

## Essai Clinique Généré le 28 avr. 2024 à partir de

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Titre	Étude de phase 2, de l'INCMGA00012 chez des participants atteints d'un carcinome à cellules de Merkel métastatique
Protocole ID	INCMGA 0012-201
ClinicalTrials.gov ID	NCT03599713
Type(s) de cancer	Peau
Phase	Phase II
Stade	Métastatique
Type étude	Traitement
Médicament	INCMGA00012
Institution	CIUSSS DU CENTRE-OUEST-DE-L'ILE-DE-MONTREAL  H HOPITAL GENERAL JUIF SIR MORTIMER B.DAVIS  3755 rue de la Côte Ste. Catherine, Montréal, QC, H3T 1E2
Ville	Montréal
Investigateur principal	Dr Wilson Miller
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Statut	Fermé
But étude	Le but de cette étude est d'évaluer l'activité clinique et l'innocuité de l'INCMGA00012 chez les participants atteints d'un carcinome métastatique à cellules de Merkel.
Critères d'éligibilité	<ul> <li>Signed informed consent.</li> <li>Diagnosis of MCC with distant metastatic disease as a component of tumor burden and no more than 3 prior systemic treatments, inclusive of systemic adjuvant therapy.</li> <li>Eastern Cooperative Oncology Group performance status of 0 to 1.</li> <li>Measurable disease according to RECIST v1.1.</li> <li>Availability of tumor tissue (fresh or archival) for central pathology review.</li> <li>Willingness to avoid pregnancy or fathering children based on protocol-defined criteria.</li> </ul>
Critères d'exclusion	<ul> <li>Prior programmed cell death protein 1 (PD-1) or programmed cell death ligand protein 1 (PD-L1)-directed therapy.</li> <li>Treatment with anticancer drugs or participation in another interventional clinical study within 21 days before the first administration of study drug.</li> <li>Has not recovered to ≤ Grade 1 or baseline from toxic effects of prior therapy (with the exceptions for anemia not requiring transfusion support and any grade of alopecia) and/or complications from prior surgical intervention within 7 days before starting study treatment.</li> <li>Radiation therapy administered within 2 weeks of first dose of study treatment or radiation therapy in the thoracic region that is &gt; 30 Gy within 6 months of the first dose of study treatment.</li> <li>Known central nervous system (CNS) metastases and/or carcinomatous meningitis.</li> <li>History of second malignancy within 3 years (with exceptions).</li> <li>Laboratory values outside the protocol-defined range at screening.</li> <li>Clinically significant pulmonary, cardiac, gastrointestinal or autoimmune disorders.</li> <li>Active bacterial, fungal, or viral infections, including hepatitis A, B, and C.</li> <li>Receipt of a live vaccine within 90 days of planned start of study therapy.</li> <li>Current use of protocol-defined prohibited medication.</li> <li>Known hypersensitivity to another monoclonal antibody that cannot be controlled with standard</li> </ul>

- measures (eg, antihistamines and corticosteroids).

   Inability or unlikely, in the opinion of the investigator, to comply with the Protocol requirements.

   Participant who is pregnant or breastfeeding.