

Capivasertib (Truqap®) Patient Management

Description: This document will help in the identification and management of patients taking capivasertib.

Background: Capivasertib is an AKT kinase inhibitor that is indicated in adult patients with:1

- Hormone receptor- positive, HER2-negative locally advanced or metastatic breast cancer with one or more PIK3CA/AKT1/PTEN-alterations
 - Administered in combination with fulvestrant
 - o Following progression on ≥ 1 endocrine-based regimen in the metastatic setting or recurrence on or within ≤ 12 months of completing adjuvant therapy

Most common adverse reactions (≥ 20%):1

- Increased: random glucose, fasting glucose, triglycerides, creatinine
- Decreased: lymphocytes, hemoglobin, leukocytes, neutrophils
- Other: diarrhea, cutaneous adverse reactions, nausea, vomiting, fatigue, stomatitis

PQI Process:2

- For pre- and peri-menopausal patients, a luteinizing hormone-releasing hormone (LHRH) agonist (according to current clinical practice standards) should be administered; for males, consider administering an LHRH agonist (according to current clinical practice standards)
- Evaluate fasting blood glucose (FBG), HbA1c and then optimize blood glucose prior to capivasertib initiation
 - Median time to first occurrence of hyperglycemia was 15 days¹
- Table 1. Dosing considerations for Capivasertib²

Dosage form	Tablet, Oral – 160 mg, 200 mg		
	Blister pack – 160 mg, 200 mg (each carton has 4 blister packs (64		
	tabs total) - each blister pack contains 16 tabs)		
Usual starting dose	400 mg twice daily (~12 hours apart) for 4 consecutive days, followed by 3 days off (administer capivasertib on days 1 to 4 of each week); in combination with fulvestrant; continue until disease progression or unacceptable toxicity		
Dose adjustments	Capivasertib has not been studied in patients with severe hepatic or		
(renal/hepatic)	renal impairment		
Dose reductions for	400 mg BID □ 320 mg BID □ 200 mg BID □ permanently discontinue		
toxicity	if unable to tolerate the final dose reduction		

- Once medication delivery is scheduled, ensure complete counseling on administration, proper handling, storage, missed dose management, side effect information, and all other pertinent information
- Assess the patient's understanding of the regimen complexity and provide tools to assist with adherence
- Monitor for signs/symptoms of cutaneous adverse reactions, diarrhea, and hyperglycemia; monitor for adverse reactions in patients with moderate hepatic impairment
- Monitor adherence

IMPORTANT NOTICE: NCODA has developed this Positive Quality Intervention platform. This platform is intended as an educational aid, does not provide individual medical advice, and does not substitute for the advice of a qualified healthcare professional. This platform does not cover all existing information related to the possible uses, directions, doses, precautions, warning, interactions, adverse effects, or risks associated with the medication. The materials contained in this platform do not constitute or imply endorsement, recommendation, or favoring of this medication by NCODA. NCODA does not ensure the accuracy of the information presented and assumes no liability relating to its accuracy. All decisions related to taking this medication should be made with the guidance and under the direction of a qualified healthcare professional. It is the individual's sole responsibility to seek guidance from a qualified healthcare professional. *Updated 05/25/25 PQI-108*

Capivasertib (TruQap®) Patient Management

Written By: Nancy Augustine, PharmD, CSP Institution: Shields Health Solutions

- Monitoring parameters ¹
 - FBG prior to treatment, on day 3 or 4 of the dosing week during weeks 1, 2, 4, 6, and 8; then monthly while on treatment; and as clinically indicated
 - HbA1C prior to treatment and every 3 months of treatment
 - o If hyperglycemia occurs during treatment:
 - Monitor FBG ≥ 2x/week, on days on and off capivasertib, until FBG decreases to baseline.
 - During treatment with anti-diabetic medications, monitor FBG ≥ 1x/week for 2 months, followed by once every 2 weeks, or as clinically indicated
 - If ketoacidosis suspected, hold capivasertib and permanently discontinue if confirmed

Patient-Centered Activities:²

- Administer with or without food, approximately every 12 hours on scheduled days; swallow whole; do not chew, crush, or split tablets
- If a dose is missed within 4 hours of the scheduled time, administer the missed dose; if a dose is missed by more than 4 hours of the scheduled time, skip the dose and administer the next dose at its usual scheduled time
- If a dose is vomited, do not administer an additional dose; administer the next dose at the usual scheduled time
- Avoid grapefruit, star fruit, pomegranate and Seville oranges products
- This medication is considered hazardous counsel on appropriate precautions for handling, administration, and disposal
 - Wash hands before and after handling; caregivers should wear gloves while handling
 - Do not dispose of any medication in trash or flush down sink or toilet contact pharmacist for disposal locations
- Store in the original bottle at room temperature
- Check blood glucose levels more frequently as medication can cause high blood sugar
- Significant drug interactions exist, requiring dose/frequency adjustment or avoidance let healthcare team know of any new medications
- Side effects to monitor
 - Skin changes that include inflammation, redness, rash, hives, itching, discoloration, sun sensitivity
 - Decreased appetite, diarrhea, nausea, vomiting, mouth sores
 - o Signs of urinary tract infection (fever, burning or pain when passing urine, lower stomach, or pelvic pain)
 - Signs of hyperglycemia (confusion, fatigue, flushing, fast breathing, unusual thirst or hunger, urinating more frequently)
 - o Fatigue, headache
- Ensure patient has access to supportive medications such as loperamide, moisturizing cream and antihistamine treatment
- Consider providing a blood glucose meter to the patient
- MyTRUQAP Support Program
 patients can enroll to receive helpful resources, emails, and a starter kit³
- Patient Assistance: NCODA Financial Assistance Tool

References:

- 1. Truqap (capivasertib) Prescribing Information.
- 2. Lexicomp. Capivasertib (Lexi-Drugs).
- 3. Truqap website. https://www.truqap.com/.



Capivasertib (TruQap®) Patient Management

Supplemental Information:²

Capivasertib Dose Reduction Levels			
Dose level	Capivasertib dose and schedule		
Initial (usual) dose	400 mg twice daily for 4 days, followed by 3 days off		
First dose reduction	320 mg twice daily for 4 days, followed by 3 days off		
Second dose reduction	200 mg twice daily for 4 days, followed by 3 days off		
Permanently discontinue if unable to tolerate the second dose reduction.			

Recommended Capiv	Recommended Capivasertib Dosage Modifications				
Adverse reaction	Severity	Capivasertib dosage modification ^a			
Dermatologic toxicity: Cutaneous adverse	Any	Early consultation with a dermatologist is recommended. May require corticosteroids (topical or systemic, depending on the severity) to manage.			
	Grade 2	Withhold capivasertib until recovery to ≤ Grade 1. Resume capivasertib at the same dose.			
		Persistent or recurrent Grade 2 toxicity: Reduce capivasertib by one dose level.			
	Grade 3	Withhold capivasertib until recovery to ≤ Grade 1.			
reactions		Resolution ≤28 days after interruption: Resume capivasertib at the same dose.			
		Resolution >28 days after interruption: Resume capivasertib at one lower dose level.			
		Recurrent Grade 3 toxicity: Permanently discontinue capivasertib.			
	Grade 4	Permanently discontinue capivasertib.			
GI toxicity: Diarrhea	Any	May require anti-diarrheal medications to manage symptoms. Advise patients to increase oral fluids and start antidiarrheal treatment at the first sign of diarrhea.			
	Grade 2	Withhold capivasertib until recovery to ≤ Grade 1.			
		Resolution ≤28 days after interruption: Resume capivasertib at the same or at one lower dose level as clinically indicated.			
		Resolution >28 days after interruption: Resume capivasertib at one lower dose level as clinically indicated.			
		Recurrence: Reduce capivasertib by one dose level.			
		Withhold capivasertib until recovery to ≤ Grade 1.			
	Grade 3	Resolution ≤28 days after interruption: Resume capivasertib at the same or at one lower dose level as clinically indicated.			
		Resolution >28 days after interruption: Permanently discontinue capivasertib.			
	Grade 4	Permanently discontinue capivasertib.			
Hyperglycemia	Any	Consider consultation with a health care practitioner with expertise in hyperglycemia management. Counsel patients on lifestyle modifications.			
	FBG ^b > ULN to 160 mg/dL or	Consider initiation or intensification of oral antidiabetic therapy.			
	FBG > ULN to 8.9 mmol/L or HbA _{1c} >7%				
	FBG 161 to 250 mg/dL or	Withhold capivasertib until FBG decreases to ≤160 mg/dL (or ≤8.9 mmol/L).			



Capivasertib (TruQap®) Patient Management

Recommended Capiva	Recommended Capivasertib Dosage Modifications				
Adverse reaction	Severity	Capivasertib dosage modification ^a			
	FBG 9 to 13.9 mmol/L	Resolution ≤28 days after interruption: Resume capivasertib at the same dose.			
		Resolution >28 days after interruption: Resume capivasertib at one lower dose level.			
	FBG 251 to 500 mg/dL or FBG 14 to 27.8 mmol/L	Withhold capivasertib until FBG decreases to ≤160 mg/dL (or ≤8.9 mmol/L).			
		Resolution ≤28 days after interruption: Resume capivasertib at one lower dose level.			
		Resolution >28 days after interruption: Permanently discontinue capivasertib.			
	FBG >500 mg/dL or	Withold capivasertib.			
		Life-threatening hyperglycemia sequelae or if FBG persists at ≥500 mg/dL after 24 hours: Permanently discontinue capivasertib.			
		If FBG is \leq 500 mg/dL (or \leq 27.8 mmol/L) within 24 hours: Follow the guidance in this table for the relevant grade.			
Other adverse reactions [see Adverse Reactions (6.1) in Package Insert] ¹	Grade 2	Withhold capivasertib until recovery to ≤ Grade 1. Resume capivasertib at the same dose.			
	Grade 3	Withhold capivasertib until recovery to ≤ Grade 1.			
		Resolution ≤28 days after interruption: Resume capivasertib at the same dose.			
		Resolution >28 days after interruption: Resume capivasertib at one lower dose level.			
	Grade 4	Permanently discontinue capivasertib.			

