

Open label, comparative, randomized, multicenter, study of trastuzumab given with docetaxel versus sequential single agent therapy with trastuzumab followed by docetaxel as first-line treatment for Her2neu+++ metastatic breast cancer patients.

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Ethische beoordeling	Positief advies
Status	Werving gestopt
Type aandoening	-
Onderzoekstype	Interventie onderzoek

Samenvatting

ID

NL-OMON26229

Bron

NTR

Verkorte titel

HERTAX, BOOG 2002-02

Aandoening

Breast cancer patients with metastases with HER2neu overexpression (3+ assessed by IHC DAKO HercepTest), previously untreated by chemotherapy, except for neoadjuvant or adjuvant (non-taxane containing) chemotherapy.

Ondersteuning

Primaire sponsor: BOOG

Overige ondersteuning: SanofiAventis

Roche

Onderzoeksproduct en/of interventie

Uitkomstmaten

Primaire uitkomstmaten

Progression free survival of total sequential versus combined treatment.

Toelichting onderzoek

Achtergrond van het onderzoek

N/A

Doel van het onderzoek

Although combined treatment will probably lead to higher response rates, sequential treatment may result in a similar time to progression in the presence of less side effects and a better quality of life in a significant number of patients.

Onderzoeksopzet

N/A

Onderzoeksproduct en/of interventie

Arm A: Comb. of trastuzumab + docetaxel;

Arm B: Trastuzumab followed by docetaxel.

Trastuzumab:

Loading dose of 4 mg/kg IV on day 1, administered as 90-minute infusion, followed by a weekly dose of 2 mg/kg.

Docetaxel:

TXT 100 mg/m² IV infusion over one hour repeated in cycles, every 3 weeks for 6 cycles.

Contactpersonen

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Wetenschappelijk

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Deelname eisen

Belangrijkste voorwaarden om deel te mogen nemen (Inclusiecriteria)

1. Histologically documented invasive adenocarcinoma of the breast;
2. Women with previously chemotherapeutically untreated metastatic breast cancer with HER2neu overexpression (defined as 3+ IHC by DAKO HercepTest);
3. Patients having previously received adjuvant treatment with an anthracycline/anthraquinone (maximum cumulative dose:
doxorubicin 360 mg/m², epirubicin 750 mg/m² or equivalent dose of other anthracycline/anthraquinone);
4. Patients over the age of 18;
ECOG performance status < = 2 and life expectancy >12 weeks;

5. Patients with evaluable disease or patients having at least one measurable target outside previously irradiated field;
6. Adequate bone marrow, hepatic and renal functions as evidenced by the following;
7. Hemoglobin > 6 mmol / l and no blood transfusion within the previous 2 weeks;
8. WBC count > 3.0×10^9 cells/l and neutrophils > 1.5×10^9 cells/l;
9. Platelets count > 100×10^9 cells/l;
10. No evidence of myelodysplastic syndrome or abnormal bone marrow reserve;
11. Creatinine < 1.5 upper normal limit (UNL) or creatinine clearance > 60 ml / min;
12. Total bilirubin < 1 x UNL;
13. ASAT (SGOT) and/or ALAT (SGPT) <2.5 x UNL;
14. Alkaline phosphatase < 5 x UNL;
15. ASAT and/or ALAT< 1.5 x UNL in combination with elevated alkaline phosphatase < 2.5 x UNL;
16. Previous radiotherapy is allowed if :
End of radiotherapy more than 14 days prior to study entry, in case RT was given on relevant areas;
17. Patient has fully recovered from all acute toxic effects;
18. Normal Cardiac Function with LVEF by ECHO or MUGA > 50% or within UNL of the institution;
19. Written informed consent and accessible for treatment and follow up.

Belangrijkste redenen om niet deel te kunnen nemen (Exclusiecriteria)

1. Operable local relapse alone after conservative treatment or contra-lateral tumour, (mastitis or inoperable local recurrence is acceptable for inclusion);
2. Pregnant or lactating women (females of childbearing potential must use adequate contraception);
3. History or presence of brain or leptomeningeal metastases;

4. Current peripheral neuropathy
5. Other prior malignancies, except for cured non melanoma skin cancer, curatively treated in situ carcinoma of the cervix;
6. Other serious illness or medical condition:

Cardiac insufficiency (NYHA III or IV), myocardial infarction within previous 6 months, unstable angina pectoris, uncontrolled arrhythmia at time of inclusion;
7. Patients with severe dyspnoea at rest due to complications of advanced malignancy or requiring supplementary oxygen therapy;
8. Clinically significant active infections;
9. Poorly controlled diabetes mellitus;
10. Uncontrolled hypertension;
11. Active peptic ulcer or other contraindication to high dose of corticosteroid therapy such as herpes zoster, cirrhosis;
12. History of allergy to drugs containing polysorbate 20, or the excipient TWEEN 80;
13. Patient with a history of a psychological illness or condition such as to interfere with the patients ability to understand the requirements of the study;
14. Patients who had received an investigational new drug within the last 30 days;
15. Patients having received prior therapy with taxoids or anti-HER2 therapies.

Onderzoeksopzet

Opzet

Type:	Interventie onderzoek
Onderzoeksmodel:	Parallel
Toewijzing:	Gerandomiseerd
Blinding:	Open / niet geblindeerd
Controle:	Geneesmiddel

Deelname

Nederland
Status: Werving gestopt
(Verwachte) startdatum: 01-02-2003
Aantal proefpersonen: 100
Type: Werkelijke startdatum

Ethische beoordeling

Positief advies
Datum: 09-09-2005
Soort: Eerste indiening

Registraties

Opgevolgd door onderstaande (mogelijk meer actuele) registratie

Geen registraties gevonden.

Andere (mogelijk minder actuele) registraties in dit register

Geen registraties gevonden.

In overige registers

Register	ID
NTR-new	NL270
NTR-old	NTR308
Ander register	: N/A
ISRCTN	ISRCTN13770586

Resultaten

Samenvatting resultaten

N/A