

<b>Öffentlicher Titel</b>	Phase I-Studie mit intrakranieller Injektion von NK-Zellen bei rezidiviertem HER2-positivem Glioblastom
<b>Wissenschaftl. Titel</b>	Monozentrische, offene Phase I-Studie mit intrakranieller Injektion von NK-92/5.28.z (HER2.taNK) Zellen bei Patienten mit rezidiviertem HER2-positivem Glioblastom
<b>Kurztitel</b>	Car2Brain
<b>Studienart</b>	prospektiv, Therapiestudie, offen/unverblindet, einarmig, Investigator Initiated Trial (IIT)
<b>Studienphase</b>	Phase I
<b>Erkrankung</b>	NEURO: Glioblastom (WHO Grad IV): Zweitlinie oder höher
<b>Einschlusskriterien</b>	<ul style="list-style-type: none"> <li>- Recurrent or refractory HER2-positive glioblastoma or its variant gliosarcoma in which a relapse surgery (partial or total) is being planned.</li> <li>- Prior therapy must include the standard of care for glioblastoma (radiotherapy plus concomitant and adjuvant chemotherapy with temozolomide according to the EORTC 26981 trial).</li> <li>- Life expectancy <math>\geq</math> 3 months</li> <li>- Bilirubin <math>\leq</math> 3x normal, AST <math>\leq</math> 5x normal, ALT <math>\leq</math> 5x, serum creatinine <math>\leq</math> 2x upper limit of normal for age, leukocyte count = 3nl, thrombocyte count = 100nl and Hb = 8.0 gdl</li> <li>- Blood oxygenation of 90% as measured by pulse oximetry on room air</li> <li>- Women must have a negative serum pregnancy test within 72h prior to the start of the first NK-925.28.z cell injection.</li> <li>- Sexually active patients must be willing to utilize effective birth control methods throughout the study and for 24 weeks after the last NK-925.28.z cell injection. This includes two different forms of effective contraception (e.g. hormonal contraceptive and condom, IUDIUS and condom) or sterilization.</li> <li>- Patients should have been off other antineoplastic therapy for two weeks prior to entry in this study. Temozolomide will be allowed up to 48h preinjection. At the the time of inclusion, dexamethasone up to a total dose of 4 mg per day will be allowed if medically indicated.</li> <li>- Informed consent explained to and signed by patient; patient given copy of informed consent.</li> <li>- Karnofsky performance score of <math>\geq</math> 50%</li> </ul>
<b>Ausschlusskriterien</b>	<ul style="list-style-type: none"> <li>- Previous anti-angiogenic therapy e.g. with bevacizumab (Avastin)</li> <li>- Coagulation disorder (INR<math>&gt;</math>1.4 or PTT<math>&gt;</math>50sec) or anticoagulation at therapeutic dosage</li> <li>- Active autoimmune disease</li> <li>- Patients with clinical or laboratory signs for immunodeficiency or under immunosuppressive medication other than corticosteroids</li> <li>- Severe intercurrent infection</li> <li>- Known HIV, HBV or HCV positivity</li> <li>- Chronic heart failure NYHA <math>\geq</math>III</li> <li>- Patients with a prior solid organ transplantation or allogenic haematopoietic stem cell transplantation</li> <li>- Unable to undergo MRI</li> <li>- Drug or alcohol abuse</li> <li>- Pregnancy or breastfeeding</li> <li>- Severe psychiatric disorder which might interfere with the study treatment or examination</li> </ul>

- Simultaneous participation in another clinical trial. If a subject participated in a trial testing another IMP, such IMP should have been terminated at least 30 days before inclusion of the subject.

<b>Alter</b>	18 Jahre und älter
<b>Molekularer Marker</b>	HER2/neu pos.
<b>Status</b>	Aktiv
<b>Beginn der Rekrutierung</b>	01.12.2017
<b>Prüfzentren</b>	<b>Universitätsklinikum Frankfurt</b> Dr. Senckenbergisches Institut für Neuroonkologie Theodor-Stern-Kai 7 60590 Frankfurt am Main Dr. med. Michael Burger <a href="mailto:michael.burger@kgu.de">michael.burger@kgu.de</a>
<b>Registrierung in anderen Studienregistern</b>	ClinicalTrials NCT03383978 (primäres Register) EUDRACT 2016-000225-39