**Principal Investigator: Name and Contact Research Coordinator: Name and Contact**

**Baseline/day 1– Could be same day as screening.**

-PI: fdg

Mardsf dsf lkdlk

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Information for Inpatient Nursing

**Huddle sheets should answer the following questions:**

**1) Will this study utilize CHPS scattersite?**

**2) What does the inpatient bedside nurse need to know?**

**3) What does the inpatient bedside nurse need to do related to the study? If this is a CHPS scattersite, differentiate between CHPS nurse procedures and inpatient nurse procedures**

Example:

Infusion/Medication Information

Review of supplies and equipment that may be needed for investigational product administration

Documentation needed in Penn Chart or on paper form

Vital signs, monitoring, ECG, labs

**Supplemental sheets may be necessary for IP/medications and should include the following information as necessary on a separate word document (*the supplemental sheet does not take the place of provide orders*)**

Medication name

Mechanism of action

Side effect

Dosing notes

Additional exclusions

Medication/IP administration instructions

IV Administration: \*\**Must be administered within 4 hours of preparation*\*\*

IV Administration: Administer with non-DEHP tubing via dedicated line; flush with 10 ml of saline before and after infusion

NGT administration: Study drug should be administered with food or feeds. Administer the study drug immediately after feeding the patient. If patient is NPO except meds, may be administered alone.

After administering tube feeding, flush the tube with water prior to study drug administration. Transfer the contents of one 400 mg capsule (400 mg total) to a clean medicine cup.

Add 30 ml of room temperature water to the medicine cup and stir vigorously to suspend the material.

Draw the suspension into a large syringe (eg, 60 mL syringe) and administer via the feeding tube.

After this first administration, flush the tube with room temperature water.

To ensure all study drug has been delivered, add 30 ml of room temperature water to medicine cup and stir to include any potentially remaining material.

Draw the fluid into the same large syringe (eg, 60 mL syringe) and transfer to the patient via the feeding tube.

After this final administration of the study drug, flush the tube with room temperature water.

**Title of the study:** Please include the IRB number and the full title of the protocol with abbreviations spelled out

**Study Design:** Observation or Intervention

**Target Population:**

· Provide a brief overview of disease type and diagnosis of patients that will be included

· Please do not include all of inclusion/ exclusion criteria here

· Avoid specific pathology and molecular information example: do not include “patients with histologically confirmed R-RCHL or R– R TCL according to the 2016 WHO Classification”

- Instead say Non Hodgkin's Lymphoma who failed multiple lines therapy