairment (Child-Pugh C). Eight healthy, demographically matched subjects served as controls and received a 10-mg dose. Lenvatinib exposure, based on dose-adjusted AUC_{0-t} and AUC_{0-inf} data, was 119%, 107%, and 180% of normal for subjects with mild, moderate, and severe hepatic impairment, respectively. It is unknown whether there is a change in the plasma protein binding in hepatically impaired subjects. See section 4.2 for dosing recommendation.

There are not sufficient data for HCC patients with Child-Pugh B (moderate hepatic impairment, 3 patients treated with lenvima in the pivotal trial) and no data available in Child Pugh C HCC patients (severe hepatic impairment). Lenvatinib is mainly eliminated via the liver and exposure might be increased in these patient populations.

The median half-life was comparable in subjects with mild, moderate, and severe hepatic impairment as well as those with normal hepatic function and ranged from 26 hours to 31 hours. The percentage of the dose of lenvatinib excreted in urine was low in all cohorts (<2.16% across treatment cohorts).

Renal impairment

The pharmacokinetics of lenvatinib following a single 24-mg dose were evaluated in 6 subjects each with mild, moderate, and severe renal impairment, and compared with 8 healthy, demographically matched subjects. Subjects with end-stage renal disease were not studied.

Lenvatinib exposure, based on AUC_{0-inf} data, was 101%, 90%, and 122% of normal for subjects with mild, moderate, and severe renal impairment, respectively. It is unknown whether there is a change in the plasma protein binding in renally impaired subjects. See section 4.2 for dosing recommendation.

Age, sex, weight, race

Based on a population pharmacokinetic analysis of patients receiving up to 24 mg lenvatinib once daily, age, sex, weight, and race (Japanese vs. other, Caucasian vs. other) had no significant effects on clearance (see section 4.2).

Paediatric Population

Paediatric patients have not been studied.

5.3 Preclinical safety data

In the repeated-dose toxicity studies (up to 39 weeks), lenvatinib caused toxicologic changes in various organs and tissues related to the expected pharmacologic effects of lenvatinib including glomerulopathy, testicular hypocellularity, ovarian follicular atresia, gastrointestinal changes, bone changes, changes to the adrenals (rats and dogs), and arterial (arterial fibrinoid necrosis, medial degeneration, or haemorrhage) lesions in rats, dogs, and cynomolgus monkeys. Elevated transaminase levels associated with signs of hepatotoxicity, were also observed in rats, dogs and monkeys. Reversibility of the toxicologic changes was observed at the end of a 4-week recovery period in all animal species investigated.

Genotoxicity

Lenvatinib was not genotoxic.

Carcinogenicity studies have not been conducted with lenvatinib.

Reproductive and developmental toxicity

No specific studies with lenvatinib have been conducted in animals to evaluate the effect on fertility. However, testicular (hypocellularity of the seminiferous epithelium) and ovarian changes (follicular atresia) were observed in repeated-dose toxicity studies in animals at exposures 11 to 15 times (rat) or 0.6 to 7 times (monkey) the anticipated clinical exposure (based on AUC) at the maximum tolerated human dose. These findings were reversible at the end of a 4-week recovery period.

Administration of lenvatinib during organogenesis resulted in embryoletality and teratogenicity in rats (foetal external and skeletal anomalies) at exposures below the clinical exposure (based on AUC) at the maximum tolerated human dose, and rabbits (foetal external, visceral or skeletal anomalies) based on body surface area; mg/m² at the maximum tolerated human dose. These findings indicate that lenvatinib has a teratogenic potential, likely related to the pharmacologic activity of lenvatinib as an antiangiogenic agent.

Lenvatinib and its metabolites are excreted in rat milk.

Juvenile animal toxicity studies

Mortality was the dose-limiting toxicity in juvenile rats in which dosing was initiated on postnatal day (PND) 7 or PND21 and was observed at exposures that were respectively 125- or 12-fold lower compared with the exposure at which mortality was observed in adult rats, suggesting an increasing sensitivity to toxicity with decreasing age. Therefore mortality may be attributed to complications related to primary duodenal lesions with possible contribution from additional toxicities in immature target organs.

The toxicity of lenvatinib was more prominent in younger rats (dosing initiated on PND7) compared with those with dosing initiated on PND21 and mortality and some toxicities were observed earlier in the juvenile rats at 10 mg/kg compared with adult rats administered the same dose level. Growth retardation, secondary delay of physical development, and lesions attributable to pharmacologic effects (incisors, femur [epiphyseal growth plate], kidneys, adrenals, and duodenum) were also observed in juvenile rats.

6. PHARMACEUTICAL PARTICULARS

6.1 List of excipients

Capsule contents

Calcium carbonate

Mannitol

Microcrystalline cellulose

Hydroxypropylcellulose

Low-substituted hydroxypropylcellulose

Talc

Capsule shell

Hypromellose

Titanium dioxide (E171)

Yellow iron oxide (E172)

Red iron oxide (E172)

Printing ink

Shellac

Black iron oxide (E172)

Potassium hydroxide

Propylene glycol

6.2 Incompatibilities

Not applicable.

6.3 Shelf life

4 years.

6.4 Special precautions for storage

Do not store above 30°C. Store in the original blister in order to protect from moisture.

6.5 Nature and contents of container

Polyamide/Aluminium/PVC/Aluminium blisters containing 10 capsules. Each carton contains 20 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.

Caregivers should not open the capsule, in order to avoid repeated exposure to the contents of the capsule.

Reg. No. : DKI1747700101A1 (4 mg)
DKI1747700101B1 (10 mg)

DO NOT DISPENSE WITHOUT PHYSICIAN'S PRE-SCRIPTION
(HARUS DENGAN RESEP DOKTER)

Manufactured by :
Patheon Inc. Toronto Region Operations
2100 Syntex Court, Mississauga, Ontario L5N 7K9, Canada
Under license of Eisai Co., Ltd. TOKYO, JAPAN

Packaged by:
Eisai Co., Ltd
GIFU-KEN, JAPAN



Imported by:
PT Eisai Indonesia
BOGOR, INDONESIA

Lembar informasi untuk pasien
LENVIMA 4 mg kapsul
LENVIMA 10 mg kapsul
lenvatinib

Obat ini masih dalam pemantauan lanjutan. Pemantauan tersebut bertujuan untuk mendeteksi dengan cepat jika ada informasi keamanan baru. Anda dapat membantu dengan melaporkan efek samping yang Anda alami. Lihat bagian akhir Bagian 4 mengenai cara pelaporan efek samping.

Baca seluruh isi lembaran ini dengan hati-hati sebelum Anda mulai meminum obat ini karena informasi di dalamnya penting bagi Anda

- Simpan lembaran ini. Anda mungkin perlu membacanya kembali.
- Jika Anda memiliki pertanyaan lebih lanjut, tanyakan pada dokter atau apoteker Anda.
- Obat ini hanya diresepkan untuk Anda. Jangan memberikannya pada orang lain. Efeknya mungkin berbahaya bagi mereka, meskipun jika tanda-tanda penyakit mereka sama dengan Anda.
- Jika Anda mengalami efek samping, bicarakanlah dengan dokter, apoteker, atau perawat Anda. Termasuk jika Anda mengalami efek samping di luar yang tercantum dalam lembaran ini. Lihat Bagian 4.

Apa isi lembaran ini:

1. Apa itu LENVIMA dan untuk kondisi apa LENVIMA digunakan
2. Apa yang perlu Anda ketahui sebelum mulai meminum LENVIMA
3. Bagaimana cara penggunaan LENVIMA
4. Efek samping yang mungkin timbul
5. Bagaimana cara menyimpan LENVIMA
6. Isi kemasan dan informasi lainnya

1. Apa itu LENVIMA dan untuk kondisi apa LENVIMA digunakan

Apa itu LENVIMA

LENVIMA adalah obat yang mengandung zat aktif lenvatinib. LENVIMA digunakan untuk terapi kanker tiroid yang bersifat progresif atau lanjut pada dewasa ketika terapi yodium radioaktif tidak lagi membantu menghentikan penyakit.

LENVIMA juga dapat digunakan untuk mengobati kanker hati (*hepatocellular carcinoma*) pada penderita yang sebelumnya belum pernah menjalani pengobatan dengan anti-kanker lain yang melewati aliran darah. Selain itu, LENVIMA juga dapat digunakan ketika kanker hati telah menyebar dan tidak dapat diatasi dengan tindakan operasi.

Bagaimana LENVIMA bekerja

LENVIMA memblokir kerja protein bernama reseptor tirosin kinase (RTK), yang berperan pada pertumbuhan sel dan pembentukan pembuluh darah baru yang mendarahi sel-sel tersebut. Protein ini dapat ditemukan dalam jumlah besar pada sel-sel kanker, sehingga dengan memblokir kerja protein tadi LENVIMA dapat memperlambat laju pertumbuhan sel-sel kanker dan membantu memutus aliran darah yang diperlukan kanker.

2. Apa yang perlu Anda ketahui sebelum mulai meminum LENVIMA

Jangan menggunakan LENVIMA jika:

- Anda alergi terhadap lenvatinib atau bahan baku lainnya dari obat ini (dicantumkan pada Bagian 6)
- Anda sedang menyusui (lihat bagian mengenai Kontrasepsi, kehamilan dan menyusui).

Peringatan dan perhatian

Konsultasikan dengan dokter Anda sebelum memulai terapi LENVIMA jika Anda:

- memiliki tekanan darah tinggi
- adalah seorang wanita yang memiliki kemungkinan untuk menjadi hamil (lihat bagian mengenai Kontrasepsi, kehamilan, dan menyusui)
- memiliki riwayat penyakit jantung atau stroke
- memiliki gangguan pada hati atau ginjal
- baru-baru ini menjalani tindakan pembedahan atau terapi radiasi
- perlu pembedahan. Dokter Anda mungkin mempertimbangkan menghentikan LENVIMA jika Anda akan menjalani pembedahan besar karena LENVIMA dapat mempengaruhi penyembuhan luka. LENVIMA dapat diberikan kembali bila penyembuhan luka sudah memadai.
- berusia lebih dari 75 tahun
- termasuk kelompok etnis selain Asia atau Kaukasia
- memiliki berat badan kurang dari 60 kg
- memiliki riwayat terbentuknya fistula abnormal antara beberapa organ tubuh atau dari suatu organ terhadap kulit

Sebelum meminum LENVIMA, dokter Anda mungkin akan melakukan serangkaian tes, untuk memeriksa tekanan darah Anda dan fungsi hati atau ginjal Anda dan untuk melihat apakah Anda memiliki kadar garam yang rendah dan kadar *thyroid stimulating hormone (TSH)* yang tinggi di dalam darah Anda. Dokter Anda akan mendiskusikan hasil tes darah tersebut dengan Anda dan akan menentukan apakah Anda dapat diberikan LENVIMA. Anda mungkin membutuhkan pengobatan tambahan dengan obat lain, diberikan dosis LENVIMA yang lebih rendah, atau diperlukan kehati-hatian lebih karena peningkatan efek samping.

Jika Anda tidak merasa yakin, bicarakan dengan dokter Anda sebelum meminum LENVIMA

Anak-anak dan remaja

LENVIMA tidak dianjurkan untuk digunakan pada anak dan remaja. Efek LENVIMA pada orang berusia kurang dari 18 tahun belum diketahui.

Obat-obatan lain dan LENVIMA

Beritahukan dokter atau apoteker Anda jika Anda sedang atau baru-baru ini atau kemungkinan akan meminum obat lain. Termasuk obat-obat tradisional (jamu) dan obat-obatan yang diminum tanpa resep dokter.

Kontrasepsi, kehamilan, dan menyusui

Jika Anda hamil atau menyusui, Anda berpikir bahwa Anda hamil atau berencana untuk hamil, tanyakan dokter atau apoteker Anda sebelum meminum obat ini.

- Jika Anda memiliki kemungkinan untuk hamil, gunakan kontrasepsi yang sangat efektif selama meminum obat ini, dan paling tidak sebulan setelah terapi LENVIMA Anda selesai. Karena tidak diketahui apakah LENVIMA dapat menurunkan efek pil kontrasepsi oral, jika Anda menggunakan metode kontrasepsi tersebut Anda harus menambahkan metode kontrasepsi *barrier* seperti *cap* atau kondom saat berhubungan selama Anda menjalani terapi dengan LENVIMA.
- Jangan meminum LENVIMA apabila Anda berencana untuk hamil selama masih dalam terapi LENVIMA. Larangan ini timbul karena LENVIMA dapat membahayakan janin Anda.
- Seandainya Anda hamil saat menjalani terapi LENVIMA, segera beritahukan dokter Anda. Dokter akan membantu Anda untuk memutuskan apakah terapi sebaiknya dilanjutkan.
- Jangan menyusui selama Anda menggunakan LENVIMA karena obat ini dapat masuk ke dalam air susu dan dapat sangat membahayakan bayi Anda.

Mengemudi dan mengoperasikan mesin

LENVIMA dapat menyebabkan efek samping yang memengaruhi kemampuan Anda mengemudi atau mengoperasikan mesin. Jangan mengemudi atau mengoperasikan mesin jika Anda merasa pusing atau lelah.

3. Bagaimana cara penggunaan LENVIMA

Selalu gunakan LENVIMA sesuai yang diinstruksikan dokter Anda. Tanyakan pada dokter atau apoteker Anda jika Anda tidak yakin.

Berapa banyak yang harus diminum

Kanker Tiroid

- Dosis LENVIMA yang dianjurkan biasanya adalah 24 mg sekali sehari (2 kapsul 10 mg dan 1 kapsul 4 mg)
- Jika Anda memiliki gangguan ginjal atau hati yang berat, dosis yang dianjurkan adalah 14 mg sekali sehari (1 kapsul 10 mg dan 1 kapsul 4 mg)
- Dokter Anda mungkin mengurangi dosis jika Anda mengalami masalah terkait efek samping

Kanker Hati

- Dosis LENVIMA yang dianjurkan tergantung dari berat badan Anda ketika pertama kali memulai pengobatan. Dosis umumnya sebesar 12 mg sekali sehari (3 kapsul sediaan 4 mg) jika berat badan Anda 60 kg atau lebih; dan 8 mg sekali sehari (2 kapsul sediaan 4 mg) jika berat badan Anda kurang dari 60 kg.
- Dokter dapat mengurangi dosis yang diberikan apabila Anda mengalami gangguan efek samping.

Meminum obat ini

- Anda dapat meminum kapsul ini setelah atau sebelum makan
- Telanlah kapsul dalam bentuk utuh dengan air putih atau dengan melarutkannya. Untuk melarutkannya, tuangkan satu sendok air atau jus apel ke dalam gelas kecil dan letakan kapsul ke dalam air tanpa menghancurkannya. Biarkan selama kurang lebih 10 menit, lalu aduk selama kurang lebih 3 menit untuk melarutkan cangkang kapsul. Minum larutan tersebut. Setelah diminum, tambahkan sejumlah air atau jus apel pada wadah yang sama, kemudian aduk dan diminum.
- Minumlah kapsul ini pada waktu yang kurang lebih sama setiap hari.

- *Caregiver* harus menghindari paparan isi kapsul.

Berapa lama Anda harus minum LENVIMA

Anda biasanya akan terus minum obat ini selama Anda masih mendapatkan manfaat.

Jika Anda minum LENVIMA melebihi dosis yang seharusnya

Jika Anda minum LENVIMA melebihi dosis yang seharusnya, langsung bicarakan dengan dokter atau apoteker. Bawalah obat beserta kemasannya bersama Anda.

Jika Anda lupa minum LENVIMA

Jangan minum dosis ganda (dua dosis sekaligus pada saat yang bersamaan) untuk menggantikan dosis yang lupa Anda minum.

Apa yang harus Anda lakukan jika lupa minum LENVIMA tergantung pada berapa lama waktu yang tersisa sebelum dosis berikutnya

- Jika jadwal dosis berikutnya adalah 12 jam atau lebih: minum dosis yang Anda lupakan tadi segera setelah Anda ingat. Kemudian minum dosis selanjutnya pada waktu seperti biasa.
- Jika jadwal dosis berikutnya adalah kurang dari 12 jam: abaikan dosis yang terlupa. Kemudian minum dosis selanjutnya pada waktu seperti biasa.

4. Efek samping yang mungkin timbul

Seperti halnya semua obat, LENVIMA dapat menyebabkan efek samping, meskipun tidak semua orang akan mengalaminya. Efek samping berikut mungkin timbul pada penggunaan obat ini.

Langsung beritahukan dokter Anda jika Anda mengalami salah satu/lebih dari efek samping berikut – Anda mungkin memerlukan terapi medis segera:

- merasa kebas atau lemah pada salah satu sisi tubuh Anda, sakit kepala hebat, kejang, kebingungan, kesulitan berbicara, perubahan penglihatan atau merasa pusing – gejala-gejala tersebut mungkin merupakan tanda stroke, perdarahan dalam otak Anda, atau efek pada otak Anda akibat meningkatnya tekanan darah secara drastis.
- nyeri atau rasa tertekan pada dada, nyeri pada lengan, punggung, leher atau rahang, sesak napas, detak jantung yang cepat atau tidak teratur, batuk, warna kebiruan pada bibir atau jari, merasa sangat lelah – gejala-gejala tadi mungkin merupakan tanda gangguan pada jantung atau adanya gumpalan darah di paru-paru Anda.
- nyeri hebat dalam perut (abdomen) Anda – hal ini dapat terjadi karena timbulnya lubang pada dinding usus Anda atau suatu fistula (lubang dalam usus yang menyambungkan saluran seperti selang dengan bagian lain tubuh atau kulit)
- feses berwarna hitam, seperti tar, atau berdarah, atau batuk darah – hal-hal tadi mungkin merupakan tanda perdarahan dalam tubuh Anda.
- Kulit nampak kekuningan atau warna putih pada mata nampak kekuningan (*jaundice*) atau timbul gangguan rasa kantuk, kebingungan, konsentrasi menurun – hal tersebut mungkin merupakan tanda timbulnya gangguan pada hati.
- diare, merasa mual dan muntah-muntah – hal-hal tadi merupakan efek samping yang sangat lazim yang dapat berakibat serius jika sampai timbul dehidrasi, yang dapat menyebabkan gagal ginjal. Dokter Anda dapat memberikan obat-obatan untuk mengurangi efek samping tersebut.

Segera beritahukan dokter Anda jika Anda mengalami salah satu atau lebih dari efek samping tersebut.

Efek samping lain meliputi:

Sangat lazim (dapat mengenai lebih dari 1 di antara 10 orang)

- tekanan darah tinggi atau rendah
- hilangnya nafsu makan atau penurunan berat badan
- merasa mual dan muntah-muntah, konstipasi, diare, nyeri perut, gangguan cerna
- merasa sangat lelah atau lemas
- suara serak
- pembengkakan pada tungkai
- ruam
- mulut kering, nyeri, atau meradang, rasa pengecapan yang aneh
- pembengkakan dan peradangan pada sendi; dan sendi, tulang, dan otot yang kaku
- merasa pusing
- rambut rontok
- perdarahan (paling lazim mimisan, tetapi dapat juga terjadi perdarahan di tempat lain seperti darah dalam urin, memar, perdarahan gusi atau dinding usus)
- kesulitan tidur
- hasil pemeriksaan urin yang abnormal untuk protein (terlalu tinggi) dan infeksi saluran kemih (meningkatnya kekerapan berkemih dan nyeri saat berkemih)
- nyeri – otot, sendi, nyeri kepala, punggung
- kemerahan, nyeri, dan pembengkakan pada kulit di tangan dan kaki (*hand-foot syndrome*)
- penurunan fungsi kelenjar tiroid (lemas, kenaikan berat badan, sembelit, merasa kedinginan, kulit kering)
- hasil pemeriksaan darah yang abnormal untuk kadar kalium (terlalu rendah) dan kalsium (terlalu rendah)
- menurunnya jumlah sel darah putih
- perubahan pada hasil tes darah yang menunjukkan gangguan fungsi hati
- memar dan luka yang tidak kunjung sembuh – tanda rendahnya kadar trombosit dalam darah

Lazim (dapat mengenai hingga 1 di antara 10 orang)

- hilangnya cairan tubuh (dehidrasi)
- jantung berdebar
- kulit kering, menebal dan gatal
- rasa kembung (*flatulence*)
- gangguan jantung atau gumpalan darah di paru (kesulitan bernapas, nyeri dada) atau organ lainnya
- gangguan fungsi hati
- rasa kantuk, kebingungan, konsentrasi menurun, pingsan; yang mungkin dapat menjadi tanda terjadinya gangguan fungsi hati.
- merasa tidak enak badan
- radang kandung empedu
- stroke
- fistula ani (saluran kecil yang terbentuk antara anus dan kulit sekitarnya)

- hasil pemeriksaan darah yang abnormal untuk magnesium yang terlalu rendah, kolesterol yang terlalu tinggi, dan TSH (*thyroid stimulating hormone*) yang terlalu tinggi
- perubahan hasil pemeriksaan darah untuk fungsi ginjal dan gagal ginjal
- perubahan lipase dan amylase (enzim yang berperan pada proses pencernaan)

Tidak lazim (dapat mengenai hingga 1 di antara 100 orang)

- infeksi yang nyeri atau iritasi di sekitar anus
- mini-stroke
- kerusakan hati
- nyeri hebat pada bagian atas kiri atas perut (abdomen) yang dapat disertai demam, menggigil, mual dan muntah (kemungkinan terjadinya gangguan fungsi limpa)
- peradangan pancreas
- masalah penyembuhan luka
- nyeri di bagian punggung, dada atau perut yang terkait dengan robek di dinding aorta dan pendarahan interna

Tidak diketahui (efek samping berikut telah dilaporkan sejak pemasaran LENVIMA namun frekuensi kejadian nya tidak diketahui)

- Tipe lain fistula (suatu kelainan hubungan antar organ-organ dalam tubuh atau antara kulit dan struktur yang mendasari seperti tenggorokan dan batang tenggorokan). Gejala-gejala yang muncul tergantung dimana lokasi fistula berada. Bicarakan dengan dokter Anda apabila mengalami keluhan baru atau tidak biasa seperti batuk ketika menelan.

Melaporkan efek samping

Jika Anda mengalami efek samping, bicarakan dengan dokter atau apoteker Anda. Termasuk jika Anda mengalami efek samping yang mungkin tidak tercantum dalam lembar ini.

Dengan melaporkan efek samping, Anda dapat membantu memberikan tambahan informasi mengenai keamanan obat ini.

5. Bagaimana cara menyimpan LENVIMA

- Jauhkan obat ini dari penglihatan dan jangkauan anak-anak.
- Jangan gunakan obat ini melewati tanggal kedaluwarsa yang tercantum pada karton dan tiap-tiap blister. Tanggal kedaluwarsa berarti hari terakhir pada bulan itu.
- Jangan disimpan pada suhu melebihi 30°C. Simpan dalam blister aslinya untuk melindungi kapsul dari kelembaban.
- Jangan buang obat ini melalui pembuangan air kotor atau pembuangan sampah rumah tangga biasa. Tanyakan pada apoteker bagaimana cara membuang obat yang tidak lagi Anda gunakan. Tindakan-tindakan ini akan membantu melindungi lingkungan.

6. Isi kemasan dan informasi lainnya

Apa komposisi LENVIMA

- Zat aktifnya adalah lenvatinib.
 - LENVIMA 4 mg kapsul : - Tiap kapsul mengandung 4 mg lenvatinib (dalam bentuk mesilat).
 - LENVIMA 10 mg kapsul : - Tiap kapsul mengandung 10 mg lenvatinib (dalam bentuk mesilat).
- Komposisi lainnya adalah *calcium carbonate, mannitol, microcrystalline cellulose, hydroxypropylcellulose, low-substituted hydroxypropyl cellulose, talc*. Cangkang kapsul mengandung *hypromellose, titanium dioxide (E171), yellow iron oxide (E172), red iron oxide (E172)*. Tinta cetak mengandung *shellac, black iron oxide (E172), potassium hydroxide, propylene glycol*.

Seperti apa tampilan LENVIMA dan isi kemasan

- Kapsul 4 mg memiliki badan merah kekuningan dan tutup merah kekuningan, dengan panjang kurang lebih 14,3 mm, dengan tulisan “C” berwarna hitam pada tutup dan “LENV 4 mg” berwarna hitam pada badan.
- Kapsul 10 mg memiliki badan berwarna kuning dan tutup merah kekuningan, dengan panjang kurang lebih 14,3 mm, dengan tulisan “C” berwarna hitam pada tutup dan “LENV 10 mg” berwarna hitam pada badan.
- Kapsul dikemas dalam blister *polyamide/aluminium/PVC* dengan lapisan penutup *aluminium foil* yang tembus bila ditekan, dalam dus karton berisi 20 kapsul.

Dibuat oleh: **Patheon Inc., Toronto, Kanada**

Di bawah lisensi **Eisai Co., Ltd., Tokyo, Jepang**

Dikemas Oleh: **Eisai Co., Ltd., Kakamigahara, Jepang**

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