KURZPROTOKOLL PANTHEON

Öffentlicher Titel

Studienart

Phase II Studie zum Überleben nach sequenzieller Chemotherapie des metastasierten Bauchspeicheldrüsenkrebs

Wissenschaftl. Titel

Einschlusskriterien

A Health Service Research Study to Investigate Survival of Metastatic Pancreatic Cancer Patients After Sequential Chemotherapy: An AIO Phase II Cross Over Trial (PANTHEON)

Kurztitel PANTHEON

multizentrisch, prospektiv, Therapiestudie, randomisiert, offen/unverblindet, zweiarmig, Investigator Initiated Trial (IIT)

Studienphase Phase II

Erkrankung Verdauung: Bauchspeicheldrüsenkrebs (Pankreaskarzinom): Zweitlinie oder höher

- Written informed consent and any locally-required authorization (EU Data Privacy Directive in the EU) obtained from the subject prior to performing any protocol-related procedures, including screening evaluations
- Age>= 18 years at time of study entry Unresectable adenocarcinoma of the pancreas previously treated in the palliative setting with gemcitabine and nabpaclitaxel (Abraxane®)
- Adequately documented recurrence and disease status after/under 1st line (Best response, duration of treatment, time to progression, preexisting PNP and other side effects)
- Radiologically confirmed disease progression during 1st-line therapy and measurable reference cancer site(s) as defined by RECIST1.1
- Randomization and start of 2nd-line treatment possible within 4 weeks after radiologically documented disease progression during 1st-line therapy
- ECOG performance status 0-2 8. No prior radiotherapy
- Adequate blood count, liver-enzymes, and renal function:
- Absolute neutrophil count (ANC) >= 1.5 x 10^9/L (> 1500 per mm³)
- Platelet count >= 100 x 10^9/L (>100,000 per mm³)
- AST (SGOT)/ALT (SGPT) < 2.5 x institutional upper limit of normal unless liver metastases are present, in which case it must be < 5x ULN
- - Serum creatinine CL >= 60 mL/min calculations according to local standard
- Bilirubin < 3 ULN 10. Female subjects must either be of non-reproductive potential (ie, post-menopausal by history: 60 years old and no menses for >= 1 year without an alternative medical cause; OR history of hysterectomy, OR history of bilateral tubal ligation, OR history of bilateral oophorectomy) or must have a negative serum pregnancy test upon study entry
- Subject is willing and able to comply with the protocol for the duration of the study including undergoing treatment and scheduled visits and examinations including follow up

Ausschlusskriterien

- Serious cardiovascular disease (eg, unstable coronary artery disease or myocardial infarction within 3 months prior to study start)
- Preexisting polyneuropathy (PNP) >= grade 3 [National Cancer Institute Common Toxicity Criteria grade 3 or 4 sensory or motor neuropathy]
- Prior or concurrent malignancy (other than pancreatic cancer) which either progresses or requires active treatment. Exceptions are: basal cell cancer of the skin
- History of DPD deficiency
- Morbus Gilbert
- History of hypersensitivity to any of the study drugs or any of the constituents of the products
- Medication that is known to interfere with any of the agents applied in the trial

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- Female subjects who are pregnant, breast-feeding or male or female patients of reproductive potential who are not employing an effective method of birth control (failure rate of less than 1% per year)
- Any condition that, in the opinion of the investigator, would interfere with evaluation of study treatment or interpretation of patient safety or study results
- Any medical condition that contraindicates dosing with any of the IMPs or constitutes a safety risk for the patient including but not limited to:
- - chronic inflammatory bowel disease and/or bowel obstruction.
- active uncontrolled infection
- - clinically significant bleeding or bleeding diathesis
- clinically significant stomatitis
- - active ulceration of the gastrointestinal tract
- Previous enrollment or randomization in the present study (does not include screening failure)
- Patient who has been incarcerated or involuntarily institutionalized by court order or by the authorities § 40 Abs. 1 S. 3 Nr. 4 AMG
- Patients who are unable to consent because they do not understand the nature, significance and implications of the clinical trial and therefore cannot form a rational intention in the light of the facts [§ 40 Abs. 1 S. 3 Nr. 3a AMG]

Alter

Prüfzentren

18 Jahre und älter

Agaplesion Markus Krankenhaus (Geschlossen)

Wilhelm-Epstein-Straße 4 60431 Frankfurt am Main Dr. med. Claus Bolling Tel: 069 95332206 Fax: 069 95332098

Fax: 069 95332098 claus.bolling@fdk.info

Universitätsklinikum Frankfurt (Geschlossen)

Medizinische Klinik I, Gastroenterologie/Hepatologie

Theodor-Stern-Kai 7 60590 Frankfurt am Main

Wina Hensel

Tel: 069 6301-87769 Fax: 069 6301-6580 wina.hensel@kgu.de

Sponsor

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Registrierung in anderen Studienregistern

EudraCT 2016-004640-11 ClinicalTrials.gov NCT03331640