

## Essai Clinique Généré le 12 mai 2025 à partir de

Titre	A Phase 2 Study of VS-6766 (Dual RAF/MEK Inhibitor) Alone and In Combination With Defactinib (FAK Inhibitor) in Recurrent Low-Grade Serous Ovarian Cancer (LGSOC)
Protocole ID	RAMP-201 (ENGOT-ov60)
ClinicalTrials.gov ID	NCT04625270
Type(s) de cancer	Ovaire
Phase	Phase II
Type étude	Clinique
Médicament	VS-6766 seul ou en association avec defactinib
Institution	CENTRE HOSPITALIER DE L'UNIVERSITE DE MONTREAL
Ville	
Investigateur principal	Dre Diane Provencher
Coordonnateur	Bonny Choy 514-890-8000 poste 24672
Statut	Fermé
But étude	This study will assess the safety and efficacy of VS-6766 monotherapy and in combination with Defactinib in subjects with recurrent Low-Grade Serous Ovarian Cancer (LGSOC)
Critères d'éligibilité	<ul> <li>Histologically proven LGSOC (ovarian, peritoneal)</li> <li>In Part A KRAS mutation, KRAS wt</li> <li>Progression or recurrence of LGSOC after at least one prior systemic therapy for metastatic disease.</li> <li>Measurable disease according to RECIST 1.1</li> <li>An Eastern Cooperative Group (ECOG) performance status ≤ 1.</li> <li>Adequate organ function</li> <li>Adequate recovery from toxicities related to prior treatments</li> <li>Agreement to use highly effective method of contraceptive</li> </ul>
Critères d'exclusion	<ul> <li>Systemic anti-cancer therapy within 4 weeks of the first dose of study therapy</li> <li>Co-existing high-grade ovarian cancer or another histology</li> <li>History of prior malignancy with recurrence &lt;3 years from the time of enrollment</li> <li>Major surgery within 4 weeks</li> <li>Symptomatic brain metastases requiring steroids or other interventions</li> <li>Known SARS-Cov2 infection (clinical symptoms) ≤28 days prior to first dose of study therapy</li> <li>For subjects with prior MEK exposure, Grade 4 toxicity deemed related to the MEK inhibitor</li> <li>Active skin disorder that has required systemic therapy within the past year</li> <li>History of rhabdomyolysis</li> <li>Concurrent ocular disorders</li> <li>Concurrent heart disease or severe obstructive pulmonary disease</li> <li>Subjects with the inability to swallow oral medications</li> </ul>