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TITLE: Randomized Trial of Interleukin-2 (IL-2) as Early Consolidation Following Marrow Ablative Therapy with Stem Cell Rescue for Metastatic Breast Cancer

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<td>Marrow ablative doses of chemotherapy followed by stem cell rescue (MAT/SR) produces a high frequency of objective responses in patients with metastatic breast cancer, with up to 40-50% complete responses. Unfortunately, responses tend to be short-lived. Interleukin-2 (IL-2) has the capacity to activate lymphocytes to kill multidrug resistant cancer cells. Our phase I data established the feasibility of administering a single course of low-dose IL-2 (1.6 million IU/m2/day as a continuous i.v. infusion for 18 days) as consolidation treatment to patients with metastatic breast cancer early after intensive chemotherapy. Seven patients (60%) remained in complete remission at a median of &gt;435 days post stem cell transplantation. We are therefore performing a phase II trial of AC+T chemotherapy followed by IL-2 consolidation (1 cycle as described above) in high risk stage II and III breast cancer patients. The goal is to kill residual chemotherapy-resistant cancer cells. Disease free survival and toxicity assessment represent major clinical aims (Specific aim 1). Immunologic effector mechanisms induced following MAT/SR by IL-2 infusion will be evaluated using phenotypic and functional assays for LAK cell induction (Specific Aim 2). Accrual to this study has been delayed due to a change from a randomized trial to a single arm phase II study and due to negotiations between the University of Utah and the Army NRM C concerning liability clauses in the consent document.</td>
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Introduction:

At least 46,000 women die from metastatic breast cancer each year in the United States. Median survival remains 12-18 months from the diagnosis of metastatic disease, and progression-free survival beyond 5 years is rare (<10%). This has led to the testing of escalated, marrow ablative doses of chemotherapy followed by stem cell rescue (MAT/SR). This approach produces a high frequency of objective responses in patients with metastatic breast cancer, with up to 40-50% complete responses. Unfortunately, responses tend to be short-lived, and only a minority of women (10-20%) achieve long-term disease free survival. Relapses may be due to both minimal tumor contamination of stem cells rein infused into patients, as well as residual chemotherapy resistant tumor cells not cleared by the MAT/SR regimen. IL-2 activated lymphocytes, termed lymphokine-activated killer (LAK) cells have significant cytotoxic activity against autologous breast cancer cells and breast cancer cell lines. Our own studies have demonstrated that multidrug-resistant tumor cells remain sensitive to LAK cell mediated killing. We have completed a phase I study to test the feasibility of administering a single course of low-dose IL-2 (1.6 million IU/m²/day as a continuous i.v. infusion) as consolidation treatment to patients with metastatic breast cancer early after MAT/SR. This study established that IL-2 consolidation could be safely begun starting on day +14 post MAT/SR with minimal toxicity. Substantial LAK cell induction was observed, using flow cytometric and cytotoxicity assays. Thus far, only 3 of 10 patients have had breast cancer relapse or progression, and a small second breast cancer was detected in 1 patient. Seven patients (60%) remain in complete remission at a median of 435 days (range: 224 - 720 days) post stem cell transplantation. Because patients with metastatic breast cancer transplanted with active disease have a 60% and 80% probability of relapse at 1 year and 3 years, respectively (without IL-2), we proposed to test this promising immunotherapy consolidation strategy in a single-institution randomized prospective trial. We propose to perform cyto reduction in patients with metastatic breast cancer using MAT/SR, followed by continuous infusions of low-dose IL-2 starting on day +14 to activate lymphocytes to kill residual chemotherapy-resistant cancer cells. Based on preliminary data, we hypothesize that a single course of IL-2 will result in a significant improvement in disease-free survival, with minimal toxicity. Effectiveness of this approach may correlate with the effective induction of LAK precursor and effector cells, as well as evidence for reduction in the burden of minimal residual cancer cells. In Specific Aim 1 we originally proposed to perform a prospective randomized clinical trial to test whether the addition of 1 cycle of continuous i.v. infusion of IL-2 in women with metastatic breast cancer, starting on day +14 after treatment with MAT/SR, can increase progression-free and overall survival by 25%. In Specific Aim 2 we will evaluate possible immunologic effector mechanisms induced following MAT/SR and IL-2 infusion. Phenotypic and functional assays for LAK cell induction and enzyme immunoassays for circulating pro-inflammatory cytokines will be performed. Following review by the US Army, Specific Aim 3 (detection of residual tumor cells in bone marrow and stem cell products by flow cytometry and RT-PCR) was omitted.
Body:
Patient accrual to our proposed clinical trial has been severely delayed due to four unanticipated events. First, shortly after this proposal was funded in 1999, a series of randomized trials was reported at the American Society of Clinical Oncology meetings in 5/00 comparing standard dose chemotherapy and marrow ablative therapy and stem cell rescue (MAT/SCR) for treatment of advanced breast cancer. The conclusions of all but one of these trials was that there was no advantage to stem cell transplants in breast cancer patients over standard chemotherapy(1-3). The second event was that the one trial showing benefit of MAT/SCR over chemotherapy (Bezwoda, et al) was found to contain fraudulent data(4). In combination, these findings made the proposed control arm of our randomized clinical trial (MAT/SCR alone) unacceptable. Since the goal of MAT/SCR in our trial was to provide maximal cytoreduction prior to IL-2 based immunotherapy, this goal was still felt scientifically reasonable, given our impressive phase I trial results. In order to further prove the validity of these observations, it was felt by Dr. Peterson and myself that a change from a randomized trial to a single arm phase II study (MAT/SCR followed by an 18 day infusion of IL-2) was warranted. This change was discussed with the USAMRMC and the study protocol and consent documents were rewritten. A third point holding up the clinical trials was due to negotiations between the University of Utah lawyers and the USAMRMC concerning required liability clauses in the consent document and final approval by the University of Utah IRB. After many months of negotiations, a finalized consent language and protocol was agreed upon. A final draft has been submitted to the University of Utah IRB and was approved with minor revisions. A fourth event has necessitated further revisions in the study design. Further investigation into fraudulent data published by Bezwoda was presented at the ASCO meetings in 5/01. This resulted in the general abandonment of MAT/SCR as a breast cancer treatment modality in the United States. At the end of August of 2001, I was notified by my co-investigator Dr. Peterson that we would not be able to accrue any patients to a MAT/SCR regimen based trial, since patient referral to the University of Utah BMT program had dropped to nothing over the summer.

We have spent the intervening month discussing options. Due to our exciting preliminary results, we still strongly believe that the concept of IL-2 consolidation in high risk breast cancer should be tested after maximal cytoreduction. Given the apparent equivalence of MAT/SCR and standard chemotherapy in high risk breast cancer patients, we have concluded that an alternative method to test our hypothesis is to enroll high-risk breast cancer patients who are treated systemically with surgery, followed by a standard chemotherapy regimen (doxorubicin/cyclophosphamide followed by paclitaxel), followed by a 21 day infusion of IL-2. Patients will subsequently receive irradiation to the breast and regional node areas. Patients deemed at high risk include: patients with ≥4 lymph nodes positive (40% 5 year survival with ≥4+ nodes, <20% 5 year survival with ≥10 nodes involved), inflammatory breast cancer (<20% 5 year survival) and patients with resected stage IV disease (Stage IV NED, <10-20% 5 year survival)(5).

Dr. Peterson will decrease his effort to 5% (since MAT/SCR will no longer be a primary treatment modality) and we will add two Breast Cancer Medical Oncology specialists (Dr. John H. Ward and Dr. Saundra Buys), from the Huntsman Cancer Institute as co-
investigators to ensure adequate accrual. Since this trial is compatible with the community standard of breast cancer management in the Intermountain Region, the trial will be opened to Huntsman Cancer Institute affiliated community oncologists to increase the pace of accrual. It is envisioned that patient accrual for the proposed phase II trial (60 patients) can be completed in 36 months. The Huntsman Cancer Institute Clinical Trials Office will provide the resources for the long-term follow-up patients beyond the scope of the current grant funding. This revised protocol has been completed and has been approved by the University of Utah IRB. It has been recently reviewed by US Army Medical Research and Material Command (HSRRB Log Number A-9034). Final revisions have been made and have been resubmitted to the University of Utah IRB which has approved these changes. The protocol has now been resubmitted to the HSRRB for final go-ahead for patient entry. The protocol will then open (estimated 2-4 weeks). Having overcome these unanticipated and frustrating delays, we remain enthusiastic to test the scientific hypothesis that IL-2 consolidation following chemotherapy of high risk breast cancer will produce long term disease free survival in >40% of patients with acceptable toxicity.

Personnel required for the start of this clinical trial are all in place, ready to being patient accrual once final USAMRMC approval are obtained. All methodologies required for patient IL-2 treatment and sample analysis for LAK cell induction on human samples have been worked out and are ready for use in this trial. Our proposal is to add an additional 2 years onto study accrual, due to the delayed start of the trial, as a no-cost extension, using funds carried over from preceding years (with additional support from the Huntsman Cancer Institute).
Task 1: Patient Enrollment: (months 1-36-start delayed to month 36)
- Protocol will be presented to eligible patients prior to chemotherapy for high risk breast cancer. Eligible patients will include ≥4+ nodes, inflammatory breast cancer and completely resected stage IV disease. All patients will be treated with at least 4 cycles of doxorubicin/cyclophosphamide and 4 cycles of paclitaxel chemotherapy (AC+T) at 3-4 week intervals.
- Appropriate people/departments will be notified of patient enrollment and randomization, to include: site pharmacy, Dr. Wolf Samlowski and/or lab and Dr. Wayne Green and/or lab.
- A Progress Note will be entered into the patient’s medical record regarding patient consent, enrollment, randomization, and the study requirements.

Task 2: Administration of IL-2: (months 1-36 from revised start date)
- On Day +14 following completion of 4 cycles of AC+T, the patient will come to clinic (if discharged), where they will be have their vital signs taken and be evaluated by a physician or physician extender for a baseline physical exam.
- The CADD-1 pump and supplies will be reviewed with the patient and caregiver(s).
- IL-2 will be started and the patient will remain in clinic for at least one hour to monitor vital signs and any adverse reactions.
- A patient diary will be given to the patient to help monitor and track fevers, other reactions, admissions, etc. (See attached sample.)
- Patient will be seen in clinic a minimum of once per week and also as needed. IL-2 cassettes will be changed every six days by the Research Nurse. Review of any adverse reactions or other problems will take place.

Task 3: Specimen Collection: (months 1-44 from revised start date)
- Approximately 50 cc’s of blood will be collected in heparinized, green top tubes on Day 0, +7, +14, +21, +32 and +100 and delivered to Dr. W. Samlowski’s and Dr. W. Green’s labs. (See below for details of lab procedures)
- Samples of pre-transplant and day 100 bone marrow material, as well as stem cell products will be transported to Dr. W. Samlowski’s lab for evaluation of minimal residual tumor cells (5 ml marrow or PBSC cells in a heparinized syringe).

Task 4: Analysis of LAK cell induction (months 1-44 from revised start date)
- Samples will be analyzed for T cell and LAK cell markers by flow cytometry (Dr. Green's lab)
- Analysis of patient samples for LAK precursor and cytolytic cell function will be performed (Dr. Samlowski's lab)

Task 5: Assays for tumor cell detection in bone marrow and stem cell products (deleted from funding by USAMRMC)

Task 6: Data Collection (months 1-48 from revised start date):
- The following will be collected: dates of each chemotherapy cycle and treatment doses, and side effects. While patients are receiving IL-2 infusions, side effects of
IL-2 (fevers, rash, etc), infections, readmission, relapse, death and other
significant events will be recorded.
• At day 100, the patient will be seen by the physician to evaluate their disease and
health status. Information such as infections, readmission, relapse and death will
be collected.
• Patients will then be followed at least every 3 months or as needed to monitor
disease status and/or death.

Task 7: Interim Analysis: (approximately month 24 from revised start date)
• After approximately 30 patients are enrolled, data collected from Dr. Wolf
Samlowski’s lab and Dr. Wayne Green’s lab, together with information collected
in the CRF’s will be analyzed by the principal investigators.

Task 8: Final Analysis: (month 60 from revised start date)
• After enrollment of 60 patients is complete, data collected from Dr. Wolf
Samlowski’s lab and Dr. Wayne Green’s lab and information collected in the
CRF’s will be analyzed by the principal investigators. At the completion of this
study a report will be generated.

Key research accomplishments:

We treated 20 patients with MAT/SCR in our phase I pilot trial. Patients received IL-
2 either starting on day 1 (10 patients) or day 14 (10 patients) following stem cell
infusion. A total of 17 patients were evaluable for response at the time of initial analysis.
A total of 17 patients (85%) completed the IL-2 course. Three patients receiving IL-2
from day 1 required IL-2 infusions to be terminated early (2 fever, 1 thrombocytopenia).
Relapse free survival was 45% with 580 day median follow-up (135-1175 days), with
75% overall survival.

LAK cell activation was evaluated in patients undergoing IL-2 infusions starting either
day 1 (5 patients) or day 14 post stem cell infusion (5 patients). Cytotoxicity against the
MCF-7 breast cancer line was detected in all patients, regardless of whether the IL-2
infusion was started day 1 or 14. Increased cytolytic activity was detected in cytotoxicity
assays performed with the addition of IL-2, suggesting a substantial increase in
circulating LAK cell precursors in both patient populations. Phenotypic evaluation
established that while CD56+ cell populations were expanded in both patient groups, the
absolute number of circulating CD56+ cells was 10-fold higher in patients receiving IL-2
starting on day 14.

Due to these results, our current clinical trial will treat patients beginning on day +14
after completion of 4 cycles of AC and 4 cycles of Taxol with a 18 day infusion of IL-2
to verify these exciting clinical results in this high-risk breast cancer population.

Reportable outcomes:

We have published the results of our phase I trial in abstract form (copies enclosed):

A manuscript for submission to a peer reviewed journal (J. Clinical Oncology) is currently being prepared.

Conclusions:

The proposed use of IL-2 following maximal cytoreduction of tumor by standard chemotherapy or MAT/SCR remains promising based on our preliminary data, with 45% of patients achieving >2 year disease free survival after MAT/SCR plus IL-2. As soon as regulatory approval of the revised protocol and consent documents are obtained, we will perform a phase II pilot study to evaluate the effectiveness of this regimen with standard chemotherapy in a prospective phase II trial.

References:

1) Stadtmauer EA et al. Phase III randomized trial of high dose chemotherapy and stem cell support shows no difference in overall survival or severe toxicity compared to maintenance chemotherapy with cyclophosphamide, methotrexate and 5-fluorouracil (CMF) for women with metastatic breast cancer who are responding to standard induction chemotherapy. Proc ASCO 18:1a, 1999.


3) Scandinavian Breast Cancer Study Group. Results of a randomized adjuvant breast cancer study with CTCb supported by autologous bone marrow stem cells versus dose-escalated and tailored FEC therapy. Proc ASCO 18:2a, 1999.


FEASIBILITY OF LOW-DOSE CONTINUOUS INFUSION OF IL-2 AS A CONSOLIDATION TREATMENT FOLLOWING INTENSIVE BREAST CANCER CHEMOTHERAPY

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Patients with regionally advanced or metastatic breast cancer continue to have a high risk for death from metastatic disease despite the use of intensive chemotherapy. This may be due to persistence of chemotherapy resistant tumor cells. Interleukin-2 (IL-2) has been shown to induce cytolytic lymphocytes, termed lymphokine activated killer (LAK) cells that can kill multidrug resistant cancer cells in vitro. We therefore tested the feasibility of using a single course of low-dose continuous i.v. infusion of IL-2 as consolidation treatment after intensive cytoreductive chemotherapy. To test this concept, we performed a phase I/II trial in patients with limited metastatic breast cancer. Patients received the Stamp-V marrow ablative regimen, followed by stem cell rescue. In phase I, patients received continuous i.v. infusion of IL-2 (1.8 million IU/m2/d for 18 days) starting either day 1 (5 patients) or day 14 (5 patients) following stem cell reinfusion. Endpoints evaluated included toxicity and LAK cell activation. We found that IL-2 infusions starting on day 1 resulted in a high frequency of febrile responses, along exacerbation of thrombocytopenia. A day 14 start was better tolerated with almost no detectable toxicity. Circulating lymphokine-activated killer cells could be detected in all patients, regardless of the day 14 start. To further assess clinical activity of this consolidation regimen, a total of 20 patients were treated with IL-2 consolidation beginning on day 14 post stem cell infusion (Phase II). Our data demonstrates that the majority of patients (85%) were able to complete the entire planned 18 day infusion of IL-2. Three patients ended the IL-2 infusion early due to severe malaise or rash. A total of 9/17 (45%) patients remained in clinical complete remission with a mean follow-up of >580 days (range 135-1175). Based on this data, we have initiated a clinical trial of IL-2 consolidation following doxorubicin/cyclophosphamide plus paclitaxel adjuvant chemotherapy for high-risk regional breast cancer patients. Due to the high eventual risk of death due to breast cancer in this patient population, improved treatment strategies, such as IL-2 consolidation, are urgently needed.