Abstract and Schema

This is a multicenter Phase I trial for children with recurrent, progressive or refractory primary central nervous system tumors.

Mesenchymal epithelial transition (MET) receptor, also known as c-MET, is an oncogene encoding for a trans-membrane tyrosine kinase receptor activated by the hepatocyte growth factor (HGF). MET has a normal function in organ development during embryogenesis and in tissue homeostasis during adult life.

Abnormal MET activation in cancer correlates with poor prognosis, where aberrantly active MET triggers tumor growth, formation of new blood vessels (angiogenesis) that supply the tumor with nutrients, and cancer spread to other organs (metastasis). Deregulation of HGF/MET signaling pathway is frequently observed in many cancer types of the kidney, liver, stomach, lung, breast and brain, conferring invasive growth and tendency to progression. Thus, HGF and MET are therapeutic targets and anti HGF/MET agents may represent a potential antitumor strategy.

Savolitinib is a potent and selective small molecule inhibitor of c-MET kinase. It is found to inhibit c-MET kinase at the cellular and enzymatic levels with an IC50 between 3-5 nM. In preclinical studies, it demonstrated strong in vitro and in vivo activity against c-MET kinases and its downstream signaling targets and significantly decrease tumor cell growth.

This is a dose escalation study of savolitinib administered orally once a day to patients with recurrent, progressive or refractory central nervous system tumors. There are three stages in this study: a dose escalation cohort, PK expansion cohort, and an efficacy expansion cohort. The dose escalation cohort is designed to determine the maximum tolerated dose (MTD)/recommended Phase II dose (RP2D), to evaluate the pharmacokinetics, safety and preliminary anti-tumor activity of savolitinib. Once the MTD/RP2D is determined, the enrollment for an efficacy expansion cohort will open for patients whose tumors harbor genetic MET activation as determined by CLIA tests performed at participating sites and confirmed by FDA approved tests.

Savolitinib will be given once a day orally for 28 days. 28 consecutive days will constitute one course. The starting dose will be 150 mg/m² (dose level 1). Dosing should be adjusted based on BSA calculated at the beginning of each course. Therapy may continue up to 39 courses (approximately 3 years) in the absence of disease progression or unacceptable toxicity.

<table>
<thead>
<tr>
<th>Dose Escalation Schedule</th>
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<tbody>
<tr>
<td><strong>Dose Level</strong></td>
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<tr>
<td>Level 0</td>
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<tr>
<td>Level 1*</td>
</tr>
</tbody>
</table>
| Level 2 | 240 mg/m² | Patient weight < 50kg  
| | | ≥ 0.63 m² but ≤ 2.00 m²  
| | | Patient weight ≥ 50kg  
| | | ≥ 0.63 m² |
*starting dose
# Patients ≥ 50kg do not have an upper BSA restriction. Please refer to the dosing tables in Appendix Error! Reference source not found. for drug administration guidance.

ELIGIBILITY CRITERIA FOR ENROLLMENT

Diagnosis

- Patients with a histologically confirmed diagnosis of a primary CNS tumor (medulloblastoma, high-grade glioma, or DIPG) that is recurrent, refractory or progressive. All tumors must have histologic verification at either the time of diagnosis or recurrence except patients with diffuse intrinsic brain stem tumors. These patients must have radiographic or clinical evidence of progression.

  Note: Refractory disease is defined as the presence of persistent abnormality on conventional MRI imaging that is further distinguished by histology (biopsy or sample of lesion) or advanced imaging, OR as determined by the treating physician and discussed with the primary investigator prior to enrollment.

- Efficacy Expansion cohort:
  Patients must have a recurrent, progressive or refractory primary CNS tumor with evidence of genetic activation of the MET pathway, regardless of histology. The submitted specimen can be from diagnosis or recurrence and there is no time limit from when the specimen was obtained to enrollment onto the efficacy expansion cohort. The assessment will be performed in a CLIA certified laboratory. MET pathway activation status must be confirmed using FDA approved testing prior to enrollment. MET pathway activation is defined as:

  - MET kinase domain mutations, allelic frequency ≥ 5%
    OR
  - MET or HGF amplification, ≥ 6 copies
    OR
  - Chromosome 7 gain
    OR
  - MET fusion

If a MET aberration is identified using local testing at a PBTC institution, final confirmation for eligibility to the efficacy cohort will be confirmed using MSKCC’s FDA approved IMPACT (Integrated Mutation Profiling of Actionable Cancer Targets) panel. Alternatively, if a MET aberration is identified at a PBTC site using another FDA approved panel (Foundation Medicine or Oncomine), the result will be considered sufficient for eligibility following study PI review. Refer to Appendix Error! Reference source not found. for more information on MET activation testing for the efficacy expansion cohort.
Available Pre-trial Tumor Tissue
Recruiting or refractory primary malignant CNS tumor patients must have adequate pre-trial frozen or FFPE tumor material available for the required correlative studies (section Error! Reference source not found. and 9.1.5). If target amounts of tissue or number of slides are not available, the site must obtain study Chair/co-Chair approval for adequacy of submitted tumor samples and prioritization of studies to be performed, prior to patient enrollment.

Patients with DIPG who have pre-trial tumor tissue available are requested to submit tissue; however, this is not required for eligibility.

Disease Status
Patients must have evaluable disease to be eligible. Evaluable disease is defined as the presence of at least one lesion that can be measured accurately in at least 2 (two) dimensions.

Age
Patients must be > 5 years and ≤ 21 years of age at the time of study enrollment.

BSA
Patients enrolled on 75 mg/m²/day (dose level 0) must have a BSA ≥ 1.00 m².
Patients enrolled on 150 mg/m²/day (dose level 1) must have a BSA ≥ 0.55 m².
Patients enrolled on 240 mg/m²/day (dose level 2) who weigh < 50kg must have a BSA ≥ 0.63 m² but ≤ 2.00 m².
Patients enrolled on 240 mg/m²/day (dose level 2) who weigh ≥ 50kg must have a BSA ≥ 0.63 m².

Prior Therapy
Patients must have failed prior standard therapy for their tumor. Patients with medulloblastoma must have received radiation therapy in addition to platinum and alkylator-based chemotherapy. Patients with HGG and DIPG must have at least received radiation therapy.

Patients must have recovered from the acute treatment related toxicities (defined as ≤ Grade 1 if not defined in eligibility criteria) of all prior chemotherapy, immunotherapy, radiotherapy or any other treatment modality prior to entering this study.

Myelosuppressive chemotherapy
Patients must have received their last dose of known myelosuppressive anticancer therapy at least 21 days prior to enrollment or at least 42 days if it included nitrosourea.

Investigational/Biologic Agent
Biologic or investigational agent (anti-neoplastic):
Patient must have recovered from any acute toxicity potentially related to the agent and received their last dose of the investigational or biologic agent ≥ 7 days prior to study enrollment.

- For agents that have known adverse events occurring beyond 7 days after administration, this period must be extended beyond the time during which adverse events are known to occur.
- Monoclonal antibody treatment and agents with known prolonged half-lives: At least three half-lives must have elapsed prior to enrollment.

Note: A list of the half-lives of commonly used monoclonal antibodies is available on the PBTC webpage under Generic Forms and Templates.

**Radiation**

Patients must have had their last fraction of:
- Craniospinal irradiation or total body irradiation or radiation to ≥ 50% of pelvis > 3 months prior to enrollment.
- Focal irradiation > 4 weeks prior to enrollment

**Stem Cell Transplant**

Patient must be:
- ≥ 6 months since allogeneic stem cell transplant prior to enrollment with no evidence of active graft vs. host disease
- ≥ 3 months since autologous stem cell transplant prior to enrollment

Inclusion of Women and Minorities

Both males and females of all races and ethnic groups are eligible for this study

**Neurologic Status**

- Patients with neurological deficits should have deficits that are stable for a minimum of 1 week prior to enrollment. A baseline detailed neurological exam should clearly document the neurological status of the patient at the time of enrollment on the study.
- Patients with seizure disorders may be enrolled if seizures are well controlled.
- Patients must be able to swallow whole tablets to be eligible for study enrollment.

**Performance Status**

Karnofsky Performance Scale (KPS for > 16 years of age) or Lansky Performance Score (LPS for ≤ 16 years of age) assessed within two weeks of enrollment must be ≥ 50. Patients who are unable to walk because of neurologic deficits, but who are up in a wheelchair, will be considered ambulatory for the purpose of assessing the performance score.

**Organ Function**

Patients must have adequate organ and marrow function as defined below:
- Absolute neutrophil count ≥ 1.0 x 10^9 cells/L
- Platelets ≥ 100 x 10^9 cells/L (unsupported, defined as no platelet transfusion within 7 days prior to enrollment)
- Hemoglobin ≥ 8g/dL (hemoglobin should be unsupported, i.e. red blood cell transfusions are not allowed within 14 days prior to enrollment)
- Adequate liver function defined as:
  - Alanine aminotransferase (ALT) and aspartate aminotransferase (AST) ≤ 2.5 x the upper limit of normal (ULN) with Total Bilirubin ≤ 1x ULN
  Or
  - Total Bilirubin >ULN-≤1.5x ULN with ALT and AST ≤ 1x ULN
- Albumin $\geq 2$ g/dL
- Serum creatinine based on age/gender as noted in Table 1. Patients that do not meet the criteria in Table 1 but have a 24 hour Creatinine Clearance or GFR (radioisotope or iothalamate) $\geq 70$ mL/min/1.73 m$^2$ are eligible.

<table>
<thead>
<tr>
<th>Age</th>
<th>Maximum Serum Creatinine (mg/dL)</th>
<th>Male</th>
<th>Female</th>
</tr>
</thead>
<tbody>
<tr>
<td>2 to &lt; 6 years</td>
<td>0.8</td>
<td>0.8</td>
<td></td>
</tr>
<tr>
<td>6 to &lt; 10 years</td>
<td>1</td>
<td>1</td>
<td></td>
</tr>
<tr>
<td>10 to &lt; 13 years</td>
<td>1.2</td>
<td>1.2</td>
<td></td>
</tr>
<tr>
<td>13 to &lt; 16 years</td>
<td>1.5</td>
<td>1.4</td>
<td></td>
</tr>
<tr>
<td>$\geq 16$ years</td>
<td>1.7</td>
<td>1.4</td>
<td></td>
</tr>
</tbody>
</table>

The threshold creatinine values in this Table were derived from the Schwartz formula for estimating GFR (Schwartz et al. J. Peds, 106:522, 1985) utilizing child length and stature data published by the CDC.

- Coagulation parameters: INR $< 1.5 \times$ ULN and aPTT $< 1.5 \times$ ULN unless patients are receiving therapeutic anti-coagulation which affects these parameters.

- Patients with known tumor thrombus or deep vein thrombosis are eligible if clinically stable on low molecular weight heparin for $\geq 2$ weeks.

- Cardiac Function:
  - Left Ventricular Ejection Fraction $> 55\%$ on ECHO

- Pulmonary Function
  - Oxygen saturation as measured by pulse oximetry is $> 93\%$ on room air

Corticosteroids
Patients who are receiving corticosteroids must be on a stable or decreasing dose for at least 1 week prior to enrollment.

Growth Factors
Patients must be off all colony-stimulating factor(s) (e.g., filgrastim, sargramostim or erythropoietin) for at least 1 week prior to enrollment. 2 (two) weeks must have elapsed if patients received PEG formulations.

Pregnancy Status
Female patients of childbearing potential must not be pregnant or breast-feeding. Female patients of childbearing potential must have a negative serum or urine pregnancy test.

Pregnancy Prevention
- Patients of childbearing or child fathering potential must be willing to use a medically acceptable form of birth control, which includes abstinence, while being treated on this study.
• Women of child-bearing potential should use effective contraception from the time of enrollment until 4 weeks after discontinuing study treatment.
• Male study participants should use a condom with female partners of child-bearing potential during the study and for 4 weeks after discontinuing study treatment.
• If the female partner of a male study participant is not using effective contraception, men must use a condom during the study and for 6 months after discontinuing study treatment.
• Male study participants should avoid fathering a child and refrain from sperm donation from study start to 6 months after discontinuing study treatment.

Informed Consent
The patient or parent/guardian is able to understand the consent and is willing to sign a written informed consent document according to institutional guidelines.

EXCLUSION CRITERIA

Breast-feeding
Female patients who are breast-feeding.

Concurrent Illness
• Patients with a known serious active infection including, but not limited to, viral hepatitis, human immunodeficiency virus, tuberculosis, or with any clinically significant unrelated systemic illness or significant cardiac, pulmonary, hepatic or other organ dysfunction), that in the opinion of the investigator would compromise the patient’s ability to tolerate protocol therapy, put them at additional risk for toxicity or would interfere with the study procedures or results.
• Patients with uncontrolled hypertension (i.e., a blood pressure (BP) > 95th percentile for age, height and gender, patients with values above these levels must have their blood pressure controlled with medication prior to starting study drug)
  • The normal blood pressure by height, age and gender tables can be assessed in the Generic Forms section of the PBTC member’s webpage.

• Patients with any of the following cardiac diseases
  • Congestive heart failure (New York Heart Association ≥ Grade 2) (see Appendix Error! Reference source not found.)
  • Clinically significant cardiac arrhythmia
  • Mean resting corrected QT interval (QTc) > 450 msec on screening obtained from 3 electrocardiograms (EKGs) or factors that may increase the risk of QTc prolongation such as chronic hypokalemia not correctable with supplements, congenital or familial long QT syndrome, or family history of unexplained sudden death under 40 years of age in first-degree relatives or any concomitant medication known to prolong the QT interval and cause Torsade de Pointes. Refer to Appendix 15.3 for a partial list of medications that are associated with QTc prolongation and Torsade de Pointes. These drugs must have been discontinued prior to the start of administration of study treatment in accordance with guidance provided in Error! Reference source not found. of Appendix 15.3.
• Any clinically important abnormalities in rhythm, conduction or morphology of resting EKG, e.g. complete left bundle branch block, third degree heart block, second degree heart block, PR interval >250 msec.

• Patients with a history of any other malignancy, except patients with a secondary brain tumor if the patient’s first malignancy has been in remission for at least 5 years from the end of treatment.

Concurrent Therapy
• Patients who are receiving any other anticancer or investigational drug therapy.
• Patients receiving strong inducers of CYP3A4, strong inhibitors of CYP3A4 or CYP1A2 or CYP3A4 substrates with a narrow therapeutic index within 2 weeks of the first dose of savolitinib (3 weeks for St John's Wort). Strong inducers of CYP3A4 and CYP3A4 substrates which have a narrow therapeutic range or CYP3A4 sensitive substrates should not be used during the trial or used with caution. Because the lists of these agents are constantly changing, it is important to regularly consult a frequently-updated medical reference. Appendix Error! Reference source not found. (Patient Drug Information Handout and Wallet Card) should be provided to patients.
• Prior or current treatment with a MET inhibitor (e.g., foretinib, crizotinib, cabozantinib, or onartuzumab).

Herbal preparations/medicines
Patients is currently receiving any of the following herbal preparations or medications and cannot be discontinued 1 week (7 days) prior to enrollment (3 weeks for St. John’s wort). These herbal medications include, but are not limited to: cannabis products, St. John’s wort, kava, ephedra (ma huang), gingko biloba, dehydroepiandrosterone (DHEA), yohimbe, saw palmetto, and ginseng.

Surgical procedures
Patient has undergone major surgical procedure ≤ 28 days prior to beginning study drug or a minor surgical procedure ≤ 7 days prior to beginning study drug. No waiting is required following port-a-cath placement.

Inability to Participate
Patients who in the opinion of the investigator are unwilling or unable to return for required follow-up visits or obtain follow-up studies required to assess toxicity to therapy or to adhere to drug administration plan, other study procedures, and study restrictions.

Allergy
Patients with a history of allergic reactions attributed to compounds of similar chemical or biologic composition.

Prisoners
Prisoners will be excluded from this study.