KURZPROTOKOLL AMLSG 21-13

Öffentlicher Titel Wissenschaftl. Titel Dasatinib bei Patienten mit neu diagnostizierter Core-Binding Factor AML

Randomized Phase III Study of Intensive Chemotherapy with or without Dasatinib (Sprycel™) in Adult Patients with Newly Diagnosed Core-Binding Factor Acute Myeloid

Leukemia (CBF-AML)

Studienart

AMLSG 21-13 Kurztitel multizentrisch, Therapiestudie, randomisiert, offen/unverblindet, zweiarmig

Studienphase

Phase III

Erkrankung

Blut: Akute myeloische Leukämie (AML): Neu diagnostiziert / de novo

Ziele

- To assess event-free survival (EFS) after intensive induction (daunorubicin and cytarabine) and consolidation (high-dose cytarabine) chemotherapy with or without dasatinib in patients with CBF-AML
- To assess the interaction between type of CBF-AML [t(8;21) versus inv(16)] and randomization accordingly on all survival endpoints
- To assess cumulative incidence of relapse (CIR) and death (CID)
- To assess relapse-free (RFS) and overall survival (OS)
- To assess outcome according to KIT mutational status
- To assess pharmacodynamic inhibition of KIT
- To assess toxicity

Einschlusskriterien

- Core-binding factor (CBF) AML with molecular diagnosis of RUNX1-RUNX1T1 fusion transcript resulting from t(8;21)(q22;q22) (or a variant form) or of CBFB-MYH11 fusion transcript resulting from inv(16)(p13.1q22)/t(16;16)(p13.1;q22) as assessed in one of the central AMLSG reference la-boratories (Ulm, Hannover)
- Age 18; there is no upper age limit
- No prior chemotherapy for leukemia except hydroxyurea for up to 5 days during the diagnostic screening phase
- Non-pregnant and non-nursing. Due to the unknown teratogenic potential of dasatinib in humans, pregnant or nursing patients may not be enrolled. Women of childbearing potential (WOCBP) must have a negative serum or urine pregnancy test within a sensitivity of at least 25 mIU/mL with-in 72 hours prior to registration. Women of childbearing potential must either commit to contin-ued abstinence from heterosexual intercourse or begin TWO acceptable methods of birth control - one highly effective method (e.g., IUD, hormonal, tubal ligation, or partner's vasectomy), and one additional effective method (e.g., latex condom, diaphragm, or cervical cap) - AT THE SAME TIME, at least four weeks before she begins dasatinib therapy and at least 3 months after last dasatinib administration. "Women of childbearing potential" is defined as a sexually active mature woman who has not undergone a hysterectomy or who has had menses at any time in the preced-ing 24 consecutive months.
- Men must agree not to father a child and must use a latex condom during any sexual contact with women of childbearing potential while taking dasatinib and for 3 months after therapy is stopped, even if they have undergone a successful vasectomy.
- Signed written informed consent.

Ausschlusskriterien

- Performance status WHO >2
- Pulmonary edema and/or pleural/pericardial effusion within 14 days of day 1. If edema/effusion resolves to CTC Grade 1, patients can be treated with dasatinib.
- Patients with ejection fraction <50% by echocardiography within 14 days of day 1
- Organ insufficiency (creatinine >1.5x upper normal serum level; bilirubin, AST or AP >2.5x upper normal serum level; heart failure NYHA III/IV; severe obstructive or restrictive ventilation disorder)
- Uncontrolled infection

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- Patients with a "currently active" second malignancy other than non-melanoma skin cancers. Pa-tients are not considered to have a "currently active" malignancy, if they have completed therapy and are considered by their physician to be at less than 30% risk of relapse within one year.
- Severe neurological or psychiatric disorder interfering with ability of giving an informed consent
- Known positive for HIV, active HBV, HCV, or Hepatitis A infection
- Bleeding disorder independent of leukemia
- No consent for registration, storage and processing of the individual disease characteristics and course as well as information of the family physician and/or other physicians involved in the treatment of the patient about study participation.
- No consent for biobanking.

Alter 18 Jahre und älter

Prüfzentren Klinikum Darmstadt GmbH (Rekrutierung beendet)

Grafenstraße 9 64283 Darmstadt Cora Schwebel-Kottke Tel: 06151 1076670 Fax: 06151 1076676

Onko-Studienzentrum@mail.klinikum-darmstadt.de

Sponsor Universität Ulm

Registrierung in anderen ClinicalTrials.gov NCT02013648 (primäres Register)

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