

Essai Clinique Généré le 16 mai 2025 à partir de

| Titre | A Randomized Phase 3 Trial of Nivolumab in Combination With Chemo-Immunotherapy for the Treatment of Newly Diagnosed Primary Mediastinal B-Cell Lymphoma |
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| Protocole ID | COG-ANHL1931 |
| ClinicalTrials.gov ID | NCT04759586 |
| Type(s) de cancer | Lymphome non-hodgkinien (LNH) |
| Phase | Phase III |
| Type étude | Clinique |
| Médicament | Nivolumab en association avec de la chimio-immunothérapie |
| Institution | CENTRE HOSPITALIER UNIVERSITAIRE SAINTE-JUSTINE |
| Ville | |
| Investigateur principal | Dre Josette Champagne |
| Coordonnateur | |
| Statut | Actif en recrutement |
| But étude | This phase III trial compares the effects of nivolumab with chemo-immunotherapy versus chemo-immunotherapy alone in treating patients with newly diagnosed primary mediastinal B-cell lymphoma (PMBCL). Immunotherapy with monoclonal antibodies, such as nivolumab, may help the body's immune system attack the cancer, and may interfere with the ability of cancer cells to grow and spread. Treatment for PMBCL involves chemotherapy combined with an immunotherapy called rituximab. Chemotherapy drugs work in different ways to stop the growth of cancer cells, either by killing the cells, by stopping them from dividing, or by stopping them from spreading. Rituximab is a monoclonal antibody. It binds to a protein called CD20, which is found on B cells (a type of white blood cell) and some types of cancer cells. This may help the immune system kill cancer cells. Giving nivolumab with chemo-immunotherapy may help treat patients with PMBCL. |
| Critères d'éligibilité | Age >= 2 years Patient must have histologically confirmed primary mediastinal B-cell lymphoma (PMBCL) as defined by World Health Organization (WHO) criteria Eastern Cooperative Oncology Group (ECOG) performance status of 0, 1, or 2 or ECOG performance status of 3 if poor performance is related to lymphoma Children's Oncology Group (COG) Institutions: Use Karnofsky for patients >= 17 and < 18 years of age and Lansky for patients < 17 years of age Adults (age 18 or older): Creatinine clearance >= 30 mL/min, as estimated by the Cockcroft and Gault formula. The creatinine value used in the calculation must have been obtained within 28 days prior to registration. Estimated creatinine clearance is based on actual body weight Pediatric Patients (age < 18 years): The following must have been obtained within 14 days prior to registration: Measured or calculated (based on institutional standard) creatinine clearance or radioisotope glomerular filtration rate (GFR) >= 70 ml/min/1.73 m^2, or Serum creatinine =< 1.5 x institutional upper limit of normal (IULN), or a serum creatinine based on age/gender as follows: Age : 2 to < 6 year; Maximum serum creatinine (mg/dL): 0.8 (male; 0.8 (female) Age : 10 to < 13 years; Maximum serum creatinine (mg/dL): 1.2 (male); 1.2 (female) Age : 13 to < 16 years; Maximum serum creatinine (mg/dL): 1.5 (male); 1.4 (female) Age :> = 16 years to < 18 years; Maximum serum creatinine (mg/dL): 1.7 (male); 1.4 (female) |

Patients with abnormal liver function will be eligible to enroll if the lab abnormality is thought to be due to the lymphoma or Gilbert's syndrome • Age >= 18 years: Ejection fraction of >= 50% by echocardiogram Age < 18 years: Shortening fraction of >= 27% by echocardiogram, or ejection fraction of >= 50% by radionuclide angiogram Human immunodeficiency virus (HIV)-infected patients on effective anti-retroviral therapy with undetectable viral load within 6 months are eligible for this trial • For patients with evidence of chronic hepatitis B virus (HBV) infection, the HBV viral load must be undetectable on suppressive therapy, if indicated Patients with a history of hepatitis C virus (HCV) infection must have been treated and cured. For patients with HCV infection who are currently on treatment, they are eligible if they have an undetectable HCV viral load • All patients and/or their parents or legal guardians must sign a written informed consent All institutional, Food and Drug Administration (FDA), and National Cancer Institute (NCI) requirements for human studies must be met Critères d'exclusion Administration of prior anti-cancer therapy except as outlined below: • A short course (=< 2 weeks) of corticosteroids for the relief of lymphoma-related • A single course of COP (cyclophosphamide, vincristine, and prednisone)

• Active ischemic heart disease or heart failure

will be permitted if in remission x 3 years

the duration of their study participation

rituximab

• Central nervous system (CNS) involvement of lymphoma

insufficiency is not considered a form of systemic treatment

mg/dL) unless thought to be due to lymphoma or Gilbert's syndrome

• Active uncontrolled infection

 One cycle of chemo-immunotherapy including R-CHOP, DA-EPOCH-R, or a pediatric mature B-cell non-Hodgkin lymphoma (B-NHL) induction therapy (such as ANHL1131)

Previous cancer that required systemic chemotherapy and/or thoracic radiation. Other cancers

 Active autoimmune disease that has required systemic treatment (such as disease modifying agents, corticosteroids, or immunosuppressive agents) in the past 2 years. Replacement therapy such as thyroxine, insulin or physiologic corticosteroid for adrenal or pituitary

In patients < 18 years of age hepatitis B serologies consistent with past or current infections
Patients with severe hepatic impairment (Child-Pugh class C or serum total bilirubin > 5.0

• Lactating females are not eligible unless they have agreed not to breastfeed their infants starting with the first dose of study therapy and for at least 6 months after the last dose of

Female patients who are pregnant since fetal toxicities and teratogenic effects have been noted for several of the study drugs. A pregnancy test is required for female patients of childbearing potential
Sexually active patients of reproductive potential who have not agreed to use a highly effective contraceptive method (failure rate of < 1% per year when used consistently and correctly) for

that has not started more than 21 days prior to enrollment