

ETUDE TAS-120-202

Identité patient (coller étiquette patient)

Version 1.0 du 10/03/2015

Investigateur:

Arc: Hélène

VALIDATION DES CRITERES DE SELECTION

Critères d'inclusion

onteres a metasion	
Provide written informed consent	□ oui
	□ non
2. ≥18 years of age (or meets the country's regulatory definition for legal adult age, whichever is	□ oui
greater)	□ non
3.Eastern Cooperative Oncology Group (ECOG) performance status 0 or 1	□ oui
	□ non
4. Has recovered from the acute toxic effects of prior anticancer therapy to baseline or	□ oui
Grade 1 (except toxicities which are not clinically significant such as alopecia)	□ non
5. Known FGFR aberration status and tumor type that meet all of the criteria for 1 of the	□ oui
following cohorts:	□ non
a : Cohort A	
i. Histologically-confirmed, locally-advanced, advanced, or metastatic solid tumors harboring a FGFR1-4 rearrangement determined in tumor tissue using nextgeneration	
sequencing (NGS), fluorescence in situ hybridization (FISH), or other	
assays that can determine gene rearrangements in tumor tissues. Patients with primary	
brain tumor or intrahepatic cholangiocarcinoma are not eligible.	
ii. Measurable disease per RECIST 1.1	
iii. Had disease progression/recurrence after standard treatment for their advanced or metastatic cancer	
b : Cohort B	
i. Histologically-confirmed, locally-advanced, advanced, or metastatic gastric or GEJ	
cancer harboring a FGFR2 amplification. The tumor must have an FGFR2/CEN10	
ratio of ≥5 or an FGFR2 copy number ≥10 signals per cell determined in tumor tissue	
using NGS, FISH, or other assays that can determine gene amplifications in tumor	
tissues. ii. Measurable disease per RECIST 1.1	
iii. Received at least 2 prior systemic regimens for advanced/metastatic disease	
iv. Experienced disease progression/recurrence during or after the most recent prior	
systemic treatment for advanced/metastatic gastric or GEJ cancer	
6. Has archival or fresh tumor tissue (preferably in block format) available to send to central	□ oui
laboratory.	□ non



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7. Adequate organ function as defined by the following criteria:	□ oui
	□ non
a. Cohorts A and B:	
i. Absolute neutrophil count (ANC) ≥ 1.0 × 109/L	
ii. Platelet count ≥ 75,000/mm3 (≥ 75 × 109/L)	
iii. Hemoglobin ≥ 9.0 g/dL	
iv. ALT and aspartate aminotransferase (AST) ≤ 3.0 × upper limit of normal (ULN); if	
liver function abnormalities are due to underlying liver metastasis, AST and ALT ≤ 5.0 × ULN.	
v. Total bilirubin $\leq 1.5 \times \text{ULN}$, or $\leq 3.0 \times \text{ULN}$ for patients with Gilbert's syndrome.	
vi. Creatinine clearance (CrCl) (calculated or measured value): ≥40 mL/min. For	
calculated CrCl, use the Cockcroft-Gault formula (Section 6).	
vii. Phosphorus < 1.5 ULN	
8. Women of child-bearing potential (WOCBP) must have a negative serum pregnancy test	□ oui
prior to administration of the first dose of futibatinib. Female patients are not considered	□ non
to be of child-bearing potential if they are post-menopausal (no menses for 12 months	
without an alternative medical cause) or permanently sterile (hysterectomy, bilateral	
salpingectomy, or bilateral oophorectomy).	
9. Both males and females of reproductive potential must agree to use effective birth control	□ oui
during the study prior to the first dose and for 90 days after the last dose or longer based	□ non
on local requirements.	
10. Ability to take medications orally (feeding tube is not permitted).	□ oui
	□ non
11. Willing and able to comply with scheduled visits and study procedures.	□ oui
	□ non



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Critères de non inclusion

<u> </u>	
1. Currently receiving an investigational drug in a clinical trial or participating in any other	□ oui
type of medical research judged not to be scientifically or medically compatible with this	□ non
study. If a patient is currently enrolled in a clinical trial involving non-approved use of a	
device, then agreement with the investigator and Taiho Medical monitor is required to	
establish eligibility.	
History and/or current evidence of any of the following disorders:	□ oui
2. Thistory analytic current evidence of any of the following alsoraers.	□ non
a. Non-tumor related alteration of the calcium-phosphorus homeostasis that is considered clinically	
significant in the opinion of the Investigator.	
significant in the opinion of the investigator.	
h Ectanic minoralization/calcification, including but not limited to coft tissue, kidneys, intesting, or	
b. Ectopic mineralization/calcification, including but not limited to soft tissue, kidneys, intestine, or	
myocardia and lung, considered clinically significant in the opinion of	
the Investigator.	
c. Retinal or corneal disorder confirmed by retinal/corneal examination and considered clinically	
significant in the opinion of the Investigator.	
3. Corrected QT interval using Fridericia's formula (QTcF) >470 msec. Patients with an atrioventricular	□ oui
·	□ non
pacemaker or other condition (for example, right bundle branch block) that renders the QT	
measurement invalid are an exception and the criterion does not apply.	
4. Treatment with any of the following within the specified time frame prior to the first dose	□ oui
of futibatinib:	□ non
a. Major surgery within 4 weeks (surgical incision should be fully healed)	
b. Radiotherapy for extended field within 4 weeks or limited field radiotherapy within 2 weeks	
c. A drug that has not received regulatory approval for any indication within 14 or 21 days of treatment	
for a nonmyelosuppressive or myelosuppressive agent, respectively	
5. Received strong inhibitors and inducers of CYP3A4 within 2 weeks	□ oui
	□ non
6. Prior treatment with an FGFR inhibitor	□ oui
	□ non
7. A serious illness or medical condition(s) including, but not limited to, the following:	□ oui
	□ non
a. Known acute systemic infection	
b. Myocardial infarction, severe/unstable angina, or symptomatic congestive heart failure within the	
previous 6 months	
c. History or current evidence of uncontrolled ventricular arrhythmia	
d. Chronic diarrhea diseases considered to be clinically significant in the opinion of the Investigator	
e. Congenital long QT syndrome, or any known history of torsade de pointes, or family history of	
unexplained sudden death	
f. Other severe acute or chronic medical or psychiatric condition or laboratory abnormality that may	
increase the risk associated with study participation or futibatinib administration, or may interfere	
with the interpretation of study results, and in the judgment of the Investigator would make the	
patient inappropriate for entry into this study	
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8. Active central nervous system (CNS) metastasis and/or carcinomatous meningitis.	□ oui
Patients with previously treated brain metastases that are clinically and radiologically stable (for at	□ non
least 4 weeks prior to enrollment) are eligible.	
9. Known additional malignancy that is progressing or has required active treatment within	□ oui
the past 2 years. Patients with basal cell carcinoma of the skin, squamous cell carcinoma	□ non
of the skin, or carcinoma in situ (eg, breast carcinoma, cervical cancer in situ) that have undergone	
potentially curative therapy are not excluded.	
10. Pregnant or breastfeeding.	□ oui
	□ non
	□ oui
	□ non
Date:	
Date :	
Signature de l'investigateur :	

Formulaire PC BECT OPC 03 - Version 01