

Essai Clinique Généré le 25 avr. 2024 à partir de

Titre	Essai clinique ouvert de phase III à répartition aléatoire visant à comparer le REGN2810 à une chimiothérapie au choix du chercheur dans le traitement du cancer du col de l'utérus récidivant ou métastatique, réfractaire au platine
Protocole ID	R2810-ONC-1676
ClinicalTrials.gov ID	NCT03257267
Type(s) de cancer	Col
Phase	Phase III
Stade	Métastatique
Type étude	Traitement
Médicament	Cemiplimab (REGN2810)
Institution	CIUSSS DE L'EST-DE-L'ILE-DE-MONTREAL PAV. MAISONNEUVE/PAV. MARCEL-LAMOUREUX 5415 boul. de l'Assomption, Montréal, QC, H1T2M4
Ville	
Investigateur principal	Dre Suzanne Fortin
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Statut	Fermé
But étude	The primary objective is to compare overall survival (OS) for patients with recurrent or metastatic platinum-refractory cervical cancer treated with either REGN2810 or investigator's choice (IC) chemotherapyThe secondary objectives are: • To compare progression-free survival (PFS) of REGN2810 versus IC chemotherapy • To compare overall response rate (ORR) (partial response [PR] + complete response [CR]) of REGN2810 versus IC chemotherapy per Response Evaluation Criteria in Solid Tumors 1.1 • To compare the duration of response (DOR) of REGN2810 versus IC chemotherapy • To compare the safety profiles of REGN2810 versus IC chemotherapy by describing adverse events (AE) • To compare quality of life (QOL) for patients treated with REGN2810 versus IC chemotherapy using the European Organization for Research and Treatment of Cancer Quality of Life Questionnaire-Core 30 (EORTC QLQ-C30)
Critères d'éligibilité	 Key Inclusion Criteria: Recurrent, persistent, and/or metastatic cervical cancer, for which there is not a curative intent option (surgery or radiation therapy with or without chemotherapy). Acceptable histologies are squamous carcinoma, adenocarcinoma, and adenosquamous carcinoma. Sarcomas and neuro-endocrine carcinomas are not eligible histologies. Tumor progression or recurrence within 6 months of last dose of platinum therapy that was used to treat metastatic, persistent or recurrent cervical cancer Patient must have measurable disease as defined by RECIST 1.1. Eastern Cooperative Oncology Group (ECOG) performance status ≤1 ≥18 years old Adequate organ or bone marrow function Received prior bevacizumab therapy or had clinically documented reason why not administered Received prior paclitaxel therapy or had clinically documented reason why not administered

Critères d'exclusion

Key Exclusion Criteria:

- Ongoing or recent (within 5 years) evidence of significant autoimmune disease that required treatment with systemic immunosuppressive treatments
- Prior treatment with an agent that blocks the PD-1/PD-L1 pathway
- Prior treatment with systemic immune modulating agents (other than anti-PD-1/PD-L1 agents)
 that was within 28 days prior to enrollment, or within 90 days prior to enrollment if there was an
 immune related adverse event, or associated with toxicity that resulted in discontinuation of the
 immune modulating agent
- Active or untreated brain metastases
- Immunosuppressive corticosteroid doses (>10 mg prednisone daily or equivalent) within 4 weeks prior to the first dose of REGN2810
- · Active infection requiring therapy
- History of pneumonitis within the last 5 years
- · Documented allergic or acute hypersensitivity reaction attributed to antibody treatments
- Known allergy to doxycycline or other tetracycline antibiotics
- Concurrent history of malignancy other than cervical cancer within 3 years of first planned dose
 of REGN2810, except for tumors with negligible risk of metastasis