

## **KURZPROTOKOLL** **RASH**

<b>Öffentlicher Titel</b>	Gemcitabin plus Erlotinib bei Rash-positivem, metastasiertem Pankreaskarzinom und günstigen Risikofaktoren
<b>Wissenschaftl. Titel</b>	Phase II Studie zur Bestimmung der Effektivität von Gemcitabin plus Erlotinib bei Rash-positiven Patienten mit metastasiertem Pankreaskarzinom und günstigen Risikofaktoren
<b>Kurztitel</b>	RASH
<b>Studienart</b>	multizentrisch, prospektiv, offen/unverblindet, zweiarmig, Investigator Initiated Trial (IIT)
<b>Studienphase</b>	Phase II
<b>Erkrankung</b>	Verdauung: Bauchspeicheldrüsenkrebs (Pankreaskarzinom): Erstlinie
<b>Ziele</b>	<ul style="list-style-type: none"><li>- die 1-Jahresüberlebensrate von "good-risk" Patienten, die unter einer Behandlung mit Gemcitabin/Erlotinib einen Rash entwickeln</li><li>- ORR, DCR, PFS, OS</li><li>- Verträglichkeit</li><li>- Translationales Projekt</li></ul>
<b>Einschlusskriterien</b>	<ul style="list-style-type: none"><li>- Histologically (not cytologically) confirmed metastatic pancreatic adenocarcinoma (stage IV according to UICC, each T, each N, M1 according to TNM)</li><li>- At least one measurable index lesion (CT or MRI) according to RECIST criteria (V 1.1)</li><li>- ECOG PS 0 and 1</li><li>- Age 18-75 years</li><li>- Serum bilirubin <math>\leq 1,5</math>x ULN (a placed biliary tract stent without concurrent cholangitis is not considered a contraindication)</li><li>- Availability of tumour samples (no cytologic samples)</li><li>- Written informed consent by the patient for collecting blood- and tumour-samples for translational research according to study protocol</li><li>- Live expectancy of at least three months</li><li>- Written informed consent</li><li>- Negative pregnancy test in women with childbearing potential (to be performed within 7 days prior to treatment start)</li><li>- Adequate kidney-, liver- and bone-marrow function: neutrophils <math>\geq 1500/\mu\text{l}</math>, platelets <math>\geq 100.000/\mu\text{l}</math>, and hemoglobin <math>\geq 8\text{g/dl}</math>, liver transaminases <math>\leq 2,5</math>x ULN, in case of liver metastases <math>\leq 5</math>x ULN, serum creatinine <math>\leq 1,25</math>x ULN, creatinine clearance <math>\geq 30</math> ml/min</li><li>- Legal capacity of the patient</li><li>- Option for constant long-term follow-up</li></ul>
<b>Ausschlusskriterien</b>	<ul style="list-style-type: none"><li>- Resectable pancreatic carcinoma</li><li>- Locally advanced pancreatic cancer (non-resectable tumour without distant metastasis)</li><li>- Previous palliative chemotherapy for metastatic or locally advanced, non-resectable pancreatic cancer</li><li>- Previous palliative radiation or chemoradiation for locally advanced, non-resectable pancreatic cancer</li><li>- Radiation therapy within four weeks prior to study enrolment or radiation of indicator lesions</li><li>- Adjuvant Chemotherapy or Radiochemotherapy for pancreatic cancer <math>\leq 6</math> months prior to study enrolment</li><li>- All previously occurred metastatic cancers or cured neoplasias diagnosed within the last 5 years before study enrolment</li></ul>

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- Major surgery within 2 weeks before study start
- Chronic diarrhea
- Known glucuronidation-deficiency (Gilbert´s syndrome)
- Acute or subacute ileus or chronic inflammatory bowel disease
- Preexisting polyneuropathy > Grade I according to NCI-CTCAE v.4.0
- Relevant comorbidities which might impair patient eligibility or safety for study participation like active infections, hepatic, renal or metabolic diseases
- Clinically significant cardiovascular diseases within 12 months prior to study start (e.g. unstable angina pectoris, myocardial infarction, heart failure  $\geq$  NYHA II, cardiac arrhythmias requiring treatment)

<b>Alter</b>	18 - 75 Jahre
<b>Status</b>	Rekrutierung beendet
<b>Prüfzentren</b>	<b>Krankenhaus Nordwest GmbH</b> Institut für klinisch-onkologische Forschung Steinbacher Hohl 2-26 60488 Frankfurt am Main Prof. Dr. med. Salah-Eddin Al-Batran Tel: 069 7601 4420 <a href="mailto:albatran@khnw.de">albatran@khnw.de</a>
<b>Sponsor</b>	Universitätsklinikum München (Hauptsponsor)
<b>Förderer</b>	Universitätsklinikum München
<b>Registrierung in anderen Studienregistern</b>	ClinicalTrials.gov NCT01729481 EudraCT 2011-005471-17