For Patients with Hodgkin Lymphoma

E4412 Available Through ECOG-ACRIN Cancer Research Group

A Phase I Study with an Expansion Cohort/Randomized Phase II Study of the Combinations of Ipilimumab, Nivolumab and Brentuximab Vedotin in Patients with Relapsed/Refractory Hodgkin Lymphoma

Patient Population
See Section 3.0 for Complete Eligibility Details

 Age ≥ 18 years, ECOG PS 0-2, and adequate lab values
 Must have pathologically confirmed relapsed/refractory classical HL, and measurable disease per protocol
 Must have relapsed after first line chemotherapy; may have relapsed after autologous/allogeneic stem cell transplant, or have primary refractory disease
 May have received prior brentuximab vedotin, ipilimumab, and other prior activating immunotherapies per protocol; no prior nivolumab or PD1/PDL1 axis agents
 No evidence of dyspnea at rest and a pulse oximetry > 92% while breathing room air
 No current or prior history of CNS involvement
 All prior therapy must have been completed at least 21 days prior to enrollment; no concomitant anti lymphoma therapy, including systemic corticosteroids for the purpose of treatment of lymphoma
 No history of Steven’s Johnson’s/TENs syndrome, or motor neuropathy
 Patients with poorly controlled HIV or other chronic active viral infections will be excluded
 Must not have autoimmune disorders or conditions of immunosuppression that require current ongoing treatment with systemic corticosteroids
 No history of symptomatic autoimmune disease, motor neuropathy considered of autoimmune origin, or other CNS autoimmune disease
 No ≥ grade 2 peripheral sensory neuropathy
 No NYHA Class III/IV heart failure, uncontrolled angina, severe uncontrolled ventricular arrhythmias, or electrocardiographic evidence of acute ischemia
 Arms D, E, F, G, H, I, X, Y: cannot have smoked tobacco or other substances within the past 6 months

Phase II (Arms K, L): See protocol for criteria, including modifications for pediatric patients age ≥ 12 years

Treatment Plan
See Section 5.0 for Complete Treatment Details

Refer to protocol for details on the Phase I dose escalation treatment (CLOSED TO ACCRUAL):
 Arms A, B, C, Z: Brentuximab + Ipilimumab
 Arms D, E, F, Y: Brentuximab + Nivolumab
 Arms G, H, I, X: Brentuximab + Nivolumab + Ipilimumab

Treatment design– Phase II:
 1 cycle = 21 days
   Arm K: Brentuximab + Nivolumab
    ◊ Nivolumab 360 mg IV (adult patients) or 3 mg/kg (pediatric patients) day 1 cycles 1-34
    ◊ Brentuximab vedotin 1.8 mg/kg (max 180 mg) IV days 1 cycles 1-16
   Arm L: Brentuximab + Nivolumab + Ipilimumab
    ◊ Ipilimumab 1 mg/kg IV day 1 beginning cycle 1, every 12 weeks for up to 9 doses
    ◊ Nivolumab 360 mg (adult patients) or 3 mg/kg (pediatric patients) day 1 cycles 1-34
    ◊ Brentuximab vedotin 1.8 mg/kg IV (max 180 mg) day 1 cycles 1-16
   A maximum of 16 cycles of brentuximab vedotin will be given, for a maximum of 16 cycles
   A maximum of up to 34 doses of nivolumab (2 years) will be given for patients who are deriving clinical benefit (SD, PR, CR, IR) without excessive toxicity
   Patients who do not have a transplant or other curative therapy option may remain on nivolumab for up to 1 year or 17 further doses after completing brentuximab

Note: 30 minute observation period surrounding therapies per protocol; prophylactic premedication per protocol

Patient Enrollment
All Sites: Oncology Patient Enrollment Network (OPEN) https://open.ctsu.org

Protocol Information

Please Enroll Your Eligible Patients!
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Refer to the protocol for the Phase I dose escalation schemas for Brentuximab Vedotin + Ipilimumab/Nivolumab/Nivolumab + Ipilimumab

ALL CLOSED TO ACCRUAL

Schema – Phase II

Am\text{K} (Nivolumab 360 mg IV (adult patients) or 3 mg/kg (pediatric patients) day 1 cycles 1-34, Brentuximab vedotin 1.8 mg/kg [max 180 mg] IV day 1 cycles 1-16)

Stratify
- Prior Brentuximab-vedotin (BV) use: Yes vs No
- Age: <18 vs ≥18 years old

Am\text{L} (Ipilimumab 1 mg/kg IV day 1 beginning cycle 1, every 12 weeks for up to 9 doses from start of therapy, Nivolumab 350 mg (adult patients) or 3 mg/kg (pediatric patients) day 1 cycles 1-34, Brentuximab vedotin 1.8 mg/kg [max 180 mg] IV day 1 cycles 1-16)

Phase II Accrual Goal = 146 patients
Cycle = 21 days