## **Eligibility CHECKLIST**

		INVESTI	INVESTIGATOR / CENTRE		
		С	Dr. P.J. Lugtenburg		
	PATIENT INITIALS	DATE OF	BIRTH		
			_][_]/[_][_	_]	
F	PATIENTS SOURCE NUMBER	GENDER	DATE INFO	RMED CON	NSENT
		Male / Female		[_][_] / 20	0[][]
INC	LUSIE CRITERIA				
				YES	NO
1.	Execute an informed consent.				
2.	Patients age ≥18 years at signing the inf				
3.	Diagnosis of relapsed or refractory lymp available therapies.				
4.	Eastern Cooperative Oncology Group (E (Appendix A, Section 15.1).	ECOG) performance stat	:us ≤2		
5.	Neutrophils >1,000 μL				
6.	Platelets ≥75,000 μL				
7.	Aspartate aminotransferase/alanine ami limit of normal (ULN).	`	, ,,		
8.	Total bilirubin <2.0 mg/dL unless elevate	ed due to known Gilbert's	s syndrome.		
9.	Creatinine ≤1.5 ULN.				
	Serum potassium and magnesium within Males and females of child bearing pote				
	willing to use at least two effective forms administration and for at least ninety (90 study drug to be eligible to participate. V must be willing to use a secondary meth abstinence is considered a highly effectifrom heterosexual intercourse during the the study treatment. The reliability of sex in relation to the duration of the clinical to lifestyle of the patient.	) days after the administ assectomized partners at lod of effective birth confeve we method only if define e entire period of risk ass kual abstinence needs to	tration of the nd patients trol. Sexual d as refraining sociated with be evaluated		
INC	LUSIE CRITERIA – <u>ALLEEN</u> voor c	ohort 2: relapsed/ref	ractory Hodgk	in's lymph	oma
		_		YÉS	NO
1.	At least three lines of prior therapy and r with proven clinical benefit.	no other standard therap	y available		
INIC	LUSIE CRITERIA – ALLEEN voor c	ohort 2: rolanced/ref	ractory parinh	oral T call	
	phoma (PTCL)	onort 3. relapsed/ret	ractory periph	erar i-ceil	
7				YES	NO
1.	Only PTCL patients with histologically or AITL or ALCL.	-	·		
2.	At least one line of prior combination the available with proven clinical benefit.	rapy and no other stand	lard therapy		
INC	LUSIE CRITERIA – <u>ALLEEN</u> voor c	ohort 4: relansed/ref	ractory cutane	OUS T-CALL	
	phoma (CTCL), subtypes mycosis				
		` ` ` `		YEŚ	NO
1.	Only CTCL patients with histologically or stage IIb to IVb disease based on modifi	ed ISCL/EORTC staging	g.		
2.	At least one line and a maximum of four no other standard therapy available with		therapies and		

		YES	NO
1.	A maximum of two lines of prior therapy and no other standard therapy available with proven clinical benefit. Patients ineligible for treatment with alemtuzumab can be included.		
FX(	CLUSIE CRITERIA		
	SEGGIE GITTERIA	YES	NO
1.	Patients with any central nervous system (CNS) involvement		
2.	Diagnosis of acute leukemia or any patient that has been treated with fludarabine.		
3.	Any patient who has relapsed within 100 days of stem cell infusion following an allogenic or autologous bone marrow transplant.		
4.	Patients with QTc interval > 450 msec at baseline.		
5.	Patients who are on treatment with drugs known to prolong the QT/QTc interval. Refer to CredibleMeds list of drugs with known risk of Torsade des pointes (TdP): https://crediblemeds.org/new-drug-list		
6.	Any serious medical condition that interferes with adherence to trial procedures.		
7.	Patients with a history of a second malignancy diagnosed within three (3) years of trial enrollment excluding basal cell carcinoma of the skin, squamous cell carcinoma of the skin, or in situ cervical cancer that has undergone potentially curative therapy.		
8.	Pregnant or breast feeding females.		
9.	New York Heart Association (NYHA) stage III/IV congestive heart failure. The following arrhythmias: atrial fibrillation/flutter with poor rate control, documented sustained ventricular tachycardia (defined as >30 seconds or requiring cardioversion before 30 seconds have elapsed) or TdP.		
10.	Active infections, or other significant co-morbidities (e.g., active central nervous system metastases and/or carcinomatous meningitis, active infection requiring systemic therapy, history of human immunodeficiency virus (HIV) infection, or active Hepatitis B or Hepatitis C).		
11.	Previous cancer therapies within four (4) weeks or 5 half-lives, whichever is shorter, of dosing unless the patient has recovered to eligibility levels prior to treatment in this study.		
12.	Use of other investigational agents within 30 days or 5 half-lives prior to the first dose of study drug unless patient has recovered from any related toxicities ≥ Grade 1.		
13.	Steroid treatment within seven (7) days prior to study treatment. Patients that require intermittent use of bronchodilators, topical steroids or local steroid injections will not be excluded from the trial. Patients who have been stabilized to 10 mg PO QD or less seven (7) days prior to trial drug administration are allowed.		
14.	Patients on Valproic Acid for any indication (epilepsy, mood disorder) must be excluded from the trial or must stop using the medication and have a wash out period of 3.5 days prior to first dose of EDO-S101 (C1D1).		
cke	ed by:Signature		

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