

PRODUCT INFORMATION

Lenvima[®] lenvatinib (as lenvatinib mesilate) hard capsules

NAME OF THE MEDICINE

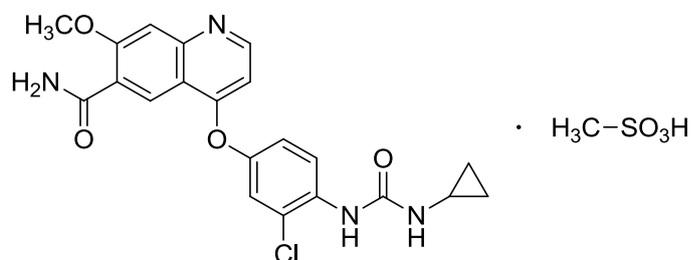
Lenvatinib as lenvatinib mesilate

Chemical Structure

Lenvatinib is a multiple receptor tyrosine kinase (RTK) inhibitor.

Chemical Name: 4-[3-chloro-4-(*N*'-cyclopropylureido)phenoxy]-7-methoxyquinoline-6-carboxamide methanesulfonate.

The empirical formula of lenvatinib is $C_{21}H_{19}ClN_4O_4 \cdot CH_4O_3S$



CAS Number: 857890-39-2

DESCRIPTION

Lenvatinib mesilate is a white powder and is sparingly soluble in acetic acid and slightly soluble in water, *N,N*-dimethylformamide, methanol, *N*-methylpyrrolidone, and pyridine. It is very slightly soluble in 1,3-dimethyl-2-imidazolidinone and practically insoluble in acetonitrile, dehydrated ethanol, 1-propanol, 2-propanol, 1-octanol and isopropyl acetate. In aqueous solutions, lenvatinib mesilate is very slightly soluble in 0.1 mol/L HCl and practically insoluble in Britton-Robinson buffer, pH 3-11.

Each 4 mg hard capsule contains lenvatinib mesilate equivalent to 4 mg lenvatinib.

Each 10 mg hard capsule contains lenvatinib mesilate equivalent to 10 mg lenvatinib.

The capsules contain the excipients Calcium carbonate, Mannitol, Microcrystalline cellulose, Hydroxypropylcellulose, and Purified talc. The capsule shell contains the excipients Hypromellose, Titanium dioxide, Iron oxide yellow and Iron oxide red. The printing ink on the capsules contains the excipients Shellac, Iron oxide black, Potassium hydroxide and Propylene glycol.

PHARMACOLOGY

Pharmacodynamic Properties

Mechanism of Action

Lenvatinib is a multiple receptor tyrosine kinase (RTK) inhibitor that 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. The combination of lenvatinib and everolimus showed increased antiangiogenic and antitumour activity as demonstrated by decreased human endothelial cell proliferation, tube formation, and VEGF signalling *in vitro* and tumour volume in mouse xenograft models of human renal cell cancer greater than each drug alone.

Pharmacodynamic effects

Cardiac electrophysiology

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 LENVIMA than in patients treated with placebo (see ADVERSE EFFECTS, Selected Adverse Reactions).

Pharmacokinetics

Absorption

Lenvatinib is rapidly absorbed after oral administration with T_{max} typically observed from 1 to 4 hours post-dose. 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.

A high degree of inter-individual variability in average exposure at steady state was observed, with a 6-fold range when used as monotherapy at the 24 mg dose, and 7-fold range when LENVIMA 18 mg dose is administered in combination with 5mg everolimus.

Distribution

In vitro binding of lenvatinib to human plasma proteins was high and ranged from 98% to 99% (0.3 – 30 μ g/mL, mesilate). This binding was mainly to albumin with minor binding to α 1-acid glycoprotein and γ -globulin.

In vitro, the lenvatinib blood-to-plasma concentration ratio ranged from 0.589 to 0.608 (0.1 - 10 μ g/mL, mesilate). *In vitro* studies indicate that lenvatinib is a substrate for P-gp and BCRP. Lenvatinib is not a substrate for OAT1, OAT3, OATP1B1, OATP1B3, OCT1, OCT2, or the BSEP.

Metabolism

In vitro, cytochrome P450 3A4 was the predominant (>80%) cytochrome isoform involved in the P450-mediated metabolism of lenvatinib. *In vivo*, inducers and inhibitors of CYP 3A4 had a minimal effect on lenvatinib exposure (see INTERACTIONS WITH OTHER MEDICINES). Patients should avoid strong inducers of CYP 3A4 and exercise caution with

mild or moderate inhibitors or inducers when using everolimus (see Everolimus Product Information) in combination with LENVIMA.

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–inf), 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 (chlorbenzyl moiety), and combinations of these pathways followed by further biotransformations (eg, 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.

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-fourth of the radiolabel were eliminated in the faeces and urine, respectively. The M2 metabolite was the predominant analyte in excreta (~5% of the dose) with lenvatinib the second most prominent (~2.5%).

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 (QD).

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).

Special populations

Hepatic impairment

The pharmacokinetics of lenvatinib following a single 10-mg dose were evaluated in 6 subjects each with mild or 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. 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).

Lenvatinib exposure, based on dose-adjusted $AUC_{0-t,unbound}$ and $AUC_{0-inf,unbound}$, was approximately 65%, 122%, and 273% of normal for subjects with mild, moderate, and severe hepatic impairment, respectively. Based on the analogous AUC_{0-t} and AUC_{0-inf} data, lenvatinib exposure was 119%, 107%, and 180% of normal for subjects with mild, moderate, and severe hepatic impairment, respectively (See DOSAGE AND ADMINISTRATION).

Renal impairment

The pharmacokinetics of lenvatinib following a single 24 mg dose were evaluated in 6 subjects each with mild, moderate, or severe renal impairment, and compared with 8 healthy, demographically matched subjects. Subjects with end-stage renal disease were not studied. The percentage of unbound lenvatinib was similar between subjects with normal renal function ($8\% \pm 3\%$, mean \pm SD) and those with severely impaired renal function ($9\% \pm 2\%$). $AUC_{0-inf,unbound}$ estimates for subjects with mild, moderate, or severe renal impairment were 54%, 129%, and 184%, respectively, compared with normal subjects. Additionally, a linear equation was fit to the creatinine clearance vs. $AUC_{0-inf,unbound}$ data and exposure was predicted. Subjects with severe renal impairment were predicted to have a 2.4-fold increase in exposure. Therefore dosage needs to be reduced in patients with severe renal impairment (See DOSAGE AND ADMINISTRATION).

Age, sex, weight, race

Based on a population pharmacokinetic analysis of patients receiving up to 24 mg LENVIMA once daily, weight showed a statistically significant effect but only explained 2.8% of the inter-individual variability, on apparent clearance. Subjects weighing < 60 kg had 36% higher exposure to lenvatinib than subjects weighing ≥ 60 kg. Simulations showed that the small effect of body weight on lenvatinib exposure does not warrant any dose adjustment. After accounting for body weight, neither age, sex, nor race (Japanese vs. other, Caucasian vs. other) influenced lenvatinib PK.

Paediatric Population

Paediatric patients have not been studied.

Genomic assessment of lenvatinib pharmacokinetic parameters

Because of lenvatinib's extensive metabolism, the effect of selected drug-metabolising enzyme phenotypes on lenvatinib clearance was investigated using data derived from the Affymetrix drug-metabolising enzyme and transporter (DMET Plus) microarray genotyping platform. None of the phenotypes for CYP3A5, CYP1A2, CYP2A6, or CYP2C19 had a significant impact on lenvatinib clearance.

CLINICAL TRIALS

Radioactive iodine refractory differentiated thyroid cancer

The SELECT study was a multicentre, randomised, double-blind, placebo-controlled trial that was conducted in 392 patients with radioactive iodine refractory differentiated thyroid cancer with independent, centrally reviewed, radiographic evidence of disease progression within 12 months (+1 month window) prior to enrolment. Radioiodine-refractory status 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 (OS). Patients in the placebo arm could opt to receive LENVIMA treatment at the time of confirmed disease progression.

Eligible patients with measurable disease according to RECIST 1.1 were randomised 2:1 to receive LENVIMA 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 (54%) had an Eastern Cooperative Oncology Group (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 LENVIMA-treated patients compared with those receiving placebo ($p < 0.0001$). The positive effect on PFS was similar in the subgroups that received 0 or 1 prior VEGF/VEGFR-targeted therapy (see Table 1). In addition, the positive effect on PFS was seen across the subgroups of age, sex, race, histological subtype, and geographic region. Following independent review confirmation of disease progression, 109 (83.2%) patients randomised to placebo crossed over to receive open-label LENVIMA.

There was no statistically significant difference in overall survival in the treatment arm compared to the placebo group at the primary analysis (HR (95% CI): 0.73 (0.59, 1.07)). However, the SELECT study was not powered to demonstrate an improvement in OS, and the high rate of crossover of patients in the placebo arm to the treatment arm after confirmed disease progression made demonstration of a statistically significant difference in OS difficult.

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 LENVIMA, 70.4% were observed to develop the response on or within 30 days of being on the 24-mg dose.

The study did not measure quality of life (QoL). The effect of treatment on QoL can therefore not be assessed and QoL may not be improved with LENVIMA treatment.

Table 1 Efficacy Results in radioactive iodine refractory differentiated thyroid cancer

	LENVIMA 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-target 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) ^{bc}	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) ^{bc}	0.22 (0.12, 0.41)	
Overall 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 (20.3, 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 year vs >65 years), and previous VEGF/VEGFR-targeted therapy (0 vs. 1).

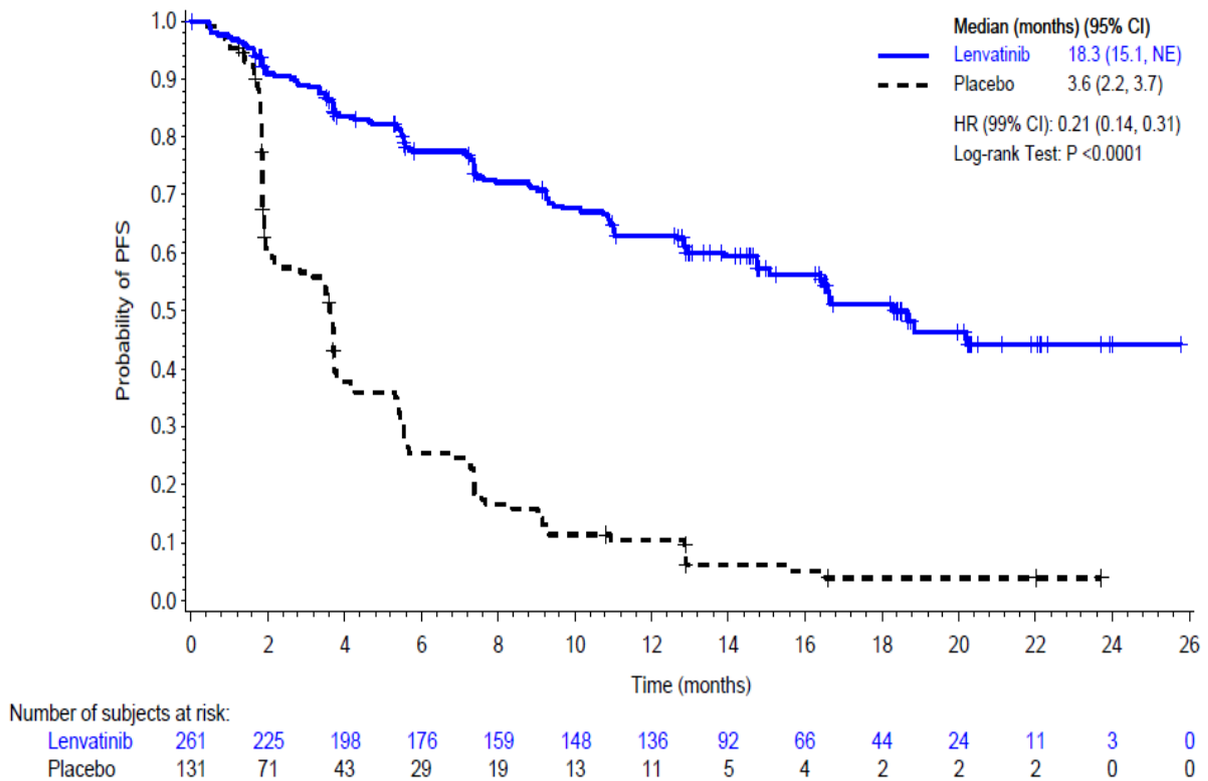
c: Estimated with Cox proportional hazard model.

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 Plot of Progression-Free Survival - DTC



Renal Cell Carcinoma

A multicentre, randomised, open-label, trial was conducted to determine the safety and efficacy of LENVIMA administered alone or in combination with everolimus in subjects with unresectable advanced or metastatic Renal Cell Carcinoma (RCC). The study consisted of a Phase 1b dose finding and a Phase 2 portion. The Phase 1b portion included 11 patients who received the combination of 18 mg of LENVIMA plus 5 mg of everolimus. The Phase 2 portion enrolled a total of 153 patients with unresectable advanced or metastatic RCC, who had previously received 1 prior VEGF-targeted treatment, 1:1:1 to LENVIMA 18 mg plus everolimus 5 mg, LENVIMA 24 mg monotherapy, or everolimus 10 mg monotherapy. All medications were administered orally once daily. Patients were required to have histological confirmation of predominant clear cell RCC, and ECOG Performance Status of 0 or 1. Patients were stratified by haemoglobin level (≤ 13 g/dL vs. > 13 g/dL for males and ≤ 11.5 g/dL vs. > 11.5 g/dL for females) and corrected serum calcium (≥ 10 mg/dL vs. < 10 mg/dL).

Of the 101 patients randomly allocated to the LENVIMA plus everolimus arm and everolimus monotherapy, 72% were male, the median age was 60 years, 31% were 65 years or older, and 96% were Caucasian. All patients were classified as having Stage IV RCC. All patients had a baseline ECOG PS of either 0 (54%) or 1 (46%) with similar distribution across the 2 treatment arms. Memorial Sloan Kettering Cancer Center (MSKCC) favourable, intermediate, and poor risk categories were observed respectively, in 24%, 37%, and 39% of patients in the LENVIMA plus everolimus arm, and 24%, 38%, and 38% of patients in the everolimus arm.

The primary efficacy outcome measure was investigator assessed PFS evaluated according to RECIST 1.1. Efficacy results are summarized in Table 2 and Figures 2 and 3. The treatment

effect of the combination on PFS was supported by a retrospective independent blinded review of radiographs with an observed hazard ratio (HR) of 0.43 (95% CI: 0.24, 0.75) compared with the everolimus arm.

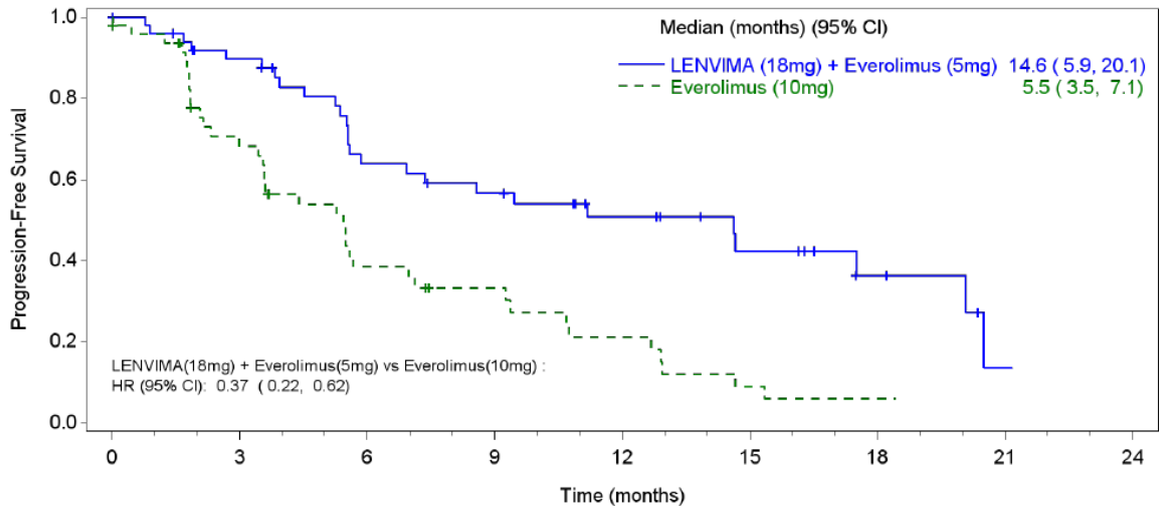
Table 2 Efficacy results in renal cell carcinoma (investigator assessment)

	LENVIMA 18 mg + Everolimus 5 mg (N=51)	Everolimus 10 mg (N=50)
Progression-Free Survival (PFS)^{ab}		
Number of events, n (%)	26 (51)	37 (74)
Progressive disease	21 (41)	35 (70)
Death	5 (10)	2 (4)
Median PFS in months (95% CI)	14.6 (5.9, 20.1)	5.5 (3.5, 7.1)
Hazard Ratio (95% CI) ^c LENVIMA + Everolimus vs Everolimus	0.37 (0.22, 0.62)	-
Overall Survival^d		
Number of deaths, n (%)	32 (63)	37 (74)
Median OS in months (95% CI)	25.5 (16.4, 32.1)	15.4 (11.8, 20.6)
Hazard Ratio (95% CI) ^c LENVIMA + Everolimus vs Everolimus	0.59 (0.36, 0.97)	-
Objective Response Rate (Confirmed)^b		
Objective response rate, n (%)	19 (37)	3 (6)
(95% CI)	(24, 52)	(1, 17)
Number of complete responses, n (%)	1 (2)	0
Number of partial responses (%)	18 (35)	3 (6)

Tumour assessments were based on RECIST v1.1 criteria for progression but only confirmed responses are included for ORR.
CI = confidence interval

- a Point estimates are based on Kaplan-Meier method and 95% CIs are based on the Greenwood formula using log-log transformation.
- b Data cutoff date = 13 Jun 2014
- c Hazard ratio is based on a stratified Cox regression model including treatment as a covariate factor and haemoglobin and corrected serum calcium as strata.
- d Data cutoff date = 31 Jul 2015

Figure 2 Kaplan-Meier Plot of Progression-Free Survival (Investigator Assessment - RCC)

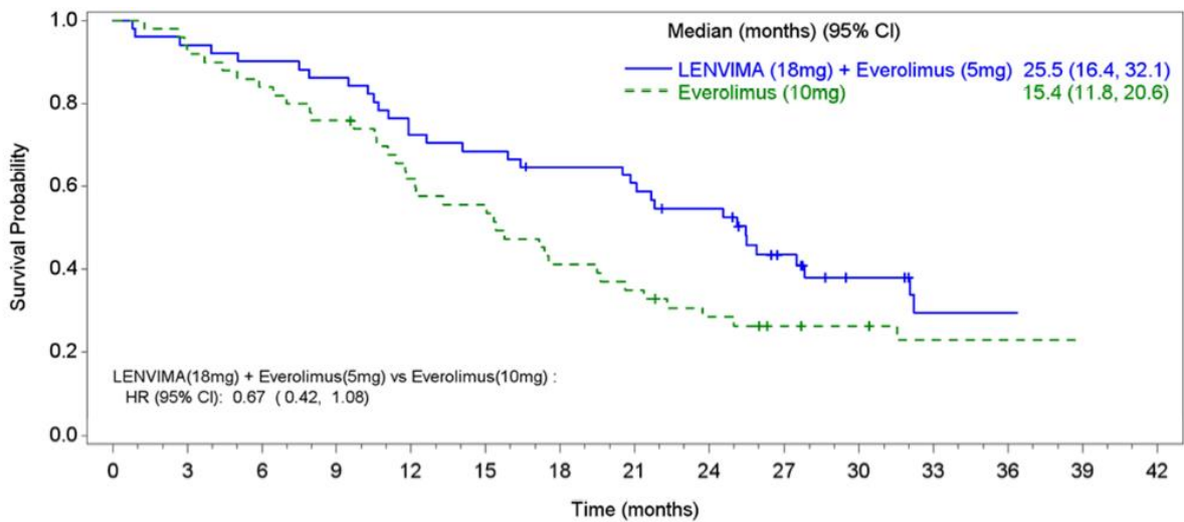


Number of Subjects at risk:

L(18mg) + E(5mg)	51	41	27	23	16	10	5	1	0
E(10mg)	50	29	15	11	7	3	1	0	0

L(18mg) + E(5mg)=LENVIMA 18mg + Everolimus 5mg; E(10mg)=Everolimus 10mg
Data Cutoff Date: 13JUN2014

Figure 3 Kaplan-Meier Plot of Overall Survival - RCC



Number of Subjects at risk:

L(18mg) + E(5mg)	51	48	46	44	37	35	32	30	26	17	11	7	2	0	0
E(10mg)	50	46	42	38	30	27	20	17	13	10	9	5	1	0	0

L(18mg) + E(5mg)=LENVIMA 18mg + Everolimus 5mg; E(10mg)=Everolimus 10mg
Data Cutoff Date: 31JUL2015

INDICATIONS

LENVIMA is indicated for the treatment of patients with progressive, locally advanced or metastatic, radioactive iodine refractory differentiated thyroid cancer.

LENVIMA is indicated in combination with everolimus for the treatment of adult patients with advanced renal cell carcinoma whose disease has progressed following one prior vascular endothelial growth factor targeted therapy.

CONTRAINDICATIONS

Hypersensitivity to the active substance or to any of the excipients (see DESCRIPTION)

PRECAUTIONS

Diarrhoea and dehydration

Diarrhoea has been reported frequently in patients treated with LENVIMA usually occurring early in the course of treatment (See ADVERSE EFFECTS, Selected Adverse Reactions). Prompt medical management of diarrhoea should be instituted in order to prevent dehydration. LENVIMA should be discontinued in the event of persistence of Grade 4 diarrhoea despite medical management (see DOSAGE AND ADMINISTRATION).

Renal failure and impairment

Patients with baseline renal function <60ml/minute experienced more adverse events, including fatal and serious adverse events and Grade 3 or 4 events, than those with normal renal function and were more likely to require a treatment interruption, dose reduction or discontinuation of treatment. The recommended starting dose is lower for patients with renal impairment (see DOSAGE AND ADMINISTRATION) and it is also recommended these patients be monitored closely during treatment. There is no clinical trial experience of patients with severe renal impairment.

Renal impairment (including renal failure) has been reported in patients treated with LENVIMA (see ADVERSE EFFECTS, Selected Adverse Reactions). The primary risk factors identified were pre-existing renal impairment and 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. Serious adverse events of both hypokalaemia and hyperkalaemia have occurred and renal function and electrolytes should be monitored closely. Caution should be taken in patients receiving agents acting on the renin-angiotensin aldosterone system given a potentially higher risk for acute renal failure with the combination treatment. Dose interruptions, adjustments, or discontinuation may be necessary (see DOSAGE AND ADMINISTRATION).

If patients have severe renal impairment, the initial dose of LENVIMA should be adjusted (see DOSAGE AND ADMINISTRATION).

Hypertension

Hypertension has been reported in patients treated with LENVIMA, usually occurring early in the course of treatment (see ADVERSE EFFECTS, Selected Adverse Reactions). Blood pressure (BP) should be well controlled prior to treatment with LENVIMA and, if patients are known to be hypertensive they should be on a stable dose of an antihypertensive therapy for at least 1 week prior to treatment with LENVIMA. The early detection and effective management of hypertension are important to minimise the need for LENVIMA dose interruptions and reductions. Antihypertensives should be started as soon as elevated BP is confirmed. Blood pressure should be monitored after 1 week of treatment with LENVIMA, 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 3.

Table 3 Recommended Management of Hypertension

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

Proteinuria

Proteinuria has been reported in patients treated with LENVIMA, usually occurring early in the course of the treatment (see ADVERSE EFFECTS, Selected Adverse Reactions). Monitor urine protein regularly. If urine dipstick proteinuria \geq 2+ is detected, dose interruptions, adjustments, or discontinuation may be necessary (see DOSAGE AND

ADMINISTRATION). LENVIMA should be discontinued in the event of nephrotic syndrome.

Cardiac dysfunction

Cardiac failure and decreased left ventricular ejection fraction have been reported in patients treated with LENVIMA (see ADVERSE EFFECTS, 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 DOSAGE AND ADMINISTRATION).

LENVIMA has not been studied in patients who have had cardiac failure within the previous 6 months and therefore should be used with caution in such patients.

Posterior reversible encephalopathy syndrome (PRES) / Reversible Posterior Leucoencephalopathy Syndrome (RPLS)

Posterior reversible encephalopathy syndrome (PRES, also known as RPLS) has been reported in patients treated with LENVIMA (observed in < 1% of patients; ADVERSE EFFECTS, 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 PRECAUTIONS, Hypertension). In patients with signs or symptoms of PRES, dose interruptions, adjustments, or discontinuation may be necessary (see DOSAGE AND ADMINISTRATION).

Hepatotoxicity

Liver-related adverse reactions most commonly reported in patients treated with LENVIMA included increases in alanine aminotransferase, increases in aspartate aminotransferase, and increases in blood bilirubin (see ADVERSE EFFECTS, Selected Adverse Reactions). Hepatic failure and acute hepatitis (observed in < 1% of patients) have been reported in patients treated with LENVIMA. The hepatic failure events 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 (see DOSAGE AND ADMINISTRATION).

If patients have any degree of liver impairment they need to be monitored closely for liver related adverse reactions. If patients have severe hepatic impairment, the initial dose of LENVIMA should be adjusted (see DOSAGE and ADMINISTRATION).

Arterial thromboembolic events

Arterial thromboembolic events (cerebrovascular accident, transient ischaemic attack, and myocardial infarction) have been reported in patients treated with LENVIMA (see ADVERSE EFFECTS, Selected Adverse Reactions). LENVIMA has not been studied in patients who have had an arterial thromboembolic event 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 LENVIMA should be discontinued following an arterial thrombotic event (see DOSAGE AND ADMINISTRATION).

Haemorrhagic events and thrombocytopenia

Serious haemorrhagic events have been reported in patients treated with LENVIMA. The most frequently reported haemorrhagic event was mild epistaxis. Serious events of thrombocytopenia have also been reported in patients treated with LENVIMA and thrombocytopenia may increase risk of developing haemorrhagic events. (see ADVERSE EFFECTS, Selected Adverse Reactions)

Serious tumour related bleeds have been reported, including fatal haemorrhagic events in LENVIMA treated patients and there have been reports of haemorrhage associated with thrombocytopenia.

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 LENVIMA therapy. In the case of haemorrhagic events/thrombocytopenia, dose interruptions, adjustments, or discontinuation may be necessary (see DOSAGE AND ADMINISTRATION).

Gastrointestinal perforation and fistula formation

Gastrointestinal perforation or fistulae have been reported in patients treated with LENVIMA (see ADVERSE EFFECTS, Selected Adverse Reactions). 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 DOSAGE AND ADMINISTRATION).

Non-Gastrointestinal fistula

Patients may be at increased risk for the development of fistulae when treated with LENVIMA. Cases of fistula formation or enlargement that involve other areas of the body 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. LENVIMA should not be started in patients with fistulae to avoid worsening and LENVIMA should be permanently discontinued in patients with oesophageal or tracheobronchial tract involvement and any Grade 4 fistula (see DOSAGE AND ADMINISTRATION); 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. LENVIMA may adversely affect the wound healing process as do other agents of the same class.

QT interval prolongation

QT/QTc interval prolongation has been reported at a higher incidence in patients treated with LENVIMA than in patients treated with placebo (see ADVERSE EFFECTS, Selected Adverse Reactions). The median time to onset of QTc prolongation was 16.1 weeks in the

DTC study for patients on LENVIMA monotherapy and 30 weeks in the RCC study for combination patients. Electrocardiograms should be monitored in patients on an ongoing basis with a special attention for those with congenital long QT syndrome, congestive heart failure, bradyarrhythmias, and those taking drugs known to prolong the QT interval, including Class Ia and III antiarrhythmics. LENVIMA should be withheld in the event of development of QT interval prolongation greater than 500 ms. LENVIMA should be resumed at a reduced dose when QTc prolongation is resolved to < 480 ms or baseline (see DOSAGE AND ADMINISTRATION).

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. Periodic monitoring of ECG and electrolytes (magnesium, potassium and calcium) should be considered during treatment. Blood calcium levels should be monitored at least monthly and calcium should be replaced as necessary during LENVIMA treatment. LENVIMA 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

LENVIMA impairs exogenous thyroid suppression (see ADVERSE EFFECTS).

Hypothyroidism has been reported as very common in patients treated with LENVIMA in the RCC trial (see ADVERSE EFFECTS, Selected Adverse Reactions).

Thyroid function should be monitored before initiation of treatment, and periodically at least monthly throughout treatment with LENVIMA. Hypothyroidism should be treated according to standard medical practice to maintain euthyroid state.

Special Populations

Limited data are available for patients of ethnic origin other than Caucasian or Asian, in patients aged ≥ 75 years. LENVIMA should be used with caution in such patients, given the reduced tolerability of LENVIMA in Asian and elderly patients (see ADVERSE EFFECTS, Other Special Populations).

There are no data on the use of LENVIMA 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 of 4 weeks.

Patients with poor ECOG performance status

Patients with an ECOG performance status of 2 or higher were excluded from the RCC studies (see CLINICAL STUDIES). Patients with an ECOG performance 3 or higher were excluded from the DTC studies (see CLINICAL STUDIES). Benefit-risk in these patients has not been evaluated.

Effects on Fertility

Effects in humans are unknown. However, testicular and ovarian toxicity has been observed in rats, dogs, and monkeys.

No specific studies with lenvatinib have been conducted in animals to evaluate the effect on fertility. However, testicular and ovarian changes 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.

Use in Pregnancy (Category D)

There is limited information on the use of LENVIMA in pregnant women. Lenvatinib was embryotoxic and teratogenic when administered to rats and rabbits during organogenesis at exposures below the clinical exposure (based on body surface area) at the maximum recommended human dose. Fetal anomalies included parietal oedema, cryptophthalmia, abnormal tail (rats), retroesophageal subclavian artery, fused ribs, and vertebral abnormalities (rabbits). These embryofetal findings are probably related to the pharmacologic activity of lenvatinib as an antiangiogenic agent.

LENVIMA 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.

Women of childbearing potential

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

Use In Lactation

It is not known whether lenvatinib is excreted in human milk. Lenvatinib and its metabolites are excreted in rat milk and neonatal rats were more sensitive to the toxicity of lenvatinib compared to adults (See Paediatric Use below). Therefore, a risk to newborns or infants cannot be excluded and LENVIMA should not be used during breastfeeding.

Paediatric Use

Clinical data are not yet available in this population.

Mortality was the dose-limiting toxicity in juvenile rats in which dosing was initiated on postnatal day (PND) 7 or PND21. Mortality occurred at lower doses in neonatal rats (dosing initiated on PND7), or after a shorter duration of treatment in juvenile rats (dosing initiated on PND21). The exposure (as AUC) to lenvatinib in juvenile rats was lower compared to adults, suggesting increased susceptibility to the toxic effects of lenvatinib in young animals. 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.

Carcinogenicity

Carcinogenicity studies have not been conducted with LENVIMA.

Genotoxicity

Lenvatinib was not mutagenic in the *in vitro* Ames and mouse lymphoma tests and not clastogenic in an *in vivo* micronucleus assay in rats. These studies indicate a low genotoxic potential for LENVIMA.

Effects on ability to drive and use machines

No studies on the effects on the ability to drive and use machines have been performed. LENVIMA may cause side effects such as fatigue and dizziness. Patients who experience these symptoms should use caution when driving or operating machines.

INTERACTIONS WITH OTHER MEDICINES

Effect of other medicinal products on LENVIMA

CYP3A, P-gp, and BCRP inhibitors or inducers

LENVIMA may be administered regardless of co-administration with CYP3A, P-gp, and BCRP inhibitors. In healthy subjects, ketoconazole (400 mg for 18 days) increased lenvatinib (administered as a single dose on Day 5) $AUC_{0-\text{inf}}$ and AUC_{0-t} approximately 15% while C_{max} increased 19%. This is supported by a population PK analysis which found CYP3A4 inhibitors decreased CI/F by 7.8%.

LENVIMA may be co-administered without dose adjustment with CYP3A and P-gp inducers, based on a study in which healthy subjects were administered repeated doses of rifampicin (600 mg for 21 days) and a single dose of lenvatinib (24 mg, Day 15). $AUC_{0-\text{inf}}$ and AUC_{0-t} decreased approximately 18% while C_{max} did not change. The effect of CYP3A induction alone was estimated by comparing the PK parameters for lenvatinib following single and multiple doses of rifampicin. Lenvatinib AUC and C_{max} were predicted to decrease by 30% and 15%, respectively, after strong induction in the absence of acute P-gp inhibition. This is supported by a population PK analysis which found CYP3A4 inducers increased CI/F by 30%.

Gastric pH-altering agents

In a population pharmacokinetic analysis of patients receiving LENVIMA up to 24 mg once daily, agents which increase gastric pH (H₂ receptor blockers, proton pump inhibitors, antacids) did not have a significant effect on lenvatinib exposure.

Other chemotherapeutic agents

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

Effect of LENVIMA on other medicinal products

Cytochrome P450 or UGT enzyme substrates

Lenvatinib is considered neither a strong inhibitor nor an inducer of cytochrome P450 or uridine 5'-diphosphoglucuronosyl transferase (UGT) enzymes.

P-gp and BCRP substrates

Lenvatinib showed minimal inhibitory activities toward P-gp-mediated and BCRP-mediated transport activities. Similarly, no induction of P-gp mRNA expression was observed.

OAT, OCT, OATP, BSEP, and aldehyde oxidase substrates

Lenvatinib showed inhibitory effects on organic anion transporter (OAT)1, OAT3, organic cation transporter (OCT)1, OCT2, organic anion transporting polypeptide (OATP)1B1, and bile salt export pump (BSEP), but minimal or no inhibitory effect on OATP1B3. In human liver cytosol, lenvatinib did not inhibit aldehyde oxidase activity.

ADVERSE EFFECTS

Clinical Trials

Summary of safety profile

The safety profile of LENVIMA is based on a pooled analysis of safety data from clinical trials in which 1,166 patients were treated with LENVIMA including 458 patients with radioactive iodine refractory differentiated thyroid cancer (RAI - Refractory DTC), who received the recommended dose in the pivotal Phase 3 SELECT trial and two Phase 2 clinical trials, 62 patients with RCC who received the recommended dose of the combination of LENVIMA and everolimus, and 656 patients with other cancer types.

Radioactive iodine refractory differentiated thyroid cancer

The safety of LENVIMA was evaluated in 392 patients from the Phase 3 SELECT trial with radioactive iodine-refractory differentiated thyroid cancer (RAI-refractory DTC) randomized to receive LENVIMA 24 mg once daily (n=261) or placebo (n=131) (see CLINICAL TRIALS).

In the SELECT study, the most common adverse reactions observed in LENVIMA-treated patients (greater than or equal to 30%) were, in order of decreasing frequency, hypertension, fatigue, diarrhoea, arthralgia/myalgia, decreased appetite, weight decreased, nausea, stomatitis, headache, vomiting, proteinuria, palmar-plantar erythrodysesthesia (PPE) syndrome, abdominal pain, and dysphonia. The most common serious adverse reactions (at least 2%) were pneumonia (4%), hypertension (3%), and dehydration (3%).

Adverse reactions led to dose reductions in 68% of patients receiving LENVIMA and 5% of patients receiving placebo; 18% of patients discontinued LENVIMA and 5% discontinued placebo for adverse reactions. The most common adverse reactions (at least 10%) resulting in dose reductions of LENVIMA were hypertension (13%), proteinuria (11%), decreased appetite (10%), and diarrhoea (10%); the most common adverse reactions (at least 1%) resulting in discontinuation of LENVIMA were hypertension (1%) and asthenia (1%).

Table 4 presents the incidence rates of treatment-emergent adverse events observed in the double blind phase of the DTC study. All adverse events occurring with a treatment difference of at least 5% over placebo are included in the Table. Clinically significant events (CSEs) that were observed more frequently than placebo are also included based on an assessment of the known pharmacology of LENVIMA and class effects

Table 4 Treatment-Emergent Adverse Events reported for LENVIMA in the double-blind phase of the DTC Study*

System Organ Class Preferred Term	LENVIMA 24 mg N=261		Placebo N=131	
	All Grades (%)	Grades 3-4 (%)	All Grades (%)	Grades 3-4 (%)
Blood & Lymphatic System Disorders				
Thrombocytopenia ^a	13.8	1.9	2.3	0
Lymphopenia ^b	10.7	2.3	4.6	0.8
Splenic infarction	0.8	0	0	0
Cardiac Disorders				
Ejection fraction decreased	5.4	1.1	0.8	0
Myocardial infarction ^{c,d}	1.1	1.1	0.8	0.8
Cardiac failure	0.8	0	0	0
Endocrine Disorders				
Hypothyroidism	5.4	0	0	0
Gastrointestinal Disorders				
Diarrhoea	67.4	9.2	16.8	0
Nausea	46.7	2.3	25.2	0.8
Stomatitis ^e	41.0	4.6	8.4	0
Vomiting	35.6	1.9	14.5	0
Abdominal pain ^f	31.4	2.3	10.7	0.8
Constipation	28.7	0.4	15.3	0.8
Oral pain ^g	24.9	1.1	2.3	0
Dry mouth	16.9	0.4	8.4	0
Dyspepsia	13.0	0.4	3.8	0
Flatulence	6.1	0	0.8	0
Anal fistula	1.1	0.4	0	0
General Disorders and Administration Site Conditions				
Fatigue	42.5	4.6	24.4	1.5
Asthenia	25.3	6.1	13.0	2.3
Oedema peripheral	20.7	0.4	7.6	0
Malaise	5.4	0	0	0
Hepatobiliary Disorders				
Hepatocellular damage / hepatitis ^h	1.1	0.8	0	0
Infections and Infestations				
Urinary tract infection	11.5	1.1	5.3	0
Perineal abscess	0.8	0.8	0	0
Investigations				
Weight decreased	51.3	13.4	14.5	0.8
Electrocardiogram QT prolonged	8.8	1.5	1.5	0
Alanine aminotransferase increased	7.7	1.5	0	0
Blood creatinine increased	7.3	0	1.5	0
Aspartate aminotransferase increased	6.9	1.9	1.5	0
Blood thyroid stimulating hormone	6.5	0	0	0

Table 4 Treatment-Emergent Adverse Events reported for LENVIMA in the double-blind phase of the DTC Study*

System Organ Class Preferred Term	LENVIMA 24 mg N=261		Placebo N=131	
	All Grades (%)	Grades 3-4 (%)	All Grades (%)	Grades 3-4 (%)
increased				
Blood alkaline phosphatase increased	6.1	0.8	2.3	0.8
Blood urea increased	3.1	0	0	0
Hepatic function abnormal	2.3	0.4	0	0
Blood bilirubin increased	1.9	0	0	0
Gamma-glutamyltransferase increased	1.5	0.8	0.8	0
Metabolism and Nutrition Disorders				
Decreased appetite	54.4	6.9	18.3	0.8
Hypokalaemia	13.8	3.4	3.8	0
Hypocalcaemia	12.6	5.0	0	0
Hypoalbuminaemia	9.6	0.4	1.5	0
Dehydration	8.8	2.3	2.3	0.8
Hypomagnesaemia ⁱ	6.5	0.4	1.5	0
Hypercholesterolaemia ^j	5.0	0.4	0	0
Musculoskeletal and Connective Tissue Disorders				
Arthralgia	26.1	0.4	6.9	0.8
Myalgia	19.2	1.5	4.6	0
Back pain	17.6	1.9	9.2	0
Musculoskeletal pain	16.1	0.4	8.4	0.8
Pain in extremity	15.3	1.1	6.9	1.5
Nervous System Disorders				
Headache	38.3	3.1	11.5	0.8
Dysgeusia	18.0	0	3.1	0
Dizziness	15.3	0.4	9.2	0
Monoparesis	1.1	0.8	0	0
Cerebrovascular accident	0.8	0.4	0	0
Transient ischemic attack	0.8	0	0	0
Reversible posterior leucoencephalopathy syndrome	0.4	0	0	0
Psychiatric Disorders				
Insomnia	11.9	0	3.1	0
Renal and Urinary Disorders				
Proteinuria	33.7	10.7	3.1	0
Renal failure events ^{d,k}	5.0	2.7	0.8	0.8
Renal impairment	1.9	0.4	0	0
Respiratory, Thoracic, and Mediastinal Disorders				
Dysphonia	31.4	1.1	5.3	0
Cough	23.8	0	17.6	0
Pulmonary embolism ^d	3.1	3.1	1.5	1.5
Skin and Subcutaneous Tissue Disorders				
Palmar-plantar erythrodysesthesia syndrome	32.2	3.4	0.8	0
Rash	18.8	0.4	1.5	0
Alopecia	12.3	0	5.3	0
Hyperkeratosis	6.9	0	1.5	0
Palmar erythema	1.1	0	0	0

Table 4 Treatment-Emergent Adverse Events reported for LENVIMA in the double-blind phase of the DTC Study*

System Organ Class Preferred Term	LENVIMA 24 mg N=261		Placebo N=131	
	All Grades (%)	Grades 3-4 (%)	All Grades (%)	Grades 3-4 (%)
Vascular Disorders				
Haemorrhage ^{d,1}	34.9	1.5	18.3	3.1
Hypertension ^m	72.8	44.4	16.0	3.8
Hypotension	8.8	1.5	2.3	0

- a: Includes the following terms: thrombocytopenia, platelet count decreased
- b: Includes the following terms: lymphopenia, lymphocyte count decreased
- c: Includes the following terms: acute myocardial infarction, myocardial infarction
- d: includes fatal events and these are counted in all Grade column
- e: Includes the following terms: aphthous stomatitis, stomatitis, glossitis, mouth ulceration, mucosal inflammation
- f: Includes the following terms: abdominal discomfort, abdominal pain, abdominal pain lower, abdominal pain upper, abdominal tenderness, epigastric discomfort, gastrointestinal pain
- g: Includes the following terms: oral pain, glossodynia, oropharyngeal pain
- h: Includes the following terms: drug-induced liver injury, cholestatic liver injury, hepatic steatosis
- i: Includes the following terms: hypomagnesaemia, blood magnesium decreased
- j: Includes the following terms: hypercholesterolaemia and blood cholesterol increased
- k: Includes the following terms: acute prerenal failure, renal failure, renal failure acute, renal tubular necrosis
- l: Includes the following terms: epistaxis, haematuria, contusion, gingival bleeding, haematochezia, pulmonary haemorrhage, vaginal haemorrhage, rectal haemorrhage, haematoma, haemorrhoidal haemorrhage, laryngeal haemorrhage, petechiae, intracranial tumour haemorrhage, haemorrhagic stroke, pleural haemorrhage, splenic haemorrhage, blood urine present, conjunctival haemorrhage, eye haemorrhage, gastroduodenitis haemorrhagic, haematemesis, increased tendency to bruise, proctitis haemorrhagic, purpura, renal haematoma, skin haemorrhage, splinter haemorrhages
- m: Includes the following terms: hypertension, hypertensive crisis, blood pressure diastolic increased, blood pressure increased

*TEAEs reported at 4 months after the cut-off for the final PFS analysis

Renal Cell Carcinoma

The most common adverse reactions observed in the LENVIMA in combination with everolimus-treated group (> 30%) were, in order of decreasing frequency, diarrhoea, fatigue, arthralgia/myalgia, decreased appetite, vomiting, nausea, stomatitis/oral inflammation, hypertension, peripheral oedema, cough, abdominal pain, dyspnoea, rash, weight decreased, haemorrhagic events, and proteinuria. The most common serious adverse reactions ($\geq 5\%$) were renal failure (11%), dehydration (10%), anaemia (6%), thrombocytopenia (5%), diarrhoea (5%), vomiting (5%), and dyspnoea (5%).

Adverse reactions led to dose reductions or interruption in 89% of patients receiving LENVIMA + everolimus and 54% in patients receiving everolimus. The most common adverse reactions ($\geq 5\%$) resulting in dose reductions in the LENVIMA + everolimus-treated group were diarrhoea (21%), fatigue (8%), thrombocytopenia (6%), vomiting (6%), nausea (5%), and proteinuria (5%).

Treatment discontinuation due to an adverse reaction occurred in 29% of patients in the LENVIMA + everolimus-treated group and 12% of patients in the everolimus-treated group.

Table 5 presents the adverse reactions in > 15% of patients in the LENVIMA + Everolimus arm.

Table 5 Grades 1-4 Adverse Reactions in > 15% of Patients in the LENVIMA + Everolimus Arm

System Organ Class Preferred Term	LENVIMA 18 mg + Everolimus 5 mg (N=62)		Everolimus 10 mg (N=50)	
	Grade 1-4 (%)	Grade 3-4 (%)	Grade 1-4 (%)	Grade 3-4 (%)
Endocrine Disorders				
Hypothyroidism	24	0	2	0
Gastrointestinal Disorders				
Constipation	16	0	18	0
Diarrhoea	81	19	34	2
Dyspepsia/Gastro-oesophageal reflux	21	0	12	0
Abdominal pain ^a	37	3	8	0
Nausea	45	5	16	0
Oral pain ^b	23	2	4	0
Stomatitis/Oral inflammation ^c	44	2	50	4
Vomiting	48	7	12	0
General Disorders and Administration Site Conditions				
Fatigued	73	18	40	2
Peripheral oedema	42	2	20	0
Pyrexia/Increased body temperature	21	2	10	2
Investigations				
Weight decreased	34	3	8	0
Metabolism and Nutrition Disorders				
Decreased appetite	53	5	18	0
Musculoskeletal and Connective Tissue Disorders				
Arthralgia/Myalgia ^e	55	5	32	0
Musculoskeletal chest pain	18	2	4	0
Nervous System Disorders				
Headache	19	2	10	2
Psychiatric Disorders				
Insomnia	16	2	2	0
Renal and Urinary Disorders				
Proteinuria/Urine protein present	31	8	14	2
Renal failure event ^f	18	10	12	2
Respiratory, Thoracic and Mediastinal Disorders				
Cough	37	0	30	0
Dysphonia	18	0	4	0
Dyspnoea/Exertional dyspnoea	35	5	28	8
Skin and Subcutaneous Tissue Disorders				
Rash ^g	35	0	40	0
Vascular Disorders				
Haemorrhagic events ^h	32	6	26	2
Hypertension/Increased blood pressure	42	13	10	2

- a: Includes abdominal discomfort, gastrointestinal pain, lower abdominal pain, and upper abdominal pain
- b: Includes gingival pain, glossodynia, and oropharyngeal pain
- c: Includes aphthous stomatitis, gingival inflammation, glossitis, and mouth ulceration
- d: Includes asthenia, fatigue, lethargy and malaise
- e: Includes arthralgia, back pain, extremity pain, musculoskeletal pain, and myalgia
- f: Includes blood creatinine increased, blood urea increased, creatinine renal clearance decreased, nephropathy toxic, renal failure, renal failure acute, and renal impairment
- g: Includes erythema, erythematous rash, genital rash, macular rash, maculo-papular rash, , papular rash, pruritic rash, pustular rash, and septic rash
- h: Includes haemorrhagic diarrhea, epistaxis, gastric haemorrhage, haemarthrosis, haematoma, haematuria, haemoptysis, lip haemorrhage, renal haematoma, and scrotal haematocele

Table 6 Grade 3-4 Laboratory Abnormalities in \geq 3% of Patients in the LENVIMA + Everolimus Arm^{a,b}

Laboratory Abnormality	LENVIMA 18 mg + Everolimus 5 mg N=62	Everolimus 10 mg N=50
	Grades 3-4 (%)	Grades 3-4 (%)
Chemistry		
Aspartate aminotransferase (AST) increased	3	0
Alanine aminotransferase (ALT) increased	3	2
Alkaline phosphatase increased	3	0
Hyperkalaemia	6	2
Hypokalaemia	6	2
Hyponatraemia	11	6
Hypocalcaemia	6	2
Hypophosphataemia	11	6
Hyperglycaemia	3	16
Hypertriglyceridaemia	18	18
Elevated cholesterol	11	0
Creatine kinase increased	3	4
Lipase increased	13	12
Hematology		
Haemoglobin decreased	8	16
Platelet count decreased	5	0
Lymphocyte count decreased	10	20

- a: With at least 1 grade increase from baseline
- b: Subject with at least 1 post baseline laboratory value

Post-marketing adverse drug reactions

The following adverse reactions have been identified during post approval use of LENVIMA. Because these reactions are reported voluntarily from a population of uncertain size, it is not always possible to reliably estimate their frequency or establish a causal relationship to drug exposure.

Gastrointestinal Disorders: amylase increased, lipase increased, pancreatitis

Hepatobiliary Disorders: cholecystitis

Selected Adverse Reactions in DTC and RCC

Hypertension

In the pivotal DTC Phase 3 SELECT trial (see CLINICAL TRIALS), hypertension (including hypertension, hypertensive crisis, blood pressure diastolic increased, and blood pressure increased) was reported in 72.8% of LENVIMA-treated patients and 16.0% of patients in the placebo-treated group. The median time to onset in LENVIMA-treated patients was 16 days. Events of Grade 3 or higher (including 1 event of Grade 4) occurred in 44.4% of LENVIMA-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.

In the RCC study (see CLINICAL TRIALS) Phase 1b plus Phase 2 population, hypertension was reported in 41.9% of patients in the LENVIMA plus everolimus-treated group (the incidence of Grade 3 or Grade 4 hypertension was 12.9%) and 10.0% of patients in the everolimus-treated group (the incidence of Grade 3 or Grade 4 hypertension was 2.0%). The median time to onset was 4.9 weeks (any grade) and 6.9 weeks (Grade \geq 3) in the LENVIMA plus everolimus-treated group.

Proteinuria

In the pivotal DTC Phase 3 SELECT trial (see CLINICAL TRIALS), proteinuria was reported in 33.7% of LENVIMA treated patients and 3.1% of patients in the placebo-treated group. The median time to onset was 6.7 weeks. Grade 3 events occurred in 10.7% of LENVIMA-treated patients and no 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.

In the RCC study (see CLINICAL TRIALS) Phase 1b plus Phase 2 population, proteinuria was reported in 30.6% of patients in the LENVIMA plus everolimus-treated group (8.1% were Grade \geq 3) and 14.0% of patients in the everolimus-treated group (2.0% were Grade 3 or greater). The median time to onset of proteinuria was 6.1 weeks (any grade) and 20.1 weeks (Grade \geq 3) in the LENVIMA plus everolimus-treated group. Proteinuria led to permanent treatment discontinuation in 4.8% of patients.

Renal failure and impairment

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

In the RCC study (see CLINICAL TRIALS) Phase 1b plus Phase 2 population, 8.1% of patients in the LENVIMA plus everolimus treated group developed renal failure and 3.2% developed renal impairment (9.7% of patients had a Grade 3 event of renal failure or impairment). In the everolimus monotherapy group 2.0% of patients developed renal failure (2.0% were Grade 3).

Cardiac dysfunction

In the pivotal DTC Phase 3 SELECT trial (see CLINICAL TRIALS), decreased ejection fraction/cardiac failure was reported in 6.5% of patients (1.5% were Grade \geq 3) in the LENVIMA treated group, and 2.3% in the placebo group (none were Grade \geq 3).

In the RCC study (see CLINICAL TRIALS) Phase 1b plus Phase 2 population, decreased ejection fraction/cardiac failure was reported in 4.8% of patients (3.2% were Grade \geq 3) in the LENVIMA plus everolimus treated group, and 4.0% in the everolimus group (2.0% were Grade \geq 3). The median time to onset of decreased ejection fraction and cardiac failure was 15.7 weeks (any grade) and 32.8 weeks (Grade \geq 3) in the LENVIMA plus everolimus-treated group.

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

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

In the RCC study (see CLINICAL TRIALS) Phase 1b plus Phase 2 population, there was 1 event of PRES (Grade 3) in the LENVIMA-treated group, occurring after 18.4 weeks of treatment. There were no reports in the LENVIMA plus everolimus or everolimus monotherapy groups.

Amongst 1,166 patients treated with LENVIMA, 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.

Hepatotoxicity

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

In the RCC study (see CLINICAL TRIALS) Phase 1b plus Phase 2 population, the most commonly reported liver-related adverse reactions in the LENVIMA plus everolimus-treated group were elevations of liver enzyme levels, including increases in alanine aminotransferase (9.7%), aspartate aminotransferase (4.8%), alkaline phosphatase (4.8%), and blood bilirubin (3.2%). The median time to onset of liver events was 6.7 weeks (any grade) and 14.2 weeks

(Grade ≥ 3) in the LENVIMA plus everolimus-treated group. Grade 3 liver-related reactions occurred in 3.2% of LENVIMA plus everolimus-treated patients. Liver-related reactions led to dose interruptions and reductions in 1.6% and 1.6% of patients, respectively, and to permanent discontinuation in 3.2% of patients.

Amongst 1,166 patients treated with LENVIMA, 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.

Arterial thromboembolisms

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

In the RCC study (see CLINICAL TRIALS) Phase 1b plus Phase 2 population, 1.6% of patients in the LENVIMA plus everolimus-treated group reported arterial thromboembolic events. The time to onset was 69.6 weeks. In the everolimus group, 6.0% of patients reported an arterial thromboembolism (4.0% were Grade ≥ 3).

Amongst 1,166 patients treated with LENVIMA, there were 4 cases (0.3%) of arterial thromboembolisms (2 cases of myocardial infarction and 2 cases of cerebrovascular accident) with a fatal outcome.

Haemorrhagic events and thrombocytopenia

In the pivotal DTC Phase 3 SELECT trial (see CLINICAL TRIALS), haemorrhagic events were reported in 34.9% of LENVIMA-treated patients versus 18.3% of placebo-treated patients. Events 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%). When adjusted to account for the 4-fold greater duration of exposure in the LENVIMA versus the placebo arm, the following reactions occurred less frequently on LENVIMA than placebo: haemoptysis (0.05 episodes/subject-year on LENVIMA vs. 0.21 episodes/subject-year on placebo) and pulmonary haemorrhage (0.02 episodes/subject year on LENVIMA vs. 0.09 episodes/subject-year on placebo).

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

Thrombocytopenia or platelet count decreased was reported in 15.3% of LENVIMA-treated patients (1.9% were Grade ≥ 3) versus 2.3% in the placebo arm (none were Grade ≥ 3). There were no Grade 4 TEAEs. The incidence of serious thrombocytopenia was 0.8%. Most reactions resolved following supportive treatment, although 5.0% of patients had events of thrombocytopenia that required dose interruption, and 1.9% required dose reduction. No patients discontinued treatment as a result of thrombocytopenia.

In the RCC study (see CLINICAL TRIALS) Phase 1b plus Phase 2 population, haemorrhage was reported in 38.7% (8.1% were Grade 3 or greater) of patients in the LENVIMA plus

everolimus-treated group. Reactions that occurred at an incidence of $\geq 2.0\%$ were: epistaxis (22.6%), haematuria (4.8%), haematoma (3.2%), and gastric haemorrhage (3.2%). The median time to first onset was 10.2 weeks (any grade) and 7.6 weeks (Grade ≥ 3) in the LENVIMA plus everolimus-treated group. The incidence of serious haemorrhage was 4.8% (cerebral haemorrhage, gastric haemorrhage and haemarthrosis). Discontinuation due to haemorrhagic events occurred in 3.2% of patients in the LENVIMA plus everolimus-treated group. There was one case of fatal cerebral haemorrhage in the LENVIMA plus everolimus-treated group and one case of fatal intracranial haemorrhage in the LENVIMA-treated group.

Thrombocytopenia or platelet count decreased was reported in 14.5% of patients in the LENVIMA plus everolimus-treated group (4.8% were Grade ≥ 3) and 10.0% of patients in the everolimus-treated group (none were Grade ≥ 3). There was one Grade 4 TEAE. The incidence of serious thrombocytopenia was 4.8%. Dose reduction or interruption due to thrombocytopenia occurred in 9.7% of patients and 3.2% of patients discontinued treatment due to thrombocytopenia.

Amongst 1166 patients treated with LENVIMA, 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.

Hypocalcaemia

In the pivotal DTC Phase 3 SELECT trial (see CLINICAL TRIALS), hypocalcaemia was reported in 12.6% of LENVIMA treated patients vs. no events in the placebo arm. The median time to first onset in LENVIMA-treated patients was 11.1 weeks. Events of Grade 3 or 4 severity occurred in 5.0% of LENVIMA-treated vs 0 placebo-treated patients. Most events 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.

In the RCC study (see CLINICAL TRIALS) Phase 1b plus Phase 2 population, hypocalcaemia was reported in 8.1% of patients in the LENVIMA plus everolimus-treated group (3.2% were Grade ≥ 3) and 4.0% of patients in the everolimus-treated group (none were Grade ≥ 3). The median time to onset of hypocalcaemia was 28.3 weeks (any grade) and 45.9 weeks (Grade ≥ 3) in the LENVIMA plus everolimus-treated group. There was one Grade 4 TEAE. No events of hypocalcaemia required dose reduction or interruption, and no patients discontinued treatment due to hypocalcaemia.

Gastrointestinal perforation and fistula formation

In the pivotal DTC Phase 3 SELECT trial (see CLINICAL TRIALS), events of gastrointestinal perforation or fistula were reported in 1.9% of LENVIMA treated patients and 0.8% of patients in the placebo group.

In the RCC study (see CLINICAL TRIALS) Phase 1b plus Phase 2 population, 1.6% of cases of perforated appendicitis (of Grade 3) occurred in the LENVIMA plus everolimus-treated group; there were no reports in the LENVIMA or everolimus groups.

Non-Gastrointestinal fistulae (see PRECAUTIONS)

LENVIMA 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 LENVIMA, with a median latency of about 3 months.

QT interval prolongation

In the pivotal DTC Phase 3 SELECT trial (see CLINICAL TRIALS), QT/QTc interval prolongation was reported in 8.8% of LENVIMA 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 LENVIMA-treated patients compared to no reports in the placebo group.

In the RCC study (see CLINICAL TRIALS) Phase 1b plus Phase 2 population, QTc interval increases greater than 60 ms were reported in 11% of patients in the LENVIMA plus everolimus-treated group. The incidence of QTc interval greater than 500 ms was 6% in the LENVIMA plus everolimus-treated group. No reports of QTc interval prolongation greater than 500 ms or increases greater than 60 ms occurred in the everolimus-treated group.

Blood thyroid stimulating hormone increased (see PRECAUTIONS Impairment of thyroid stimulating hormone suppression / Thyroid dysfunction)

In the pivotal DTC Phase 3 SELECT trial (see CLINICAL TRIALS), 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 LENVIMA-treated patients as compared with 14% of placebo-treated patients.

In the RCC study (see CLINICAL TRIALS) Phase 1b plus Phase 2 population, hypothyroidism occurred in 24% of patients in the LENVIMA plus everolimus-treated group and 2% of patients in the everolimus-treated group. All events of hypothyroidism in the LENVIMA plus everolimus-treated group were of Grade 1 or 2. In patients with a normal TSH at baseline, an elevation of TSH level was observed post baseline in 60.5% of LENVIMA plus everolimus-treated patients as compared with none in patients receiving everolimus alone.

Dyslipidaemia

In the pivotal DTC Phase 3 SELECT trial (see CLINICAL TRIALS), hypercholesterolaemia or blood cholesterol increased were reported in 5.0% of LENVIMA treated patients (0.4% were Grade \geq 3) vs. no events in the placebo arm. Hypertriglyceridaemia or blood triglycerides increased were reported in 2.7% of LENVIMA treated patients (0.8% were Grade \geq 3) vs. no events in the placebo arm. Most events resolved following supportive treatment, without dose interruption or reduction, which occurred in 0.4% and 0.4% of patients, respectively. No patients discontinued treatment as a result of dyslipidaemia.

In the RCC study (see CLINICAL TRIALS) Phase 1b plus Phase 2 population, dyslipidaemia was reported in 50.0% of patients in the LENVIMA plus everolimus-treated group (17.7% were Grade \geq 3) and 34.0% of patients in the everolimus-treated group (8.0% were Grade \geq 3). Reactions that occurred at an incidence of \geq 30.0% were: hypertriglyceridaemia (40.3%), and hypercholesterolaemia (30.6%). The median time to onset of dyslipidaemia was 4.1 weeks (any grade) and 6.1 weeks (Grade \geq 3) in the LENVIMA plus everolimus-treated group. The incidence of serious dyslipidaemia was 1.6%. Most reactions resolved following supportive treatment, although 8.1% of patients had events of dyslipidaemia that required dose interruption, and 3.2% required dose reduction. No patients discontinued treatment as a result of dyslipidaemia.

Diarrhoea

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

In the RCC study (see CLINICAL TRIALS) Phase 1b plus Phase 2 population, diarrhoea was reported in 80.6% of patients in the LENVIMA plus everolimus-treated group (21.0% were Grade \geq 3) and in 34.0% of patients in the everolimus-treated group (2.0% were Grade \geq 3). The median time to onset was 4.1 weeks (any grade) and 8.1 weeks (Grade \geq 3) in the LENVIMA plus everolimus-treated group. Diarrhoea was the most frequent cause of dose interruption/reduction and recurred despite dose reduction. Diarrhoea resulted in discontinuation in one patient.

Other special populations

Elderly

In DTC, patients of age \geq 75 years were more likely to experience Grade 3 or 4 hypertension, proteinuria, decreased appetite, and dehydration. There are limited data on patients of age \geq 75 years with RCC.

Sex

In 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.

Race

In DTC, Asian patients had a higher incidence than Caucasian patients of oedema peripheral, hypertension, fatigue, PPE, proteinuria, thrombocytopenia, and blood thyroid stimulating hormone increased. Japanese patients had a higher incidence of Grade 3 or 4 hypertension, decreased appetite, fatigue, and thrombocytopenia compared with non-Japanese subjects. There are limited data on Asian patients with RCC.

Baseline hypertension

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

Hepatic impairment

In 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. There are limited data on patients with hepatic impairment in RCC.

Renal impairment

In DTC, patients with baseline renal impairment had a higher incidence of Grade 3 or 4 hypertension, proteinuria, fatigue, stomatitis, oedema peripheral, thrombocytopenia, dehydration, prolonged electrocardiogram QT, hypothyroidism, hyponatraemia, blood thyroid stimulating hormone increased and pneumonia compared with subjects with normal

renal function. These patients also had a higher incidence of renal events and a trend towards a higher incidence of liver events.

See also PRECAUTIONS, DOSAGE AND ADMINISTRATION.

Patients with body weight < 60 kg

In DTC, patients with low body weight (< 60 kg) had a higher incidence of PPE, proteinuria, Grade 3 or 4 hypocalcaemia and hyponatraemia, and a trend towards a higher incidence of Grade 3 or 4 decreased appetite. There are limited data on patients with body weight < 60 kg in RCC.

DOSAGE AND ADMINISTRATION

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

Starting dose in RAI – Refractory DTC

The recommended dose of LENVIMA is 24 mg (two 10 mg capsules plus one 4 mg capsule) taken once daily. The daily dose is to be modified as needed according to the dose/toxicity management plan (see dose adjustment section below).

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

Starting dose in Advanced Renal Cell Carcinoma

The recommended daily dose of LENVIMA is 18 mg (one 10 mg capsule and two 4 mg capsules) once daily in combination with 5 mg everolimus once daily. The daily doses of LENVIMA, and if necessary, everolimus are to be modified as needed according to the dose/toxicity management plan (see dose adjustment section below).

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

Dose adjustment during therapy

Management of adverse reactions may require dose interruption, adjustment, or discontinuation of LENVIMA or LENVIMA and everolimus if treating in the combination (See PRECAUTIONS).

Medical management of nausea, vomiting and diarrhoea should be optimised to reduce the risk of dehydration and renal failure (see PRECAUTIONS, Renal failure and impairment) prior to any LENVIMA therapy interruption or dose reduction.

For toxicities thought to be related to LENVIMA, general advice about dose management is included in Table 7, and specific daily dose modifications for thyroid cancer patients are in Table 8, and for renal cell carcinoma patients in Table 9.

For toxicities thought to be related to everolimus, when treating renal cell carcinoma using the combination of LENVIMA and everolimus, everolimus treatment should be interrupted, reduced to alternate day dosing, or discontinued (see the everolimus PI for advice on specific adverse reactions).

For toxicities thought to be related to both LENVIMA and everolimus, when treating renal cell carcinoma using the combination, LENVIMA (See Table 7) should be reduced prior to reducing everolimus.

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

Table 7 Dose modifications for adverse reactions

Adverse Reaction	CTCAE Grade	Action	Dose Reduce and Resume LENVIMA
Hypertension	Grade 3 (despite optimal antihypertensive therapy)	Interrupt	Resolves to Grade 0, 1 or 2. See detailed guidance in Table 3 in PRECAUTIONS, Hypertension section
	Grade 4	Discontinue	Do not resume
Proteinuria	≥2 gm/24 hours	Interrupt	Resolves to less than 2 gm/24 hours
Nephrotic syndrome	-----	Discontinue	Do not resume
Renal impairment or failure	Grade 3	Interrupt	Resolves to Grade 0-1 or baseline
	Grade 4*	Discontinue	Do not resume
Cardiac failure	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 and Thrombocytopenia*	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
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)

Table 8 Dose Modifications from Recommended Daily Dose for treatment of DTC

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 one 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.

Table 9 Dose Modifications from Recommended Daily Dose for treatment of RCC

Dose Level	Daily Dose	Number of Capsules
Recommended daily dose	18 mg orally once daily	One 10 mg capsule plus two 4 mg capsules
First dose reduction	14 mg orally once daily	One 10 mg capsule plus one 4 mg capsule
Second dose reduction	10 mg orally once daily	One 10 mg capsule
Third dose reduction	8 mg orally once daily ^a	Two 4 mg capsules

a: Limited data are available for doses below 8 mg

Special populations

No data with the combination are available for most of the special populations. The following information is derived from the clinical experience on single agent LENVIMA in patients with RAI - Refractory DTC.

All patients other than those with severe hepatic or renal impairment (see below) should initiate treatment at the recommended dose of 24 mg LENVIMA taken once daily for thyroid cancer and 18 mg of LENVIMA with 5 mg of everolimus taken once daily for renal cell carcinoma, following which the dose should be further adjusted on the basis of individual tolerability.

Certain subpopulations of patients appear to have reduced tolerability to LENVIMA (see ADVERSE EFFECTS, Special Populations). Following treatment initiation at the recommended dose, the dose should be adjusted on the basis of individual tolerability.

Patients with hepatic impairment

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 for RAI - Refractory DTC is 14 mg LENVIMA taken once daily and for RCC is 10 mg of LENVIMA taken once daily in combination with the dose of everolimus recommended for patients with severe hepatic impairment in the everolimus PI. Further dose adjustments may be necessary on the basis of individual tolerability. The combination should be used in patients with severe hepatic impairment only if the anticipated benefit exceeds the risk.

Patients with renal impairment

No adjustment of the starting dose is required on the basis of renal function in patients with mild (CrCL 60 -89 mL/min) or moderate (CrCl 30-59 mL/min) renal impairment. In patients with severe renal impairment (CrCL <30 mL/min), the recommended starting dose for thyroid cancer is 14 mg LENVIMA taken once daily and for renal cell carcinoma is 10 mg of LENVIMA plus 5 mg everolimus taken once daily. Further dose adjustments may be necessary based on the individual tolerability. Patients with end-stage renal disease were not studied, therefore the use of LENVIMA in these patients is not recommended.

Patients with hypertension

Blood pressure should be well controlled prior to treatment with LENVIMA, and should be regularly monitored during treatment (see PRECAUTIONS and ADVERSE EFFECTS).

Elderly population

No adjustment of the starting dose is required on the basis of age. Limited data are available on the use in patients aged ≥ 75 years.

Paediatric population

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

Race

No adjustment of starting dose is required on the basis of race. Limited data are available on use in patients from ethnic origins other than Caucasian or Asian.

Body weight below 60 kg

No adjustment of starting dose is required on the basis of body weight. Limited data are available on patients with a body weight below 60 kg with RCC (see ADVERSE EFFECTS, Other special populations).

Method of administration

LENVIMA should be taken at about the same time each day, with or without food. The capsules should be swallowed whole with water.

Alternatively, if unable to swallow the capsule whole place the capsule, without breaking or crushing, in a glass of approximately 25 mL of water or apple juice. The capsules must be left to disintegrate in the liquid for at least 10 minutes and then gently stirred for at least 3 minutes to dissolve the capsules shells. The suspension is to be swallowed. After drinking, the same amount of water or apple juice (25 mL) must be added to the glass and swirled a few times. The additional liquid must be swallowed. Do not mix more than one medicine in the glass at the same time.

The person preparing the suspension should ensure their hands are thoroughly washed on completion of preparation and taking of the medication.

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.

OVERDOSAGE

Contact the Poisons Information Centre on telephone 13 11 26 for advice on management of overdose.

There have been reports of overdose with LENVIMA including a single administration of 144 mg, 6 times the recommended daily dose. These cases were associated with adverse reactions consistent with the known safety profile of LENVIMA, or were without adverse reactions. There is no specific antidote for overdose with LENVIMA, due to the high plasma protein binding, lenvatinib is not expected to be dialyzable. In case of suspected overdose, LENVIMA should be withheld and appropriate supportive care given as required.

PRESENTATION AND STORAGE

Presentation

LENVIMA 4 mg hard capsule: 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 4 mg hard capsules are available in polyamide/aluminium/PVC/aluminium blisters of 30 capsules.

LENVIMA 10 mg hard capsule: 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.

LENVIMA 10 mg hard capsules are available in polyamide/aluminium/PVC/aluminium blisters of 30 capsules.

Storage

Store below 30°C.

NAME AND ADDRESS OF THE SPONSOR

Eisai Australia Pty Ltd
Level 2, 437 St Kilda Road
Melbourne, VIC, 3004

POISON SCHEDULE OF THE MEDICINE

S4

DATE OF FIRST INCLUSION IN THE AUSTRALIAN REGISTER OF THERAPEUTIC GOODS (THE ARTG)

28 January 2016

DATE OF MOST RECENT AMENDMENT

26 July 2017