

KURZPROTOKOLL **Compete**

Öffentlicher Titel	Phase III Studie zur Peptid-Rezeptor-Radionuklid-Therapie mit ¹⁷⁷ Lu-edotreotide bei gastroenterologischen oder pankreatischen neuroendokrinen Tumoren
Wissenschaftl. Titel	A prospective, randomised, controlled, open label, multicentre phase III study to evaluate efficacy and safety of Peptide Receptor Radionuclide Therapy (PRRT) with ¹⁷⁷ Lu Edotreotide compared to targeted molecular therapy with Everolimus in patients with inoperable, progressive, somatostatinreceptor positive (SSTR+), neuroendocrine tumours of gastroenteric or pancreatic origin (GEP-NET)
Kurztitel	Compete
Studienart	multizentrisch, prospektiv, Therapiestudie, randomisiert, offen/unverblindet, Pharma-Studie
Studienphase	Phase III
Erkrankung	Drüsen/Hormone/Stoffwechsel: Neuroendokrine Tumoren
Einschlusskriterien	<ul style="list-style-type: none">- Histologically and clinically confirmed diagnosis of well-differentiated neuro-endocrine tumour of non-functional gastroenteric origin (GE-NET) or both functional or non-functional pancreatic origin (P-NET)- Measurable disease per RECIST 1.1- Somatostatin receptor positive (SSTR+) disease- Radiological disease progression, defined as progressive disease per RECIST 1.1. criteria
Ausschlusskriterien	<ul style="list-style-type: none">- Known hypersensitivity to edotreotide or everolimus- Known hypersensitivity to DOTA, lutetium-177, or any excipient of edotreotide or everolimus or any other Rapamycin derivative- Prior exposure to any peptide receptor radionuclide therapy (PRRT)- Prior therapy with mTor inhibitors- Prior EFR (external field radiation) to GEP-NET lesions or radioembolisation therapy- Therapy with an investigational compound and/or medical device within 30 days prior to randomisation- Indication for surgical lesion removal with curative potential- Planned alternative therapy (for the period of study participation)- Serious non-malignant disease- Renal, hepatic, cardiovascular, or haematological organ dysfunction, potentially interfering with the safety of the study treatments- Pregnant or breast-feeding women- Subjects not able to declare meaningful informed consent on their own (e.g. with legal guardian for mental disorders) or any other vulnerable population to that sense (e.g. persons institutionalised, incarcerated etc.)
Alter	18 Jahre und älter
Molekularer Marker	SSTR
Prüfzentren	Universitätsklinikum Gießen und Marburg, Standort Marburg (Rekrutierung beendet) Baldingerstraße 35043 Marburg Mariana May Tel: 06421 58 67022 mariana.may@uk-gm.de
Sponsor	ITM Solucin GmbH
Registrierung in anderen Studienregistern	EudraCT 2016-001897-13 ClinicalTrials.gov NCT03049189 (primäres Register)