HIGHLIGHTS OF PRESCRIBING INFORMATION
These highlights do not include all the information needed to use ROMVIMZA™ safely and effectively. See full prescribing information for ROMVIMZA™.

ROMVIMZA[™] (vimseltinib) capsules, for oral use Initial U.S. Approval: 2025

----INDICATIONS AND USAGE----

ROMVIMZA is a kinase inhibitor indicated for treatment of adult patients with symptomatic tenosynovial giant cell tumor (TGCT) for which surgical resection will potentially cause worsening functional limitation or severe morbidity. (1)

---DOSAGE AND ADMINISTRATION----

- <u>Recommended Dosage</u>: 30 mg orally twice weekly, with a minimum of 72 hours between doses as described in the blister package. (2.1)
- See full prescribing information for dosage modifications due to hepatotoxicity and drug interactions. (2.2, 2.3)

------Capsules: 14 mg, 20 mg, 30 mg. (3)

-----CONTRAINDICATIONS-----

None (4)

-----WARNINGS AND PRECAUTIONS-----

<u>Hepatotoxicity:</u> Elevated AST and ALT can occur. Evaluate liver tests prior to initiation of treatment and during treatment. (2.2, 5.1)

Embryo-fetal toxicity: Can cause fetal harm. Advise patients of reproductive potential of the potential risk to a fetus and to use effective contraception. (5.2, 8.1, 8.3)

Allergic Reactions to FD&C Yellow No. 5 (tartrazine) and No. 6 (Sunset Yellow FCF): 14 mg capsule contains FD&C Yellow No. 6 (Sunset Yellow FCF); 20 mg capsule contains FD&C Yellow No.5 (tartrazine) and No. 6 (Sunset Yellow FCF) as color additives, which may cause allergic reactions (including bronchial asthma) in certain susceptible patients. (5.3)

<u>Increased serum creatinine without affecting renal function:</u> Increases in serum creatinine can occur. Use alternative measures that are not based on serum creatinine to assess renal function. (5.4)

----ADVERSE REACTIONS---

Most common adverse reactions (incidence ≥20%), including laboratory abnormalities are increased AST, periorbital edema, fatigue, rash, increased cholesterol, peripheral edema, face edema, decreased neutrophils, decreased leukocytes, pruritus, and increased ALT. (6.1)

To report SUSPECTED ADVERSE REACTIONS, contact Deciphera Pharmaceuticals, LLC at 1-888-724-3274 or FDA at 1-800-FDA-1088 or www.fda.gov/medwatch.

-----DRUG INTERACTIONS------DRUG INTERACTIONS

- P-glycoprotein (P-gp) substrates: Avoid concomitant use of ROMVIMZA with P-gp substrates. If concomitant use cannot be avoided, take ROMVIMZA at least 4 hours prior to P-gp substrates. Concomitant use of vimseltinib with P-gp substrates may increase exposure of these substrates. (2.3, 7.1)
- Breast Cancer Resistance Protein (BCRP) substrates: Avoid concomitant use of ROMVIMZA with BCRP substrates. Concomitant use of vimseltinib with BCRP substrates may increase exposure of these substrates. (7.1)
- Organic Cation Transporter 2 (OCT) substrates: Avoid concomitant use of ROMVIMZA with OCT2 substrates. Concomitant use of vimseltinib with OCT2 substrates may increase exposure of these substrates. (7.1)

-----USE IN SPECIFIC POPULATIONS-----

Lactation: Advise not to breastfeed. (8.2)

See 17 for PATIENT COUNSELING INFORMATION and Medication Guide

Revised: 02/2025

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FULL PRESCRIBING INFORMATION

1 INDICATIONS AND USAGE

ROMVIMZA is indicated for treatment of adult patients with symptomatic tenosynovial giant cell tumor (TGCT) for which surgical resection will potentially cause worsening functional limitation or severe morbidity.

2 DOSAGE AND ADMINISTRATION

2.1 Recommended Dosage

The recommended dosage of ROMVIMZA is 30 mg orally taken twice weekly, with a minimum of 72 hours between doses, as directed on the blister package [see Clinical Pharmacology (12.3)]. Instruct patients to follow the schedule on the blister package and to take ROMVIMZA on the same days each week.

- ROMVIMZA may be taken with or without food.
- Swallow ROMVIMZA capsules whole. Do not open, break, or chew the capsules.
- If a dose is missed by 48 hours or less, take the missed dose as soon as possible and take the next dose on its regularly scheduled day. If a dose is missed by more than 48 hours, skip the missed dose, and take the next dose on its regularly scheduled day.
- If vomiting occurs within 30 minutes of taking a dose, repeat that dose. Otherwise, take the next dose on its regularly scheduled day.

2.2 Dose Modifications for Adverse Reactions

The recommended dose reductions for adverse reactions are provided in Table 1.

Table 1: Recommended Dose Reductions

Dose Reduction	Twice Weekly Dose
First	20 mg
Second	14 mg

Permanently discontinue ROMVIMZA in patients who are unable to tolerate 14 mg orally twice weekly.

The recommended dosage modifications for hepatotoxicity are summarized in Table 2.

Table 2: Recommended Dosage Modifications for Hepatotoxicity

Hepatotoxicity Severity	ROMVIMZA Dosage Modifications
AST and/or ALT increases >3–5 times ULN and total bilirubin increases up to 2 times ULN	Withhold ROMVIMZA until AST and ALT resolve to baseline or ≤3 times ULN, and bilirubin resolves to baseline.
OR	Resume at the next lower dose level once Hy's law has been definitively ruled out.
Total bilirubin increases up to 2 times ULN	Permanently discontinue if adverse reaction does not resolve within 4 weeks.

Hepatotoxicity Severity	ROMVIMZA Dosage Modifications
AST and/or ALT increases >3–5 times ULN, and total bilirubin increases >2 times ULN or INR >1.5 and ALP <2 times ULN OR	Withhold ROMVIMZA until AST and ALT resolve to baseline or ≤3 times ULN, and bilirubin resolves to baseline. Resume at the next lower dose level once Hy's law has been definitively ruled out.
Total bilirubin increases >2 times ULN	Permanently discontinue if adverse reaction does not resolve within 4 weeks.
AST and/or ALT increases >5–8 times ULN, and total bilirubin ≤ULN and without clinical symptoms	Withhold ROMVIMZA until AST and ALT resolve to ≤3 times ULN or baseline. Permanently discontinue if adverse reaction does not resolve within 4 weeks.
AST and/or ALT increases >5-8 times ULN and total bilirubin increase >ULN, or INR >1.5, or ALP >2 times ULN	Permanently discontinue ROMVIMZA.
AST and/or ALT increases >8 times ULN	Permanently discontinue ROMVIMZA.

ALT = alanine aminotransferase; ALP = alkaline phosphatase; AST = aspartate aminotransferase; INR = International normalized ratio; ULN = upper limit of normal

2.3 Dosage Modification for P-glycoprotein (P-gp) Substrates

Avoid concomitant use of ROMVIMZA with P-gp substrates. If concomitant use of a P-gp substrate is unavoidable, administer ROMVIMZA at least 4 hours before taking the P-gp substrate unless otherwise recommended in the substrate Prescribing Information [see Drug Interactions (7.1)].

3 DOSAGE FORMS AND STRENGTHS

14 mg capsule

Orange cap, white body size 4 capsule imprinted with "DCV14" in black ink.

20 mg capsule

Yellow cap, white body size 2 capsule imprinted with "DCV20" in black ink.

30 mg capsule

Light blue cap, white body size 1 capsule imprinted with "DCV30" in black ink.

4 CONTRAINDICATIONS

None.

5 WARNINGS AND PRECAUTIONS

5.1 Hepatotoxicity

Cases of serious and fatal liver injury have occurred with the use of another kinase inhibitor that targets CSF1R [see Clinical Pharmacology (12.1)]. Serious and fatal liver injury have not been observed with ROMVIMZA.

Across clinical trials in 253 patients treated with ROMVIMZA, 2% had Grade 3 increased AST, and 1% had Grade 3 increased ALT. Dose interruptions occurred in 2% of patients and dose reductions occurred in 1% of patients due to AST/ALT increase. One patient discontinued therapy due to Grade 3 AST increased.

Avoid ROMVIMZA in patients with pre-existing increased serum transaminases; total bilirubin or direct bilirubin (>ULN); or active liver or biliary tract disease, including ALP. Monitor liver tests, including AST, ALT, total bilirubin, direct bilirubin, ALP and gamma-glutamyl transferase (GGT), prior to initiation of ROMVIMZA, twice a month for the first two months and once every 3 months for the first year of therapy and as clinically indicated thereafter. Withhold and reduce the dose, or permanently discontinue ROMVIMZA based on the severity of the hepatotoxicity [see Dosage and Administration (2.2)].

5.2 Embryo-Fetal Toxicity

Based on data from animal studies and its mechanism of action, ROMVIMZA can cause fetal harm when administered to pregnant women. In female rats administered vimseltinib, fetal structural abnormalities occurred at exposures that were at least 3 times the recommended dose based on area under the curve (AUC). Advise pregnant women on the potential risk to the fetus.

Advise females of reproductive potential and males with female partners of reproductive potential to use effective contraception during treatment with ROMVIMZA and for 1 month after the last dose [see Use in Specific Populations (8.1, 8.3)].

5.3 Allergic Reactions to FD&C Yellow No.5 (Tartrazine) and No. 6 (Sunset Yellow FCF)

ROMVIMZA 20 mg capsule contains FD&C Yellow No. 5 (tartrazine) which may cause allergic reactions (including bronchial asthma) in certain susceptible patients. Although the overall incidence of FD&C Yellow No. 5 (tartrazine) sensitivity in the general population is low, it is frequently seen in patients who also have aspirin sensitivity.

ROMVIMZA 14 mg and 20 mg capsules contain FD&C Yellow No.6 (Sunset Yellow FCF), which may cause allergic reactions.

5.4 Increased Creatinine without Affecting Renal Function

In MOTION, serum creatinine increased (mean increase of 19 µmol/L) and reached a maximum mean increase by 10.4 weeks compared to baseline. These increases in serum creatinine may not be associated with changes in renal function. Increases in creatinine reversed upon ROMVIMZA discontinuation. The increases in serum creatinine may be due to inhibition of renal tubular secretion transporters [see *Drug Interactions (7.1) and Clinical Pharmacology (12.3)*]. During ROMVIMZA treatment, use alternative measures that are not based on serum creatinine to assess renal function.

6 ADVERSE REACTIONS

The following clinically significant adverse reactions are described elsewhere in the labeling:

Hepatotoxicity [see Warnings and Precautions (5.1)]

6.1 Clinical Trials Experience

Because clinical trials are conducted under widely varying conditions, adverse reaction rates observed in the clinical trials of a drug cannot be directly compared to rates in the clinical trials of another drug and may not reflect the rates observed in practice.

The pooled safety population described in the Warnings and Precautions reflects exposure to ROMVIMZA in 83 patients with TGCT enrolled in the double-blind portion and in 35 patients with TGCT in the open-label portion who crossed over to ROMVIMZA in MOTION, and in 135 patients with TGCT or solid tumors in other clinical trials.

The safety of ROMVIMZA was evaluated in 83 adult patients with TGCT in MOTION [see Clinical Studies (14)]. MOTION excluded patients with bilirubin, AST, or ALT >ULN. All patients received ROMVIMZA twice weekly until disease progression or unacceptable toxicity. Among these patients, 82% were exposed for 6 months or longer and 30% were exposed for greater than one year.

Serious adverse reactions occurred in 2.4% of patients who received ROMVIMZA. Serious adverse reactions in ≥1% included subcutaneous abscess (1.2%) and cellulitis (1.2%).

Permanent discontinuation due to an adverse reaction occurred in 4.8% of patients who received ROMVIMZA. Adverse reactions leading to permanent discontinuation in one patient each included periorbital edema, neuropathy, rash, and hypertension.

Dose reductions due to an adverse reaction or laboratory abnormality occurred in 39% of patients who received ROMVIMZA. Adverse reactions leading to dose reductions in ≥2% of patients receiving ROMVIMZA were rash, periorbital edema, peripheral edema, fatigue, pruritus, face edema, increased CPK, neuropathy, and hypertension.

Dose interruptions due to an adverse reaction or laboratory abnormality occurred in 40% of patients who received ROMVIMZA. Adverse reactions leading to interruptions in ≥2% of patients included rash, fatigue, peripheral edema, increased CPK, periorbital edema, face edema, pruritus, neuropathy, and hypertension.

The most common (≥20%) adverse reactions, including laboratory abnormalities that occurred in patients receiving ROMVIMZA were increased AST, periorbital edema, fatigue, rash, increased cholesterol, peripheral edema, face edema, decreased neutrophils, decreased leukocytes, pruritus, and increased ALT. Table 3 and Table 4 summarize the adverse reactions and laboratory abnormalities in MOTION during the randomized phase through Week 25.

Table 3: Adverse Reactions Occurring in ≥10% of Patients Receiving ROMVIMZA with a Difference Between Arms of >5% Compared to Placebo Through Week 25 in MOTION

Adverse Beastian*	ROMVIMZA N=83		Placebo N=39		
Adverse Reaction* All Grade (%)		Grade 3 or 4 (%)	All Grades (%)	Grade 3 or 4 (%)	
Eye disorders					
Periorbital edema ¹	60	3.6	21	0	
Lacrimation increased	12	0	0	0	
Dry eye ¹	10	0	0	0	

Advance Departions	ROMVIMZA N=83		Placebo N=39	
Adverse Reaction* -	All Grades (%)	Grade 3 or 4 (%)	All Grades (%)	Grade 3 or 4 (%)
General disorders and ac	lministration site c	onditions	.	. ,
Fatigue ¹	59	1.2	38	2.6
Peripheral edema ¹	33	1.2	8	0
Face edema	31	1.2	8	0
Skin and subcutaneous t	issue disorders			
Rash ¹	47	3.6	5	0
Pruritus	29	2.4	8	0
Vascular disorders		- 1		1
Hypertension	17	4.8	10	2.6
Nervous system disorder	rs	<u> </u>		•
Neuropathy ¹	12	1.2	2.6	0

¹ Includes multiple related terms

Other clinically significant adverse reactions occurring in <10% of patients treated with ROMVIMZA include blurred vision (6%).

Table 4: Laboratory Abnormalities Worsening from Baseline in ≥10% of Patients Receiving ROMVIMZA with a Difference Between Arms of >5% Compared to Placebo Through Week 25 in MOTION

l ale a veta ve Ale ve a veta alite d	ROMVIMZA N=83		Placebo N=39 ^b		
Laboratory Abnormality ^a	All Grades (%)	Grade 3 or 4 (%)	All Grades (%)	Grade 3 or 4 (%)	
Chemistry					
AST increased	92	0	11	0	
Cholesterol increased	43	0	16	0	
ALT increased	24	0	16	0	
Creatinine increased	17	0	2.6	0	
ALP increased	14	0	8	0	
Magnesium increased	13	1.2	2.6	0	
Calcium decreased	13	0	2.6	0	
Hematology					
Neutrophils decreased	31	1.2	2.6	0	
Leukocytes decreased	29	0	8	0	

AST – aspartate aminotransferase, ALT – alanine aminotransferase, ALP – alkaline phosphatase

Additional clinically significant laboratory abnormality: *Increased Creatine Phosphokinase (CPK)*

^a The severity of adverse reactions was assessed using CTCAE v5.0.

^b The denominator used to calculate the rate was 83 for ROMVIMZA and 38 for placebo based on the number of patients with a baseline value and at least one post-treatment value.

7 DRUG INTERACTIONS

7.1 Effects of ROMVIMZA on Other Drugs

Table 5 describes drug interactions where concomitant use with ROMVIMZA affects another drug.

Table 5: Effect of ROMVIMZA on Other Drugs

P-glycoprotein (P-gp) substrates		
Prevention or Management	Avoid concomitant use with P-gp substrates while taking ROMVIMZA. If concomitant use cannot be avoided, take ROMVIMZA at least 4 hours prior to P-gp substrates [see Dosage and Administration (2.3)] unless otherwise recommended in the substrate Prescribing Information.	
Mechanism and Clinical Effect(s)	This recommendation is based upon a mechanistic understanding of vimseltinib pharmacokinetics and it being a P-gp inhibitor in vitro [see Clinical Pharmacology (12.3)]. Concomitant use of ROMVIMZA with P-gp substrates may increase exposure of these substrates; however, this has not been studied clinically.	
Breast Cancer Resis	tance Protein (BCRP) substrates	
Prevention or Management	Avoid concomitant use with BCRP substrates while taking ROMVIMZA. Refer to the Prescribing Information of the BCRP substrate for dose modifications if concomitant use cannot be avoided.	
Mechanism and Clinical Effect(s)	This recommendation is based upon a mechanistic understanding of vimseltinib pharmacokinetics and it being a BCRP inhibitor in vitro [see Clinical Pharmacology (12.3)]. Concomitant use of ROMVIMZA with BCRP substrates may increase exposure of these substrates; however, this has not been studied clinically.	
Organic Cation Trans	sporter 2 (OCT2) substrates	
Prevention or Management	Avoid concomitant use with OCT2 substrates while taking ROMVIMZA. Refer to the Prescribing Information of the OCT2 substrate for dose modifications if concomitant use cannot be avoided.	
Mechanism and Clinical Effect(s)	This recommendation is based upon a mechanistic understanding of vimseltinib pharmacokinetics and it being an OCT2 inhibitor in vitro [see Clinical Pharmacology (12.3)]. Concomitant use of ROMVIMZA with OCT2 substrates may increase exposure of these substrates; however, this has not been studied clinically.	

8 USE IN SPECIFIC POPULATIONS

8.1 Pregnancy

Risk Summary

Based on data from animal studies and its mechanism of action, ROMVIMZA can cause fetal harm when administered to a pregnant woman. There are no available data on vimseltinib use in pregnant women to evaluate for a drug-associated risk of major birth defects, miscarriage, or adverse maternal or fetal outcomes. In female rats administered vimseltinib during the period of

organogenesis, fetal structural abnormalities occurred at exposures that were at least 3 times the recommended dose based on AUC (*see Data*). Advise pregnant women of the potential risk to a fetus.

The background risk of major birth defects and miscarriage for the indicated population is unknown. All pregnancies have a background risk of birth defect, loss, or other adverse outcomes. In the U.S. general population, the estimated background risk of major birth defects and miscarriage in clinically recognized pregnancies is 2 to 4% and 15 to 20%, respectively.

Data

Animal Data

In a rat embryo-fetal development study, pregnant female rats were dosed once daily during the period of organogenesis (gestational days 6 to 17) at doses of 2.5, 5, or 15 mg/kg/day. Structural abnormalities (skeletal variations) occurred at ≥2.5 mg/kg/day (approximately 3 times the exposure at the recommended dose based on AUC). Additional structural abnormalities (cardiac malformations) were observed at the highest dose of 15 mg/kg/day (approximately 23 times the exposure at the recommended dose based on AUC).

8.2 Lactation

Risk Summary

There are no data on the presence of vimseltinib or its metabolites in either human or animal milk or its effects on a breastfed child or on milk production. Because of the potential for serious adverse reactions in the breastfed child, advise women not to breastfeed during treatment with ROMVIMZA and for 1 month after the last dose.

8.3 Females and Males of Reproductive Potential

ROMVIMZA can cause fetal harm when administered to a pregnant woman [see Use in Specific Populations (8.1)].

Pregnancy Testing

Verify pregnancy status in females of reproductive potential prior to the initiation of ROMVIMZA [see Use in Specific Populations (8.1)].

Contraception

Females

Advise females of reproductive potential to use effective contraception during treatment with ROMVIMZA and for 1 month after the last dose [see Warnings and Precautions (5.2), Nonclinical Toxicology (13.1)].

Males

Advise males that are partnered with females of reproductive potential to use effective contraception during treatment with ROMVIMZA and for 1 month after the last dose [see Warnings and Precautions (5.2), Nonclinical Toxicology (13.1)].

Infertility

Females and Males

Based on findings from animal studies, ROMVIMZA may impair fertility [see Nonclinical Toxicology (13.1)].

8.4 Pediatric Use

The safety and effectiveness of ROMVIMZA in pediatric patients have not been established.

Animal Toxicity Data

In a 26-week repeat-dose toxicology study, rats administered vimseltinib at ≥2.5 mg/kg/day had physeal thickening and decay of the incisors and molars. Bone and tooth toxicities occurred at exposures at least 8 times the recommended dose based on AUC.

8.5 Geriatric Use

Clinical studies of ROMVIMZA did not include a sufficient number of patients aged 65 years and older to determine whether they respond differently from younger patients.

8.6 Hepatic Impairment

No dose adjustment is recommended for patients with mild (bilirubin ≤upper limit of normal (ULN) and aspartate aminotransferase (AST) >ULN or bilirubin >1x to 1.5x ULN and any AST) hepatic impairment. ROMVIMZA has not been studied in patients with moderate (bilirubin >1.5x to 3x ULN and any AST) or severe (bilirubin >3x ULN and any AST) hepatic impairment [see Clinical Pharmacology (12.3)].

11 DESCRIPTION

Vimseltinib is a kinase inhibitor. The chemical name of vimseltinib dihydrate is 3-methyl-5-[6-methyl-5-[2-(1-methylpyrazol-4-yl)pyridin-4-yl]oxypyridin-2-yl]-2-(propan-2-ylamino)pyrimidin-4-one, dihydrate.

Vimseltinib is a white to off-white crystalline solid. Vimseltinib is a weak base, very slightly soluble in water. The molecular formula for vimseltinib dihydrate is C₂₃H₂₅N₇O₂ • 2 H₂O, and the molecular weight is 467.52 g/mol. The chemical structure is:

$$\begin{array}{c|c}
 & O & \\
 & N & \\
 &$$

ROMVIMZA (vimseltinib) capsules are supplied as printed hard gelatin capsules containing 14 mg, 20 mg, or 30 mg of vimseltinib (equivalent to 15.18 mg, 21.68 mg, or 32.52 mg of vimseltinib dihydrate, respectively). The capsule contains the following inactive ingredients:

crospovidone, lactose monohydrate, and magnesium stearate. The capsule shell contains Brilliant Blue FCF (30 mg strength), erythrosine (30 mg strength), gelatin, Sunset Yellow FCF (14 mg and 20 mg strengths), tartrazine (20 mg), and titanium dioxide.

12 CLINICAL PHARMACOLOGY

12.1 Mechanism of Action

Vimseltinib is a kinase inhibitor that inhibits colony-stimulating factor 1 receptor (CSF1R). In vitro, vimseltinib inhibited CSF1R autophosphorylation, signaling induced by CSF1 ligand binding, and proliferation of cells expressing CSF1R.

12.2 Pharmacodynamics

Exposure-Response Relationship

Higher vimseltinib exposure is associated with an increased risk of all grades of edema, rash, increased AST, and increased ALT.

Vimseltinib exposure-response relationship for efficacy and time course of pharmacodynamic response have not been fully characterized.

Cardiac Electrophysiology

At the maximum recommended dose of ROMVIMZA, clinically significant QTc interval prolongation was not observed. However, the largest mean increase in QTc interval was 8.2 ms (upper confidence internal = 12.3 ms) after administration of vimseltinib 40 mg once daily for 5 days (3.3 times the maximum recommended weekly dose). The increase in QTc interval was concentration-dependent [see Clinical Pharmacology (12.3)].

12.3 Pharmacokinetics

Vimseltinib pharmacokinetic parameters were determined following a single oral dose of 30 mg or at steady state following multiple doses of 30 mg twice weekly and are provided as mean (CV%) unless otherwise specified.

Vimseltinib peak plasma concentration (C_{max}) is 283 ng/mL (36%) or 747 ng/mL (39%) after a single dose or at steady state, respectively, and area under the time concentration curve (AUC_{0-inf}) is 46,900 ng•h/mL (45%) after a single dose and AUC_{0-24hr} is 13,400 ng•h/mL (45%) at steady state.

Vimseltinib pharmacokinetics are dose-proportional.

Absorption

Vimseltinib median time to C_{max} (T_{max}) is 1 hour (0.5 to 4 hours).

Effect of Food

No clinically significant differences in vimseltinib pharmacokinetics were observed following administration of a high-fat meal (800 to 1000 kcal, 50% fat), compared to fasted conditions.

Distribution

Vimseltinib volume of distribution (V/F) is 90 L (16%). Vimseltinib is 96.5% bound to human plasma proteins.

Elimination

Vimseltinib elimination half-life ($t_{1/2}$) is approximately 6 days (32%) with a clearance (CL/F) of 0.5 L/h (23%).

Metabolism

Vimseltinib is primarily metabolized by oxidation, *N*-demethylation, and *N*-dealkylation; secondary biotransformation pathways included *N*-demethylation, dehydrogenation, and oxidation. CYP450 enzymes are not anticipated to play a major role in the metabolism of vimseltinib.

Excretion

Approximately 43% of the dose was recovered in feces (9.1% unchanged) and 38% in urine (5.1% unchanged) after a single radiolabeled dose.

Specific Populations

No clinically significant differences in the pharmacokinetics of vimseltinib were observed based on age (20 to 91 years), sex, race (Asian, Black or African American, White), body weight (43 to 150 kg), tumor (TGCT or other malignant solid tumors), and mild to moderate renal impairment (estimated glomerular filtration rate [eGFR] \geq 30 mL/min calculated by CKD-EPI equation). The effect of severe renal impairment (eGFR <30 mL/min) or moderate to severe hepatic impairment (total bilirubin >1.5 x ULN with any AST) on vimseltinib pharmacokinetics is unknown.

Drug Interaction Studies

Clinical Studies and Model-Informed Approaches

<u>P-glycoprotein (P-gp) inhibitors:</u> Dabigatran (a P-gp substrate) AUC_{0-inf} and C_{max} are predicted to increase 2- to 3-fold with concomitant use with vimseltinib 30 mg twice weekly.

Dabigatran C_{max} and AUC_{0-inf} are predicted to increase up to 1.3-fold if administered 4 hours after administration of vimseltinib 30 mg twice weekly.

<u>Other Drugs:</u> No clinically significant differences in vimseltinib pharmacokinetics were observed when used concomitantly with itraconazole (a P-gp inhibitor) or rabeprazole (a proton pump inhibitor).

In Vitro Studies

<u>CYP 450 enzymes:</u> Vimseltinib is not a substrate of CYP1A2, CYP2B6, CYP2C8, CYP2C9, CYP2C19, CYP2D6, and CYP3A.

Vimseltinib is not an inhibitor of CYP1A2, CYP2B6, CYP2C8, CYP2C9, CYP2C19, CYP2D6, and CYP3A4.

Vimseltinib is not an inducer of CYP1A2, CYP2B6, or CYP3A4.

<u>Transporter systems:</u> Vimseltinib is a P-gp substrate but is not a substrate of BCRP, BSEP, OATP1B1, OATP1B3, OAT1, OAT3, OCT2, MATE1, and MATE2-K.

Vimseltinib inhibits P-gp, BCRP, BSEP, OATP1B1, OATP1B3, OCT2, BSEP, MATE1, and MATE2-K. Vimseltinib does not inhibit OAT1 and OAT3. Vimseltinib may increase serum creatinine by decreasing renal tubular secretion of creatinine; this may occur due to inhibition of renal transporters OCT2 and MATE1 and may not affect renal function.

13 NONCLINICAL TOXICOLOGY

13.1 Carcinogenesis, Mutagenesis, Impairment of Fertility

In a 6-month transgenic mouse carcinogenicity study at doses up to 12.5 mg/kg/day, vimseltinib was negative for carcinogenic findings.

Vimseltinib was not mutagenic in the bacterial reverse mutation assay (Ames). In an in vitro micronucleus assay, vimseltinib increased micronuclei after a 24 -hour incubation in the absence of metabolic activation. In vivo, vimseltinib administered to rats at doses up to 200 mg/kg/day did not increase bone marrow micronucleated polychromatic erythrocytes, nor did vimseltinib increase liver DNA strand breaks.

In a fertility and early embryonic development study, male rats were administered 1, 2.5, or 5 mg/kg/day of vimseltinib starting 10 weeks before cohabitation, during cohabitation with untreated females, and at least 2 weeks post-cohabitation. Lower epididymal and testes weights were observed at 5 mg/kg/day (approximately 12 times the exposure at the recommended dose based on AUC). There were no treatment-related effects on mating, fertility, or sperm parameters at any dose tested. Female rats were administered 2.5, 5, or 10 mg/kg/day of vimseltinib 2 weeks prior to cohabitation with untreated males and during cohabitation until gestational day 7. Although there were no treatment-related effects on mating or estrous cycles, vimseltinib administered daily resulted in post-implantation loss at 10 mg/kg/day in female rats (approximately 20 times the exposure at the recommended dose based on AUC).

In a 26-week repeat-dose general toxicology study, recovery male rats that were administered 2.5 or 5 mg/kg/day had moderate to marked reductions in sperm and marked testicular atrophy (1 of 5 and 2 of 5 animals, respectively) corresponding to approximately 6 and 12 times the exposure at the recommended dose based on AUC, respectively. In a 39-week repeat-dose general toxicology study, minimal to moderate epididymal mineralization occurred in male dogs administered ≥4 mg/kg/day corresponding to exposures lower than the exposure at the recommended dose based on AUC.

13.2 Animal Toxicology and/or Pharmacology

In a 26-week repeat-dose general toxicology study in rats, chronic progressive nephropathy occurred at vimseltinib doses of ≥2.5 mg/kg/day (approximately 6 times the exposure at the recommended dose based on AUC). Degeneration of blood vessels (perivascular inflammation and necrosis of arteries and arteriole walls) occurred in multiple tissues at 5 mg/kg/day (approximately 12 times the exposure at the recommended dose based on AUC).

In a 39-week repeat-dose general toxicology study in dogs, skin depigmentation in the head and legs occurred at vimseltinib doses of ≥4 mg/kg/day corresponding to exposures lower than the exposure at the recommended dose based on AUC.

14 CLINICAL STUDIES

The efficacy of ROMVIMZA was evaluated in MOTION (NCT05059262), a phase 3, double-blind, multicenter, randomized (2:1), placebo-controlled study in patients with TGCT for whom surgical resection may cause worsening functional limitation or severe morbidity. Eligible patients had a confirmed diagnosis of TGCT with measurable disease per the Response Evaluation Criteria in Solid Tumors (RECIST v1.1) with at least one lesion having a minimum size of 2 cm. Patients were randomized to placebo or ROMVIMZA 30 mg twice weekly for 24 weeks. Randomization was stratified by tumor location (lower limb versus all other) and region (United States [US] versus non-US). At Week 25, patients who completed the double-blind, randomized part of the trial were eligible to advance to an ongoing, open-label extension study in which all patients received ROMVIMZA.

The major efficacy outcome measure was overall response rate (ORR) as assessed by blinded independent radiological review (IRR) per RECIST v1.1 at Week 25. Additional efficacy outcomes measured at Week 25 included ORR as assessed using tumor volume score (TVS), mean change from baseline in active range of motion of the affected joint at Week 25 measured by goniometry assessments, change from baseline in the Patient-Reported Outcomes Measurement Information System-Physical Function (PROMIS-PF) 15-item score (upper and lower extremity items), and response of at least a 30% improvement in the mean Brief Pain Inventory (BPI) Worst Pain numeric rating scale (NRS) score without a 30% or greater increase in narcotic analgesic use.

A total of 123 patients were randomized: 83 to ROMVIMZA and 40 to placebo during the double-blind period of the study. The median age was 44 years (range 20 to 78 years); 59% of patients were female; 65% were White, 4% were Asian, 3% were Black or African American, and 28% were not reported or unknown; 69% were not Hispanic or Latino, 3% were Hispanic or Latino, and 28% were not reported or unknown; 74% of patients had prior surgery; 69% of patients had diffuse TGCT; and 23% of patients were previously treated with systemic therapy. Disease locations were knee (67%), ankle (12%), hip (10%), other (5%), foot (3.3%), and wrist (2.4%).

A statistically significant improvement in ORR was demonstrated in patients randomized to ROMVIMZA compared with placebo. Efficacy results in MOTION are summarized in Table 6.

Table 6: Efficacy Results Assessed at Week 25 for MOTION

Efficacy Parameter	ROMVIMZA N = 83	Placebo N = 40		
Overall Response Rate per RECIST v1.1 (95% CI)	40% (29%, 51%)	0% (0%, 9%)		
Complete Response	5%	0%		
Partial Response	35%	0%		
p-value	<0.0	<0.0001		
Duration of Response ¹				
Median (Range in months) ²	NR (2.5+, 19.4+)	N/A		
DOR ≥6 months	28 (85%)	-		

DOR ≥9 months	19 (58%)	-
Active ROM ³		
Patients with data at baseline and Week 25, n	73	33
Baseline mean (SD)	63.0 (29.4)	62.9 (32.2)
Mean change from baseline ⁴	10 / (5 6 21 2)	20/105 100
(95% CI)	18.4 (5.6, 31.2)	3.8 (-10.5, 18.0)
Difference in LS means (95% CI)	14.6% (4	.0, 25.3)
p-value	0.0077	
PROMIS-PF (15-Item score; ranges from 0-100) ⁵		
Patients with data at baseline and Week 25, n	63	30
Baseline mean (SD)	39.0 (6.1)	38.5 (6.0)
Mean change from baseline ⁴	4 0 (0 = 0 =)	
(95% CI)	4.6 (2.7, 6.5)	1.3 (-0.5, 3.0)
Difference in LS means (95% CI)	3.3 (1.4, 5.2)	
p-value	0.0007	
BPI-30 Response ⁶		
Patients with data at baseline and Week 25, n	68	31
Responders, n (%)	40 (48.2%)	9 (22.5%)
(95% CI)	(37.1%, 59.4%)	(10.8%, 38.5%)
Difference in responder rate (95% CI) ⁷	26.2% (9.5, 42.8)	
p-value	0.0056	

NR=Not reached; N/A=Not applicable; AMA=American Medical Association; BPI=Brief Pain Inventory; CI=confidence interval; LS=least squares; MMRM=mixed model for repeated measures; n=number of patients in the category; N=sample size; PROMIS-PF=Patient-reported Outcomes Measurement Information System-Physical Function; ROM=range of motion; SD= Standard deviation.

ORR by TVS was 67% (95% CI: 56, 77) in patients randomized to ROMVIMZA and 0% in patients randomized to placebo; p-value <0.0001.

Individual patient results of change from baseline in active ROM at Week 25 (ROMVIMZA N = 73; placebo N=33) are shown in Figure 1.

¹ DOR results are based on an additional 6 months of follow-up from the time of ORR analysis.

² The median DOR was estimated using the Kaplan-Meier method. "+" indicates that the patient's response was ongoing at last assessment as of the data cutoff date.

³ Active ROM was normalized to the AMA reference standard.

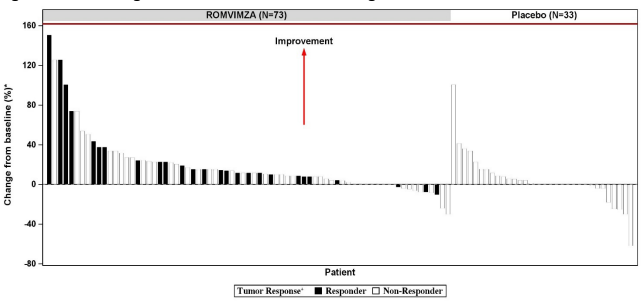
⁴ Mean change from baseline was estimated from the MMRM for each corresponding endpoint. Baseline means presented include all patients and not only the ones with data at baseline and Week 25.

⁵ Data for PROMIS-PF is largely based on lower limb extremity assessment due to tumor location as described above. Higher scores of PROMIS-PF indicate better physical functioning.

⁶ BPI response in Worst Pain is defined as at least a 30% improvement in the mean BPI Worst Pain NRS score (0-10 NRS) without a 30% or greater increase in narcotic analgesic use at Week 25.

⁷95% CI for the difference in response rates based on the stratified Mantel-Haenszel method.

Figure 1: Change from Baseline in Active Range of Motion at Week 25 for MOTION



^{*}Percent normal reference range for the affected joint.

16 HOW SUPPLIED/STORAGE AND HANDLING

Table 7: ROMVIMZA 14 mg, 20 mg, and 30 mg Capsules

Strength	Description	Package Size and Type	NDC Number
14 mg	Size 4 hard gelatin capsule with white body and orange cap with black print "DCV14", packed in oPA-film/ aluminum foil /PVC-film blisters with push-through aluminum foil lidding.	Each carton contains one child-resistant blister pack containing 8 capsules (four-week supply)	73207-302-40
20 mg	Size 2 hard gelatin capsule with white body and yellow cap with black print "DCV20", packed in oPA-film/ aluminum foil /PVC-film blisters with push-through aluminum foil lidding.	Each carton contains one child-resistant blister pack containing 8 capsules (four-week supply)	73207-303-40
30 mg	Size 1 hard gelatin capsule with white body and light blue cap with black print "DCV30", packed in oPA-film/ aluminum foil /PVC-film blisters with push-through aluminum foil lidding.	Each carton contains one child-resistant blister pack containing 8 capsules (four-week supply)	73207-304-40

Store at controlled room temperature 20°C to 25°C (68°F to 77°F); excursions permitted to 15°C to 30°C (59°F to 86°F) [See USP Controlled Room Temperature].

⁺ Response by RECIST v1.1.

Store capsules in their original blister packs until ready to be taken. Do not store ROMVIMZA in another container.

17 PATIENT COUNSELING INFORMATION

Advise the patient to read the FDA approved patient labeling (Medication Guide).

Hepatotoxicity

Advise patients there may be a potential risk of hepatotoxicity and that they will need to undergo laboratory tests to monitor liver function and to immediately report any signs or symptoms of severe liver injury to their healthcare provider [see Warnings and Precautions (5.1)].

Embryo-Fetal Toxicity

- Advise pregnant women and females of reproductive potential of the potential risk to the
 fetus. Advise females of reproductive potential to inform their healthcare provider of a
 known or suspected pregnancy [see Warnings and Precautions (5.2), Use in Specific
 Populations (8.1, 8.3)].
- Advise females of reproductive potential to avoid pregnancy and to use effective contraception during treatment with ROMVIMZA and for 1 month after the last dose [see Warnings and Precautions (5.2), Use in Specific Populations (8.1, 8.3)].
- Advise males with female partners of reproductive potential to use effective contraception during treatment and for 1 month after the last dose of ROMVIMZA [see Warnings and Precautions (5.2) and Use in Specific Populations (8.3)].

Lactation

Advise females not to breastfeed during treatment with ROMVIMZA and for 1 month after the final dose [see Use in Specific Populations (8.2)].

Infertility

Advise patients that ROMVIMZA may impair fertility [see Use in Specific Populations (8.3), Nonclinical Toxicology (13.1)].

Allergic Reactions to FD&C Yellow No. 5 (Tartrazine) and No. 6 (Sunset Yellow FCF)

Advise patients that ROMVIMZA 20 mg contains FD&C Yellow No. 5 (tartrazine), which may cause allergic-type reactions (including bronchial asthma) in certain susceptible persons or in patients who also have aspirin hypersensitivity [see Warnings and Precautions (5.3)]. Advise patients ROMVIMZA 14 mg and ROMVIMZA 20 mg contains FD&C Yellow No. 6 (Sunset Yellow FCF) which may cause allergic-type reactions [see Warnings and Precautions (5.3)].

Administration

- Instruct patients that doses should be taken twice weekly at least 72 hours apart [see Dosage and Administration (2.1)].
- Instruct patients to swallow capsules whole (do not open, break, or chew) [see *Dosage* and Administration (2.1)].

Drug Interactions

Advise patients to inform their healthcare providers of all concomitant products, including over-the-counter products and supplements [see Dosage and Administration (2.3), Drug Interactions (7.1)].

Manufactured for and marketed by: **Deciphera Pharmaceuticals, LLC** 200 Smith Street, Waltham, MA 02451