## KURZPROTOKOLL GMALL-MOLACT1-BLINA

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Öffentlicher Titel	Blinatumomab bei MRD-positiver B-Vorläufer ALL
Wissenschaftl. Titel	A multicenter, single-arm study to assess the efficacy, safety, and tolerability of the BiTE® antibody blinatumomab in adult patients with minimal residual disease (MRD) of B-precursor acute lymphoblastic leukemia (Blast Successor Trial)
Kurztitel	GMALL-MOLACT1-BLINA
Studienart	multizentrisch, prospektiv, Therapiestudie, offen/unverblindet, einarmig, Investigator Initiated Trial (IIT)
Studienphase	Phase II
Erkrankung	Blut: Akute lymphatische Leukämie (ALL): Rezidiviert/refraktär
Einschlusskriterien	<ul> <li>Patients with CD19 positive B-precursor ALL in complete hematological remission defined as less than 5% blasts in bone marrow after at least three intense chemotherapy blocks (e.g., GMALL induction I-II/consolidation I).</li> </ul>
	<ul> <li>Presence of minimal residual disease (MRD) at a level of &gt;=10-4 (molecular failure or molecular relapse) in an assay with a minimum sensitivity of 10-4 documented after an interval of at least 2 weeks from last systemic chemotherapy</li> </ul>
	<ul> <li>For evaluation of MRD patients must have at least one molecular marker based on individual rearrangements of immunoglobulin, TCR-genes or other suitable genes evaluated by the reference laboratory of the trial</li> </ul>
	<ul> <li>Bone marrow function as defined below: (a) ANC (Neutrophils) &gt;= 1,000/µL; (b)</li> <li>Platelets &gt;= 50,000/µL (transfusion permitted); (c) HB level &gt;= 9g/dl (transfusion permitted)</li> </ul>
	<ul> <li>Renal and hepatic function as defined below: (a) AST (GOT), ALT (GPT), and AP &lt; 5 x upper limit of normal (ULN); (b) Total bilirubin &lt; 1.5 x ULN (unless related to Gilbert's Meulengracht disease); (c) Creatinine &lt; 1.5x ULN; (d) Creatinine clearance &gt;= 60 mL/min (e.g. calculated according Cockroft&amp;Gault)</li> </ul>
	- Negative HIV test, negative hepatitis B (HbsAg) and hepatitis C virus (anti-HCV) test
	<ul> <li>Negative pregnancy test in women of childbearing potential</li> </ul>
	- ECOG Performance Status 0 or 1
	- Age >=18 years
	- Ability to understand and willingness to sign a written informed consent
	- Signed and dated written informed consent is available
	<ul> <li>Participation in the registry of the German Multicenter Study Group for Adult ALL (GMALL)</li> </ul>
Ausschlusskriterien	- Ph/BCR-ABL positive ALL
	<ul> <li>Presence of circulating blasts or current extramedullary involvement by ALL</li> </ul>
	<ul> <li>History or presence of clinically relevant CNS pathology (e.g. seizure, paresis, aphasia, cerebrovascular ischemia/hemorrhage, severe brain injuries, dementia, Parkinson's disease, cerebellar disease, organic brain syndrome or psychosis)</li> </ul>
	- Current detection of ALL blast cells in cerebro-spinal fluid
	- History of or active relevant autoimmune disease
	<ul> <li>Systemic cancer chemotherapy within 2 weeks prior to study treatment (except for intrathecal prophylaxis)</li> </ul>
	- Radiotherapy within 4 weeks prior to study treatment
	- Live vaccination within 2 weeks before the start of study treatment
	<ul> <li>Autologous hematopoietic stem cell transplantation (SCT) within six weeks prior to study treatment</li> </ul>
	- Allogeneic SCT within 12 weeks before the start of study treatment

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	<ul> <li>Any active acute Graft-versus-Host Disease (GvHD), grade 2-4 according to the Glucksberg criteria or active chronic GvHD requiring systemic treatment</li> </ul>
	- Any systemic therapy against GvHD within 2 weeks before start of study treatment
	<ul> <li>Therapy with monoclonal antibodies (rituximab, alemtuzumab) within 4 weeks prior to study treatment</li> </ul>
	- Treatment with any investigational product within four weeks prior to study treatment
	- Previous treatment with blinatumomab or other anti-CD19-therapy
	<ul> <li>Known hypersensitivity to immunoglobulins or to any other component of the study drug formulation</li> </ul>
	<ul> <li>History of malignancy other than ALL diagnosed within 5 years prior to start of protocol-specified therapy with the exception of: (a) Adequately treated non- melanoma skin cancer or lentigo maligna without evidence of disease; (b) Adequately treated cervical carcinoma in situ without evidence of disease; (c) Adequately treated breast ductal carcinoma in situ without evidence of disease; (d) Prostatic intraepithelial neoplasia without evidence of prostate cancer</li> </ul>
	<ul> <li>Active infection, any other concurrent disease or medical condition that are deemed to interfere with the conduct of the study as judged by the investigator</li> </ul>
	- Nursing women
	<ul> <li>Woman of childbearing potential and is not willing to use 2 highly effective methods of contraception while receiving study treatment and for an additional 3 months after the last dose of study treatment.</li> </ul>
	<ul> <li>Male who has a female partner of childbearing potential, and is not willing to use 2 highly effective forms of contraception while receiving study treatment and for at least an additional 3 months after the last dose of study treatment</li> </ul>
Alter	18 Jahre und älter
Molekularer Marker	CD19
Fallzahl	30
Prüfzentren	Innere Medizin 2 (Rekrutierung beendet) Hämatologie / Medizinische Onkologie Theodor-Stern-Kai 7 60590 Frankfurt am Main Studienkoordination GMALL-Molact-1-Blina molact1-blina@med.uni-frankfurt.de
Sponsor	Goethe-Universität Frankfurt
Förderer	AMGEN GmbH
Registrierung in anderen Studienregistern	ClinicalTrials.gov NCT03109093 EudraCT 2015-000733-76
Links	Studiendokumente zum Download (roXtra)
	Zu den Ein- und Ausschlusskriterien