

Public Title	JNJ-56022473 bei MDS und AML nach Versagen einer Therapie basierend auf hypomethylierenden Substanzen
Short Title	SAMBA
Id KN/ELN	LN_DEUTSC_2016_604
Trialgroup	Deutsche MDS
Type of Trial	multicentric, single-group, open-label
Phase	Phase II
Disease	Myelodysplastic Syndrome(MDS) Low risk and intermedia I Myelodysplastic Syndrome(MDS) Intermedia II and high risk Myelodysplastic Syndrome(MDS) RAEB I Myelodysplastic Syndrome(MDS) RAEB II
Stage of Disease	.
Outcomes	- To assess the efficacy of JNJ-56022473 for the treatment of MDS and AML patients who have relapsed after or are refractory to treatment with HMAs To assess the efficacy of JNJ-56022473 for the treatment of MDS and AML patients who have relapsed after or are refractory to treatment with HMAs (Primary Outcome)
Inclusion Criteria	<ul style="list-style-type: none">- 18 years of age- Diagnosis of AML or MDS- At least 5% BM blasts at the time of screening (done by central morphology)- At least one cytopenia (ANC < 1800/L or platelet count < 100,000/L or hemoglobin < 10 g/dL)- Failure to achieve complete or partial response or hematological improvement after at least six (azacitidine) or four (decitabine) 4-week treatment cycles administered during the past two years OR- Relapse after initial complete or partial response or hematological improvement observed after at least six (azacitidine) or four (decitabine) 4-week treatment cycles administered during the past two years OR- Intolerance to treatment with HMA (hypomethylating agents) defined by drug-related Grade 3 liver or renal toxicity leading to treatment discontinuation during the past two years- Failed to respond to, relapsed following, not eligible, or opted not to participate in bone marrow transplantation- Off all other treatments for AML/MDS for at least four weeks; Filgrastim (G-CSF) and erythropoietin are allowed before and during the study as clinically indicated- No medical need for or patient opted not to receive induction chemotherapy- ECOG performance status of 0-2- Willing to adhere to the prohibitions and restrictions specified in the protocol- Signed informed consent
Exclusion Criteria	<ul style="list-style-type: none">- Previous treatment with a CD123 agent or T- or NK cell redirecting therapy- Patients having received intensive chemotherapy to treat HMA failure- Diagnosis of acute promyelocytic leukemia (APL)- WBC > 15 GPT/L- Any active malignancy within the past year, except basal cell or squamous cell skin cancer or carcinoma in situ of the cervix or breast- Uncontrolled intercurrent illness including, but not limited to, symptomatic congestive heart failure, unstable angina pectoris, or cardiac arrhythmia- Active infection not adequately responding to appropriate therapy

- Total bilirubin > 1.5 mg/dL not related to hemolysis or Gilbert's disease
- ALT/AST > 2.5 x upper limit of normal
- Serum creatinine > 2.0 mg/dL
- Patients who are unwilling to follow strict contraception requirements (including condom use for males with sexual partners, and for females: prescription oral contraceptives, contraceptive injections, intrauterine device, double-barrier method, contraceptive patch, or surgical sterilization) before entry, throughout the study and within 3 months after last study drug administration
- Female patients with reproductive potential who do not have a negative urine β -HCG pregnancy test at screening and prior to the first study drug administration
- Female patients with reproductive potential who do not have a negative urine beta-HCG pregnancy test at screening and prior to the first study drug administration
- Female patients who are lactating

Age	>= 18 years
Status	Closed
start of Recruitment	01.09.2016
Target Sample Size	43
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Supporters	Janssen
Other Registers	European Clinical Trials Database - EUDRACT2016-000327-10